# LS-DYNA ${ }^{\circledR}$ KEYWORD USER'S MANUAL 

## VOLUME III

## Multiphysics Solvers

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## Ansys

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## LS-DYNA MULTIPHYSICS USER'S MANUAL INTRODUCTION

In this manual, there are five main solvers: two compressible flow solvers, an incompressible flow solver, an electromagnetism solver, and a battery electrochemistry solver. Each of them implements coupling with the structural solver in LS-DYNA.

The keywords covered in this manual fit into one of three categories. In the first category are the keyword cards that provide input to each of the multiphysics solvers that in turn couple with the structural solver. In the second category are keyword cards involving extensions to the basic solvers. Presently, the chemistry and stochastic particle solvers are the two solvers in this category, and they are used in conjunction with the *CESE compressible flow solver discussed below. In the third category are keyword cards for support facilities. A volume mesher that creates volume tetrahedral element meshes from bounding surface meshes is one of these tools. Another is a data output mechanism for a limited set of variables from some of the solvers in this manual. This mechanism is accessed through *LSO keyword cards.

The CESE solver is a compressible flow solver based upon the Conservation Element/Solution Element (CE/SE) method, originally proposed by Chang of the NASA Glenn Research Center. This method is a novel numerical framework for conservation laws. It has many non-traditional features, including a unified treatment of space and time, the introduction of separate conservation elements (CE) and solution elements (SE), and a novel shock capturing strategy without using a Riemann solver. This method has been used to solve many types of flow problems, such as detonation waves, shock/acoustic wave interaction, cavitating flows, supersonic liquid jets, and chemically reacting flows. In LS-DYNA, it has been extended to also solve fluid-structure interaction (FSI) problems. It does this with two approaches. The first approach solves the compressible flow equations on an Eulerian mesh while the structural mechanics is solved on a moving mesh that moves through the fixed CE/SE mesh. In the second approach (new with this version), the CE/SE mesh moves in a fashion such that its FSI boundary surface matches the corresponding FSI boundary surface of the moving structural mechanics mesh. This second approach is more accurate for FSI problems, especially with boundary layers flows. Another feature with the CESE moving mesh solver is conjugate heat transfer coupling with the solid thermal solver. The chemistry and stochastic particle solvers are two addon solvers that extend the CESE solver.

The dual CESE solver is another compressible flow solver that is also based upon the Conservation Element/Solution Element (CE/SE) method, but with improvements related to accuracy and robustness. This method follows a similar novel numerical
framework for conservation laws. In LS-DYNA, the dual CESE solver also include fluidstructure interaction (FSI) capabilities. It also does this with two approaches. The first approach solves the compressible flow equations on an Eulerian mesh while the structural mechanics is solved on a moving mesh that moves through the fixed dual CE/SE mesh. In the second approach. the dual CE/SE mesh moves in a fashion such that its FSI boundary surface matches the corresponding FSI boundary surface of the moving structural mechanics mesh. This second approach is more accurate for FSI problems, especially with boundary layers flows. One of the advances in these FSI computations with the dual CESE solver as compared with the older CESE solver is that each FSI approach (or fixed mesh Eulerian solver) may be employed in different subregions of the fluid mesh in the same problem. Unlike the *CESE solvers, the dual CESE solvers do not yet have conjugate heat transfer coupling with the solid thermal solver, nor coupling with the chemistry or stochastic particle solvers. Another advance available only with the dual CESE solvers is the availability of equations of state for pure and pseudo-pure fluids of industrial interest in the REFPROP and COOLPROP EOS libraries. These complex EOSes are generally expensive to evaluate, so a bi-cubic table look-up mechanism has been developed that greatly accelerates their use.

The third solver is the incompressible flow solver (ICFD) that is fully coupled with the solid mechanics solver. This coupling permits robust FSI analysis via either an explicit technique when the FSI is weak, or using an implicit coupling when the FSI coupling is strong. In addition to being able to handle free surface flows, there is also a bi-phasic flow capability that involves modeling using a conservative Lagrangian interface tracking technique. Basic turbulence models are also supported. This solver is the first in LSDYNA to make use of a new volume mesher that takes surface meshes bounding the fluid domain as input (*MESH keywords). In addition, during the time advancement of the incompressible flow, the solution is adaptively re-meshed as an automatic feature of the solver. Another important feature of the mesher is the ability to create boundary layer meshes. These anisotropic meshes become a crucial part of the model when shear stresses are to be calculated near fluid walls. The ICFD solver is also coupled to the solid thermal solver using a monolithic approach for conjugate heat transfer problems.

The fourth solver is an electromagnetics (EM) solver. This module solves the Maxwell equations in the Eddy current (induction-diffusion) approximation. This is suitable for cases where the propagation of electromagnetic waves in air (or vacuum) can be considered as instantaneous. Therefore, the wave propagation is not solved. The main applications are Magnetic Metal Forming, bending or welding, induced heating, ring expansions and so forth. The EM module allows the introduction of a source of electrical current into solid conductors and the computation of the associated magnetic field, electric field, as well as induced currents. The EM solver is coupled with the structural mechanics solver (the Lorentz forces are added to the mechanics equations of motion), and with the structural thermal solver (the ohmic heating is added to the thermal solver as an extra source of heat). The EM fields are solved using a Finite Element Method (FEM) for the conductors and a Boundary Element Method (BEM) for the surrounding air/insulators. Thus no air mesh is necessary.

The fifth solver is a battery electrochemistry solver. At this time, the available capability involves a one-dimensional electrochemistry solver that is coupled to the structural mechanics and structural thermal solver in each structural element identified as being part of a battery cell. It solves these one-dimensional models implicitly so as to be able to run simulations for very long physical times that are typical of battery-structure interaction problems.

As stated above, the *CHEMISTRY and *STOCHASTIC cards are only used in the CESE solver at this time.

## *BATTERY

The keyword *BATTERY provides input data for the electrochemistry solver:

*BATTERY_DATABASE_HISTORY_ALLNDS_ON_ELESET<br>*BATTERY_DATABASE_HISTORY_GLOBALS<br>*BATTERY_DATABASE_HISTORY_NODELIST_ON_ELESET<br>*BATTERY_ECHEM_CELL_GEOMETRY<br>*BATTERY_ECHEM_CONTROL_SOLVER<br>*BATTERY_ECHEM_INITIAL<br>*BATTERY_ECHEM_MAT_ANODE<br>*BATTERY_ECHEM_MAT_CATHODE<br>*BATTERY_ECHEM_MAT_ELECTROLYTE<br>*BATTERY_ECHEM_THERMAL

For now, the available capability involves a one-dimensional electrochemistry solver that is coupled to the structural mechanics and structural thermal solver in each structural element identified as being part of a battery cell.

## *BATTERY_ECHEM_CELL_GEOMETRY

Purpose: Set general-purpose geometry variables for a single cell BATTERY model.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | IMODEL | ALEN | SLEN | CLEN | ACCLEN | CCCLEN |  |  |
| Type | I | F | F | F | F | F |  |  |
| Default | none | none | none | none | none | none |  |  |


| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | NELEA | NELES | NELEC | NELECCA | NELECCC |  |  |  |
| Type | 1 | 1 | 1 | 1 | 1 |  |  |  |
| Default | none | none | none | none | none |  |  |  |

VARIABLE
IMODEL

ALEN
SLEN

CLEN

ACCLEN
CCCLEN

NELEA
NELES
NELEC
NELECCA

DESCRIPTION
Battery model identifier (see Remark 1)
Length of the anode side electrode
Length of the separator
Length of the cathode side electrode
Length of the negative current collector
Length of the positive current collector
Number of elements in the anode electrode
Number of elements in the separator
Number of elements in the cathode electrode
Number of elements in the anode current collector

## DESCRIPTION

NELECCC Number of elements in the cathode current collector

## Remarks:

1. Identifier. The battery model identifier (IMODEL) should match the IMODEL value specified in the corresponding *BATTERY_ECHEM_CONTROL_SOLVER card. If a different value is given, the value on the *BATTERY_ECHEM_CONTROL_SOLVER card will be the default.

## *BATTERY_ECHEM_CONTROL_SOLVER

Purpose: Set general purpose control variables for a battery electrochemistry simulation.

## Card Summary:

Card 1. This card is required.

| IMODEL | IDIMEN | NCYCLE | IAGING | ITRA | IGAS |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

Card 2. Include NCYCLE instances of this card, one for each cycle.

| CMODE | CTYPE | CEND | TCUT | VCUT | RCURR |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

Card 3.1. Include this card if IAGING $=1$.

| SEIMW | SEIRH0 | SEIBRUG | SEIEPS | SEIC0 | SEITO |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

Card 3.2. Include this card if IAGING $=1$.

| SEIIO | SEIRKA | SEICON | ECCO | ECDFS |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |

Card 4.1. Include this card if ITRA $=1$.

| AFI | EAT | HOFEC | HOFLI | HOFLED |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |

Card 4.2. Include this card if $\operatorname{ITRA}=1$.

| HOFC2H4 | HOFLC | HOFCO2 | HOFO2 |  |  |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

Card 5. Include this card if $\operatorname{IGAS}=1$.

| IC2H4 | IO2 | IC02 | IH2O | AG1 | AG2 | EG1 | EG2 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |

## Data Card Definitions:

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | IMODEL | IDIMEN | NCYCLE | IAGING | ITRA | IGAS |  |  |
| Type | I | 1 | 1 | 1 | 1 | 1 |  |  |
| Default | none | none | 1 | 1 | 0 | 0 |  |  |

## DESCRIPTION

IMODEL

IDIMEN

NCYCLE Number of cycles to run. Default is 1 cycle.
IAGING Aging model flag (see Remark 3):
EQ.0: Off
EQ.1: On

ITRA Thermal runaway model flag (see Remark 3):
EQ.0: Off
EQ.1: On

IGAS Gas generation model flag (see Remark 4):
EQ.0: Off
EQ.1: On

Cycle Card. Include NCYCLE instances of this card, that is, one for each cycle.

| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | CMODE | CTYPE | CEND | TCUT | VCUT | RCURR |  |  |
| Type | 1 | 1 | 1 | $F$ | $F$ | $F$ |  |  |
| Default | 1 | 0 | none | none | none | none |  |  |

VARIABLE
CMODE

CTYPE

CEND

TCUT
VCUT
RCURR Cycle operating current in the case of constant current

Aging Card 1. Include this card if IAGING $=1$.

| Card 3.1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | SEIMW | SEIRHO | SEIBRUG | SEIEPS | SEICO | SEIT0 |  |  |
| Type | F | F | F | F | F | F |  |  |
| Default | 0.0 | 0.0 | 0.0 | 0.0 | 0.0 | 0.0 |  |  |

Aging Card 2. Include this card if IAGING = 1.

| Card 3.2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | SEIIO | SEIRKA | SEICON | ECCO | ECDFS |  |  |  |
| Type | F | F | F | F | F |  |  |  |
| Default | 0.0 | 0.0 | 0.0 | 0.0 | 0.0 |  |  |  |

VARIABLE
SEIMW Molecular weight of the SEI
SEIRHO Density of the SEI

SEIEPS Initial SEI porosity

ECC0

SEIBRUG The Bruggeman constant of the SEI

SEIC0 Initial SEI concentration (units: $\mathrm{mol} / \mathrm{m}^{3}$ )
SEIT0 Initial thickness of the SEI layer (units: m)
SEII0 Exchange current density for the SEI reaction
SEIRKA Reaction rate constant for the SEI reaction (ignored if SEIIO $\neq 0.0$ )
SEICON Ionic conductivity (units: $\mathrm{S} / \mathrm{m}$ )

## DESCRIPTION

Initial concentration of EC (ethylene carbonate). This field is ignored if SEIIO $\neq 0.0$.

## VARIABLE

ECDFS

## DESCRIPTION

Diffusion coefficient of EC

Thermal Runaway Card 1. Include this card if ITRA $=1$.

| Card 4.1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | AFI | EAT | HOFEC | HOFLI | HOFLED |  |  |  |
| Type | F | F | F | F | F |  |  |  |
| Default | 0.0 | 0.0 | 0.0 | 0.0 | 0.0 |  |  |  |

Thermal Runaway Card 2. Include this card if ITRA $=1$.

| Card 4.2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | HOFC2H4 | HOFLC | HOFCO2 | HOFO2 |  |  |  |  |
| Type | F | F | F | F |  |  |  |  |
| Default | 0.0 | 0.0 | 0.0 | 0.0 |  |  |  |  |

## VARIABLE

AFI
EAT
HOFEC
HOFLI
HOFLED

HOFLC
HOFCO2
HOFO2

HOFC2H4 Formation enthalpy of ethylene (units: $\mathrm{kJ} / \mathrm{mol}$ )

## DESCRIPTION

Frequency factor for the reaction
Activation energy for the reaction
Formation enthalpy of the EC (units: $\mathrm{kJ} / \mathrm{mol}$ )
Formation enthalpy of the LI
Formation enthalpy of the SEI layer (units: $\mathrm{kJ} / \mathrm{mol}$ )

Formation enthalpy of $\mathrm{LC}\left(\mathrm{Li}_{2} \mathrm{CO}_{3}\right.$; units: $\left.\mathrm{kJ} / \mathrm{mol}\right)$
Formation enthalpy of $\mathrm{CO}_{2}$ (units: $\mathrm{kJ} / \mathrm{mol}$ )
Formation enthalpy of $\mathrm{O}_{2}$ (units: $\mathrm{kJ} / \mathrm{mol}$ )

Gas Initial Conditions Card. Include this card if IGAS $=1$.

| Card 5 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | IC2H4 | IO2 | IC02 | IH20 | AG1 | AG2 | EG1 | EG2 |
| Type | F | F | F | F | F | F | F | F |
| Default | 0.0 | 0.0 | 0.0 | 0.0 | $3.426 E 26$ | $5.028 \mathrm{E}-6$ | $2.50 e 5$ | 2.51 e 5 |
| Remarks |  |  |  |  | 5 | 5 | 5 | 5 |

VARIABLE
IC2H4

AG1
AG2
EG1
EG2

IO 2 Initial concentration of $\mathrm{O}_{2}$ gas (units: $\mathrm{mol} / \mathrm{m}^{3}$ )
ICO2 Initial concentration of $\mathrm{CO}_{2}$ gas (units: $\mathrm{mol} / \mathrm{m}^{3}$ )
$\mathrm{IH} 2 \mathrm{O} \quad$ Initial concentration of $\mathrm{H}_{2} \mathrm{O}$ gas (units: $\mathrm{mol} / \mathrm{m}^{3}$ )

## DESCRIPTION

Initial concentration of $\mathrm{C}_{2} \mathrm{H}_{2}$ gas (units: $\mathrm{mol} / \mathrm{m}^{3}$ )

Frequency factor for Ethylene oxidation (units: $\mathrm{m} / \mathrm{s}$ )
Frequency factor for Lithium hydration (units: $\mathrm{m} / \mathrm{s}$ )
Activation energy of Ethylene oxidation (units: $\mathrm{J} / \mathrm{mol}$ )
Activation energy of Lithium hydration (units: J/mol)

## Remarks:

1. Multiphysics Battery Models. With IMODEL $=2$, you can model a battery with thermal effects or a battery with both thermal and chemistry effects depending on the value of IGAS. These two battery models include battery aging, thermal runaway, and battery swelling. The battery and chemistry effects model additionally includes a gas generation model for modeling the chemistry effects.
2. Shutdown Key. For coupled battery models (IDIMEN = 101) with IMODEL = 1 or 2 , the shutdown key will work if the minimum ignition energy in the battery system reaches the critical condition like pre-thermal-runaway.
3. Input Requirements for Aging and Thermal Runaway Models. When IAG$\operatorname{ING}=1$, all the variables on Cards 3.1 and 3.2 must be filled. A similar
requirement applies when ITRA $=1$ in that all the variables of Cards 41 and 4.2 must be filled. For more details about these models, please see the Theory manual.
4. Battery Chemistry Files. When IGAS $=1$, *CHEMISTRY_BATTERY must be included in the input deck to provide the file names for the battery chemistry input file, the corresponding thermodynamics data file, and the transport properties file. Please refer to the *CHEMISTRY chapter for further details about the *CHEMISTRY_BATTERY keyword.
5. AG1, AG2, EG1, and EG2. AG1, AG2, EG1, and EG2 are used to compute the rate of reaction based on Arrhenius equation as

$$
r=A_{k} c_{k} e^{-\frac{\mathrm{EG}_{k}}{R T}} .
$$

$A_{k}$ is the frequency factor for the reaction, $c_{k}$ is the concentration, $\mathrm{EG}_{k}$ is the activation energy for the reaction, $R$ is the universal gas constant, and $T$ is the temperature.
*BATTERY_ECHEM_INITIAL
Purpose: Initializes the composite electrodes and electrolyte in every element of the BATTERY simulation mesh.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | DT0 | LICE | PHI1 | LICS | CURRIC | FLUXIC | PHI2 |  |
| Type | F | F | F | F | F | F | F |  |
| Default | none | none | none | none | none | none | none |  |

## VARIABLE

DT0

LICE

PHI1
LICS

CURRIC

FLUXIC
PHI2 Initial condition of the electrode potential

## *BATTERY_ECHEM_MAT_ANODE

Purpose: Set the battery material variables for the anode side electrode.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | PIDA | IOCPA | CAPTA | SOCA | RADA | RATEA | RANODE |  |
| Type | I | I | F | F | F | $F$ | $F$ |  |
| Default | none | none | none | none | none | none | none |  |


| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | RHOEA | RHOFA | RHOCCA | DFSA | CONDA | MWA |  |  |
| Type | F | F | F | F | F | $F$ |  |  |
| Default | none | none | none | none | none | none |  |  |


| Card 3 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | VFEA | VFPA | VFFA | VFGA |  |  |  |  |
| Type | F | F | F | F |  |  |  |  |
| Default | none | none | none | none |  |  |  |  |

## VARIABLE

## DESCRIPTION

PIDA

IOCPA

Part ID

Material type for the OCP in the anode electrode:
EQ.1: Sony carbon (petroleum coke)
EQ.2: MCMB 2510
EQ.3: MCMB 2528

VARIABLE

CAPTA
SOCA

RADA

RATEA
RANODE
RHOEA

RHOFA
RHOCCA
DFSA

CONDA

MWA

VFEA
VFPA
VFFA
VFGA

## DESCRIPTION

## EQ.4: KS6 graphite

Coulombic capacity of anode material (units: $\mathrm{mAh} / \mathrm{g}$ )
Initial lithium stoichiometric coefficient of the anode side active material. For example, $\operatorname{Li}_{x} \mathrm{C}_{6}(0<x<0.7)$.

Radius of spherical particles in the anode side active material (units: m)

Reaction rate constant for the anode electrode
Film resistance for the anode electrode
Density of anode insertion material (electrode particles) (kg/m ${ }^{3}$ )
Density of the inert filler in the anode (units: $\mathrm{kg} / \mathrm{m}^{3}$ )
Density of the current collector in the anode (units: $\mathrm{kg} / \mathrm{m}^{3}$ )
Diffusion coefficient of lithium ions in the anode electrode material (units: $\mathrm{m}^{2} / \mathrm{s}$ )

Effective electronic conductivity of the anode porous electrode (units: S/m)

Molecular weight of the anode electrode (units: $\mathrm{kg} / \mathrm{mol}$ )
Volume fraction of electrolyte in the anode electrode
Volume fraction of the polymer phase in the anode electrode
Volume fraction of the inert filler in the anode electrode
Volume fraction of the gas in the anode electrode

## *BATTERY_ECHEM_MAT_CATHODE

Purpose: Set the battery material variables for the positive electrode.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | PIDC | IOCPC | CAPTC | SOCC | RADC | RATEC | RCATHDE |  |
| Type | I | I | F | F | F | F | F |  |
| Default | none | none | none | none | none | none | none |  |


| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | RHOEC | RHOFC | RHOCCC | DFSC | CONDC | MWC |  |  |
| Type | F | F | F | F | F | F |  |  |
| Default | none | none | none | none | none | none |  |  |


| Card 3 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | VFEC | VFPC | VFFC | VFGC |  |  |  |  |
| Type | F | F | F | F |  |  |  |  |
| Default | none | none | none | none |  |  |  |  |

## VARIABLE

## DESCRIPTION

PIDC
Part ID
IOCPC Material type for the open-circuit potential:
EQ.1: $\mathrm{Mn}_{2} \mathrm{O}_{4}$ (lower plateau) $(1.1<\mathrm{y}<1.99)$
EQ.2: $\mathrm{Mn}_{2} \mathrm{O}_{4}$ (upper plateau) $(0.17<\mathrm{y}<0.99)$
EQ.3: Cobalt dioxide 1, $\mathrm{Li}_{y} \mathrm{CoO}_{2}(0.0<\mathrm{y}<0.99)$

## DESCRIPTION

EQ.4: Cobalt dioxide 2, $\mathrm{Li}_{\mathrm{y}} \mathrm{CoO}_{2}(0.0<\mathrm{y}<0.99)$
EQ.5: $\mathrm{Mn}_{2} \mathrm{O}_{4}$ (literature version) $(0.17<\mathrm{y}<0.99)$
EQ.6: NMC-111
EQ.7: NMC-811

CAPTC Coulombic capacity of the cathode material (units: $\mathrm{mAh} / \mathrm{g}$ )
SOCC Initial lithium stoichiometric coefficient for the cathode side active material. For example, $\mathrm{Li}_{y} \mathrm{WO}_{3}(0<y<0.67)$.

RADC Radius of spherical particle in the cathode side active material. (units: m)

RATEC Reaction rate constant for the cathode electrode
RCATH Film resistance for the cathode electrode
RHOEC Density of the cathode insertion material (electrode particles). (units: $\mathrm{kg} / \mathrm{m}^{3}$ )

RHOFC Density of the cathode side inert filler (units: $\mathrm{kg} / \mathrm{m}^{3}$ )
RHOCCC Density of the cathode side current collector (units: $\mathrm{kg} / \mathrm{m}^{3}$ )
DFSC Diffusion coefficient of lithium ions in the cathode insertion material (units: $\mathrm{m}^{2} / \mathrm{s}$ )

CONDC Effective electronic conductivity of the cathode porous electrode (units: S/m).

MWC Molecular weight of the cathode electrode (units: $\mathrm{kg} / \mathrm{mol}$ )
VFEC Volume fraction of electrolyte in the cathode electrode
VFPC Volume fraction of the polymer phase in the cathode electrode
VFFC Volume fraction of the inert filler in the cathode electrode
VFGC Volume fraction of the gas in the cathode electrode

## *BATTERY_ECHEM_MAT_ELECTROLYTE

Purpose: Set the battery material variables for the electrolyte and separator.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | PIDEL | IOCPE | IELTYPE | RHOEL | RHOP | RHOS |  |  |
| Type | I | I | I | F | F | F |  |  |
| Default | none | none | none | none | none | none |  |  |


| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | VFES | VFPS | VFGS |  |  |  |  |  |
| Type | F | F | F |  |  |  |  |  |
| Default | none | none | none |  |  |  |  |  |

## VARIABLE

PIDEL
IOPCE

IELTYPE

DESCRIPTION
Part ID
Material type for the open-circuit potential:
EQ.1: LiPF6 in EC : DMC (1:1).
EQ.2: LiPF6 in EC : DMC (2:1).
EQ.3: LiPF6 in EC : DMC (1:2).
EQ.4: LiPF6 in PC
EQ.5: LiClO 4 in PC
Here, EC is ethylene carbonate, DMC is dimethyl carbonate, and PC is propylene carbonate.

Type of electrolyte (units: $\mathrm{kg} / \mathrm{m}^{3}$ ):
EQ.O: Liquid electrolyte
EQ.1: Polymer electrolyte

RHOEL Density of the electrolyte (units: $\mathrm{kg} / \mathrm{m}^{3}$ )
RHOP Density of the polymer phase (units: $\mathrm{kg} / \mathrm{m}^{3}$ )
RHOS Density of the separator material (units: $\mathrm{kg} / \mathrm{m}^{3}$ )
VFES Volume fraction of electrolyte in the separator
VFPS Volume fraction of the polymer phase in the separator
VFGS Volume fraction of the gas in the separator

## *BATTERY_ECHEM_THERMAL

Purpose: Set parameters for the thermal treatment in a cell stack.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | TNAME | ITTYPE | IPRT | CP | HCONV | TEMP |  |  |
| Type | A | I | I | F | F | F |  |  |
| Default | none | none | none | none | none | none |  |  |
| Remarks |  |  | 3 | 2 | 2 | 2 |  |  |


| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |
| Variable | FILE |  |  |  |  |  |  |  |
| Type | A |  |  |  |  |  |  |  |

## VARIABLE

TNAME

ITTYPE

IPRT

CP

HCONV

## DESCRIPTION

Thermal material identifier
Flag for how temperature is determined:
EQ.0: Constant temperature mode
EQ.1: Isothermal temperature with time
EQ.2: Thermal coupling with LS-DYNA thermal solver
Data print in ASCII format:
EQ.O: No data print out.
EQ.1: Time versus heat flux print out for thermal solver
Specific heat coefficient of the cell (units: $\mathrm{J} /(\mathrm{kg} \mathrm{K})$ )
Convective heat transfer coefficient with external medium. (units: W/( $\left.\mathrm{m}^{2} \mathrm{~K}\right)$ )

TEMP

Ambient temperature around the cell stack (K)

VARIABLE
FILE
Name of the battery cell output file (ASCII)

## Remarks:

1. Battery Parts for Thermal-Mechanical Coupling. In case of thermal-mechanical coupling, the part number for the battery simulation must be specified, so only this part number is considered in the battery parts.
2. Material Properties for Thermal-Mechanical Coupling. If ITTYPE is 2 , the material properties are set through the thermal material card, including anisotropic conductivities (see *MAT_THERMAL_ORTHOTROPIC). CP, HCONV, and TEMP specified here are ignored.
3. Heat Flux Output. If IPRT = 1, then the heat flux generated by the battery solver is printed out.

## Example:

The following is a partial example for 1D Electrochemisty.


```
*BATTERY_ECHEM_CELL_GEOMETRY
$---------1----------2---------3----------4--------------------------------------
\begin{tabular}{rrrrrr}
\(\$\) & alen & slen & clen & acclen & ccclen \\
2 & \(9.6 e-5\) & \(2.5 e-5\) & \(6.0 e-5\) & \(1.0 e-5\) & \(1.0 e-5\)
\end{tabular}
$ nelea neles nelec
$
*BATTERY_ECHEM_MAT_ANODE
\begin{tabular}{|c|c|c|c|c|c|c|c|}
\hline \$ & pida & i0cpa & capta & soca & rada & ratea & ranode \\
\hline & 2 & 3 & 372.0 & 0.6 & 8.0e-6 & \(3.0 \mathrm{e}-9\) & \(0.35 \mathrm{e}-2\) \\
\hline \$ & rhoea & rhofa & rhocca & dfsa & conda & mwa & \\
\hline & 1800.0 & 1800.0 & 8954.0 & 7.0e-14 & 100.0 & 0.079 & \\
\hline \$ & vfec & vfpc & vffc & vfgc & & & \\
\hline & 0.4 & 0.0 & 0.064 & 0.0 & & & \\
\hline
\end{tabular}
$
*BATTERY_ECHEM_MAT_CATHODE
\begin{tabular}{|c|c|c|c|c|c|c|c|}
\hline \$ & pidc & iocpc & captc & SOCC & radc & ratec & rcathde \\
\hline & 2 & 3 & 274.0 & 0.8 & 5.0e-6 & 3.0e-9 & 0.0 \\
\hline \$ & rhoec & rhofc & rhoccc & dfsc & condc & mwc & \\
\hline & 5010.0 & 1800.0 & 2707.0 & 3.0e-14 & 0.5 & 0.9787 & \\
\hline \$ & vfelc & vfplc & vffic & vfgsx & & & \\
\hline & 0.36 & 0.0 & 0.106 & 0.0 & & & \\
\hline
\end{tabular}
$
*BATTERY_ECHEM_MAT_ELECTROLYTE
$---------1----------2----------3----------4--------------------------------------------
\begin{tabular}{rrrrrr}
\(\$\) & pidel & iocpe & ieltype & rhoel & rhop
\end{tabular}\(\quad\) rhos
$ vfels vfpls
$
*BATTERY ECHEM THERMAL
$--------1----------2----------3----------4----------------------------------------
```



```
heat_\overline{discharg_lco.k}
$
* END
```


## *CESE

The keyword *CESE provides input data for the Conservation Element/Solution Element (CESE) compressible fluid solver:
*CESE_BOUNDARY_AXISYMMETRIC_\{OPTION\}
*CESE_BOUNDARY_BLAST_LOAD\}
*CESE_BOUNDARY_CONJ_HEAT_\{OPTION\}
*CESE_BOUNDARY_CYCLIC_\{OPTION\}
*CESE_BOUNDARY_FSI_\{OPTION\}
*CESE_BOUNDARY_NON_REFLECTIVE_\{OPTION\}
*CESE_BOUNDARY_PRESCRIBED_\{OPTION\}
*CESE_BOUNDARY_REFLECTIVE_\{OPTION\}
*CESE_BOUNDARY_SLIDING_\{OPTION\}
*CESE_BOUNDARY_SOLID_WALL_\{OPTION1\}_\{OPTION2\}
*CESE_CHEMISTRY_D3PLOT
*CESE_CONTROL_LIMITER
*CESE_CONTROL_MESH_MOV
*CESE_CONTROL_SOLVER
*CESE_CONTROL_TIMESTEP
*CESE_DATABASE_ELOUT
*CESE_DATABASE_FLUXAVG
*CESE_DATABASE_FSIDRAG
*CESE_DATABASE_POINTOUT
*CESE_DATABASE_SSETDRAG
*CESE_DEFINE_NONINERTIAL
*CESE_DEFINE_POINT

## *CESE

*CESE_DRAG
*CESE_EOS_CAV_HOMOG_EQUILIB_
*CESE_EOS_IDEAL_GAS
*CESE_EOS_INFLATOR1
*CESE_EOS_INFLATOR2
*CESE_FSI_EXCLUDE
*CESE_INITIAL
*CESE_INITIAL_\{OPTION\}
*CESE_INITIAL_CHEMISTRY
*CESE_INITIAL_CHEMISTRY_ELEMENT
*CESE_INITIAL_CHEMISTRY_PART
*CESE_INITIAL_CHEMISTRY_SET
*CESE_MAT_000
*CESE_MAT_001 (*CESE_MAT_GAS)
*CESE_MAT_002
*CESE_PART
*CESE_SURFACE_MECHSSID_D3PLOT
*CESE_SURFACE_MECHVARS_D3PLOT

Note that when performing a chemistry calculation with the CESE solver, initialization should only be done with the *CESE_INITIAL_CHEMISTRY_... cards, not the *CESE_INITIAL... cards.

## *CESE_BOUNDARY_AXISYMMETRIC_OPTION

Available options are:
MSURF
MSURF_SET
SET

## SEGMENT

Purpose: Define an axisymmetric boundary condition on the axisymmetric axis for the 2D axisymmetric CESE compressible flow solver.

The MSURF and MSURF_SET options are used when the CESE mesh has been created using *MESH cards. The SET and SEGMENT cards are used when *ELEMENT_SOLID cards are used to define the CESE mesh.

## Card Summary:

Card 1a. This card is included if the keyword option is MSURF. Provide as many cards as necessary. This input ends at the next keyword ("*") card.

| MSURFID |  |  |  |  |  |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

Card 1b. This card is included if the keyword option is MSURF_SET. Provide as many cards as necessary. This input ends at the next keyword ("*") card.

| MSURF_S |  |  |  |  |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- |

Card 1c. This card is included if the keyword option is SET. Provide as many cards as necessary. This input ends at the next keyword ("*") card.

| SSID |  |  |  |  |  |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

Card 1d. This card is included if the keyword option is SEGMENT. Provide as many cards as necessary. This input ends at the next keyword ("*") card.

| N1 | N2 | N3 | N4 |  |  |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

## Data Card Definitions:

Surface Part Card. Card 1 format used when the MSURF keyword option is active. Provide as many cards as necessary. This input ends at the next keyword ("*") card.

| Card 1a | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | MSURFID |  |  |  |  |  |  |  |
| Type | 1 |  |  |  |  |  |  |  |
| Default | none |  |  |  |  |  |  |  |

## VARIABLE

MSURFID

## DESCRIPTION

Mesh surface part ID referenced in *MESH_SURFACE_ELEMENT cards.

Surface Part Set Card. Card 1 format used when the MSURF_SET keyword option is active. Provide as many cards as necessary. This input ends at the next keyword ("*") card.

| Card 1b | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | MSURF_S |  |  |  |  |  |  |  |
| Type | 1 |  |  |  |  |  |  |  |
| Default | none |  |  |  |  |  |  |  |

VARIABLE
MSURF_S

## DESCRIPTION

Identifier of a set of mesh surface part IDs created with a *LSO_ID_SET card, where each mesh surface part ID in the set is referenced in *MESH_SURFACE_ELEMENT cards.

Set Card. Card 1 format used when the SET keyword option is active. Provide as many cards as necessary. This input ends at the next keyword ("*") card.

| Card 1c | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | SSID |  |  |  |  |  |  |  |
| Type | 1 |  |  |  |  |  |  |  |
| Default | none |  |  |  |  |  |  |  |

## VARIABLE

SSID Segment set ID

Segment Cards. Card 1 format used when SEGMENT keyword option is active. Include an additional card for each corresponding pair of segments. This input ends at the next keyword ("*") card.

| Card 1d | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | N1 | N2 | N3 | N4 |  |  |  |  |
| Type | I | 1 | 1 | 1 |  |  |  |  |
| Default | none | none | none | none |  |  |  |  |

## VARIABLE

## DESCRIPTION

N1, N2, ... Node IDs defining a segment

## Remarks:

1. Restrictions. This boundary condition can only be used on the axisymmetric axis for the 2D axisymmetric CESE fluid solver.

## *CESE_BOUNDARY_BLAST_LOAD_OPTION

Available options include:
MSURF
MSURF_SET
SET

## SEGMENT

Purpose: For the CESE compressible flow solver, set boundary values for velocity, density, and pressure from a blast wave defined by a *LOAD_BLAST_ENHANCED card. Boundary values are applied at the centroid of elements connected with this boundary. $O P T I O N=S E T$ and $O P T I O N=$ SEGMENT are for user defined meshes whereas OPTION $=$ MSURF and OPTION $=$ MSURF_SET are associated with the automatic volume mesher (See *MESH keywords). In other words, the MSURF and MSURF_SET options are used when the CESE mesh has been created using *MESH cards. The SET and SEGMENT cards are used when *ELEMENT_SOLID cards are used to define the CESE mesh.

## Card Summary:

Card 1a. This card is included if the keyword option is MSURF.

| BID | MSURFID |  |  |  |  |  |  |
| :---: | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

Card 1b. This card is included if the keyword option is MSURF_SET.

| BID | MSURF_S |  |  |  |  |  |
| :---: | :--- | :--- | :--- | :--- | :--- | :--- |

Card 1c. This card is included if the keyword option is SET.

| BID | SSID |  |  |  |  |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

Card 1d. This card is included if the keyword option is SEGMENT.

| BID | N1 | N2 | N3 | N4 |  |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

## Data Card Definitions:

Surface Part Card. Card 1 format used when the MSURF keyword option is active.

| Card 1a | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | BID | MSURFID |  |  |  |  |  |  |
| Type | 1 | 1 |  |  |  |  |  |  |
| Default | none | none |  |  |  |  |  |  |

VARIABLE BID

MSURFID

## DESCRIPTION

Blast source ID (see *LOAD_BLAST_ENHANCED)
A mesh surface part ID referenced in *MESH_SURFACE_ELEMENT cards

Surface Part Set Card. Card 1 format used when the MSURF_SET keyword option is active.

| Card 1b | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | BID | MSURF_S |  |  |  |  |  |  |
| Type | 1 | 1 |  |  |  |  |  |  |
| Default | none | none |  |  |  |  |  |  |

## VARIABLE

BID Blast source ID (see *LOAD_BLAST_ENHANCED)
MSURF_S Identifier of a set of mesh surface part IDs created with a *LSO_ID_SET card, where each mesh surface part ID in the set is referenced in *MESH_SURFACE_ELEMENT cards.

Set Card. Card 1 format used when the SET keyword option is active.

| Card 1c | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | BID | SSID |  |  |  |  |  |  |
| Type | 1 | 1 |  |  |  |  |  |  |
| Default | none | none |  |  |  |  |  |  |

VARIABLE
BID
SSID

## DESCRIPTION

Blast source ID (see *LOAD_BLAST_ENHANCED)
Segment set ID

Segment Card. Card 1 for SEGMENT keyword option is active.

| Card 1d | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | BID | N1 | N2 | N3 | N4 |  |  |  |
| Type | I | I | 1 | 1 | I |  |  |  |
| Default | none | none | none | none | none |  |  |  |

VARIABLE
BID
N1, N2, ...

## DESCRIPTION

Blast source ID (see *LOAD_BLAST_ENHANCED)
Node ID's defining a segment

## *CESE_BOUNDARY_CONJ_HEAT_OPTION

Available options are:
MSURF
MSURF_SET
SET

## SEGMENT

Purpose: Define a conjugate heat transfer interface condition for CESE compressible flows. This condition identifies those boundary faces of the CESE mesh that are in contact with non-moving structural parts and through which heat flows. This is only possible when the structural thermal solver is also being used in the structural parts.

The MSURF and MSURF_SET options are used when the CESE mesh has been created using *MESH cards. The SET and SEGMENT cards are used when *ELEMENT_SOLID cards are used to define the CESE mesh.

## Card Summary:

Card 1a. This card is included when the keyword option is MSURF. Include as many cards as necessary. This input ends at the next keyword ("*") card.

| MSURFID |  |  |  |  |  |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

Card 1b. This card is included when the keyword option is MSURF_SET. Include as many cards as necessary. This input ends at the next keyword ("*") card.

| MSURF_S |  |  |  |  |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- |

Card 1c. This card is included when the keyword option is SET. Include as many cards as necessary. This input ends at the next keyword ("*") card.

| SSID |  |  |  |  |  |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

Card 1d. This card is included when the keyword option is SEGMENT. Include as many cards as necessary. This input ends at the next keyword ("*") card.

| N1 | N2 | N3 | N4 |  |  |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

## Data Card Definitions:

Surface Part Card. Card 1 used when the MSURF keyword option is active. Include as many cards as necessary. This input ends at the next keyword ("*") card.

| Card 1a | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | MSURFID |  |  |  |  |  |  |  |
| Type | 1 |  |  |  |  |  |  |  |
| Default | none |  |  |  |  |  |  |  |

## VARIABLE

MSURFID

## DESCRIPTION

Mesh surface part ID referenced in *MESH_SURFACE_ELEMENT cards.

Surface Part Set Card. Card 1 used when the MSURF_SET keyword option is active. Include as many cards as necessary. This input ends at the next keyword ("*") card.

| Card 1b | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | MSURF_S |  |  |  |  |  |  |  |
| Type | I |  |  |  |  |  |  |  |
| Default | none |  |  |  |  |  |  |  |

VARIABLE
MSURF_S

## DESCRIPTION

Identifier of a set of mesh surface part IDs created with an *LSO_ID_SET card, where each mesh surface part ID in the set is referenced in *MESH_SURFACE_ELEMENT cards.

Set Card. Card 1 used when the SET keyword option is active. Include as many cards as necessary. This input ends at the next keyword ("*") card.

| Card 1c | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | SSID |  |  |  |  |  |  |  |
| Type | 1 |  |  |  |  |  |  |  |
| Default | none |  |  |  |  |  |  |  |

## VARIABLE

SSID

DESCRIPTION
Segment set ID

Segment Cards. Card 1 used when SEGMENT keyword option is active. Include an additional card for each corresponding pair of segments. This input ends at the next keyword ("*") card.

| Card 1d | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | N1 | N2 | N3 | N4 |  |  |  |  |
| Type | I | 1 | 1 | 1 |  |  |  |  |
| Default | none | none | none | none |  |  |  |  |

## VARIABLE

## DESCRIPTION

N1, N2, ...
Node IDs defining a segment

## Remarks:

1. Restrictions. This boundary condition should only be imposed on a CESE mesh boundary that is in contact with non-moving structural parts. An Eulerian CESE solver is required as is use of the structural thermal solver.

## *CESE_BOUNDARY_CYCLIC_OPTION

Available options are:
MSURF
MSURF_SET
SET

## SEGMENT

Purpose: Define a cyclic (periodic) boundary condition for CESE compressible flows. This cyclic boundary condition can be used on periodic boundary surfaces.

The MSURF and MSURF_SET options are used when the CESE mesh has been created using ${ }^{*}$ MESH cards. The SET and SEGMENT cards are used when *ELEMENT_SOLID cards are used to define the CESE mesh.

## Card Summary:

Include as many sets of the following cards as needed. This input ends at the next keyword ("*") card.

Card 1a. This card is included when the keyword option is set to MSURF.

| MSURFID1 | MSURFID2 | CYCTYP |  |  |  |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

Card 1 lb . This card is included when the keyword option is set to MSURF_SET.

| MSRF_S1 | MSRF_S2 | CYCTYP |  |  |  |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

Card 1c. This card is included when the keyword option is set to SET.

| SSID1 | SSID2 | CYCTYP |  |  |  |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

Card 1d. This card is included when the keyword option is set to SEGMENT.

| ND1 | ND2 | ND3 | ND4 | NP1 | NP2 | NP3 | NP4 |
| :---: | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

Card 2a. This card is included for the MSURF, MSURF_SET, and SET options when CYCTYP $=1$.

| AXISX1 | AXISY1 | AXISZ1 | DIRX | DIRY | DIRZ | ROTANG |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

Card 2b. This card is included for the MSURF, MSURF_SET, and SET options when CYCTYP $=2$.

| TRANSX | TRANSY | TRANSZ |  |  |  |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

## Data Card Definitions:

Surface Part Card. Card 1 format when the MSURF keyword option is active

| Card 1a | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | MSURFID1 | MSURFID2 | CYCTYP |  |  |  |  |  |
| Type | 1 | 1 | 1 |  |  |  |  |  |
| Default | none | none | 0 |  |  |  |  |  |
| Remarks | 2 | 2 | 1 |  |  |  |  |  |

## VARIABLE

MSURFID1, MSURFID2

CYCTYP

## DESCRIPTION

Mesh surface part IDs referenced in *MESH_SURFACE_ELEMENT cards.

Relationship between the two cyclic boundary condition surfaces:

EQ.0: None assumed (default)
EQ.1: The first surface is rotated about an axis to match the second surface.

EQ.2: The faces of the first surface are translated in a given direction to obtain the corresponding faces on the second surface.

Surface Part Set Card. Card 1 format when the MSURF_SET keyword option is active

| Card 1b | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | MSRF_S1 | MSRF_S2 | CYCTYP |  |  |  |  |  |
| Type | 1 | 1 | 1 |  |  |  |  |  |
| Default | none | none | 0 |  |  |  |  |  |
| Remarks | 3 | 3 | 1 |  |  |  |  |  |

## VARIABLE

MSRF_S1,
MSRF_S2

## DESCRIPTION

Identifiers of two sets of mesh surface part IDs, each created with a *LSO_ID_SET card, where each mesh surface part ID in each set is referenced in *MESH_SURFACE_ELEMENT cards.

Relationship between the two cyclic boundary condition surfaces:

EQ.0: None assumed (default)
EQ.1: The first surface is rotated about an axis to match the second surface.

EQ.2: The faces of the first surface are translated in a given direction to obtain the corresponding faces on the second surface.

Set Card. Card 1 format when the SET keyword option is active

| Card 1c | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | SSID1 | SSID2 | CYCTYP |  |  |  |  |  |
| Type | 1 | 1 | 1 |  |  |  |  |  |
| Default | none | none | 0 |  |  |  |  |  |
| Remarks | 4 | 4 | 1 |  |  |  |  |  |

VARIABLE
SSID1, SSID2
CYCTYP

## DESCRIPTION

A pair of segment set IDs.
Relationship between the two cyclic boundary condition surfaces:

EQ.0: None assumed (default)
EQ.1: The first surface is rotated about an axis to match the second surface.

EQ.2: The faces of the first surface are translated in a given direction to obtain the corresponding faces on the second surface.

Segment Card. Card 1 format when SEGMENT keyword option is active. Include an additional card for each corresponding pair of segments. This input ends at the next keyword ("*") card.

| Card 1d | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | ND1 | ND2 | ND3 | ND4 | NP1 | NP2 | NP3 | NP4 |
| Type | 1 | 1 | 1 | 1 | 1 | 1 | 1 | 1 |
| Default | none | none | none | none | none | none | none | none |

## VARIABLE

NDi, NPi

## DESCRIPTION

Node IDs defining a pair of segments: ND1, ND2, ND3, ND4 define the first segment, while NP1, NP2, NP3, NP4 define the second segment. This pair of segments must match either through a geometric translation or rotation.

Rotation Case Card. Additional card for the MSURF, MSURF_SET, and SET options when CYCTYP = 1 .

| Card 2a | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | AXISX1 | AXISY1 | AXISZ1 | DIRX | DIRY | DIRZ | ROTANG |  |
| Type | $F$ | $F$ | $F$ | $F$ | $F$ | $F$ | $F$ |  |
| Default | 0.0 | 0.0 | 0.0 | none | none | none | none |  |

## VARIABLE

AXIS[Z,Y,Z]1
DIR[X,Y,Z]

ROTANG The angle of rotation (in degrees) that transforms the centroid of each face on the first surface to the centroid of the corresponding face on the second surface

Translation Case Card. Additional card for the MSURF, MSURF_SET, and SET options when CYCTYP $=2$.

| Card 2b | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | TRANSX | TRANSY | TRANSZ |  |  |  |  |  |
| Type | F | F | F |  |  |  |  |  |
| Default | none | none | none |  |  |  |  |  |

## VARIABLE

TRANS[X,Y,Z]

## DESCRIPTION

The translation direction that enables the identification of the segment in the second surface that matches a segment in the first surface

## Remarks:

1. CYCTYP. For the MSURF, MSURF_SET, or SET options with CYCTYP $=0$, the code examines the geometry of two faces of the two surfaces in order to determine if the surfaces are approximately parallel (CYCTYP $=2$ ) or related through a rotation $(C Y C T Y P=1)$. The geometric parameters required are then computed.
2. MSURF. For the MSURF option, the number of mesh surface elements must be the same in each mesh surface part. The mesh surface elements in each mesh surface part are then internally ordered to match pairwise between the two mesh surface parts.
3. MSURF_SET. For the MSURF_SET option, the number of mesh surface elements must be the same in each mesh surface part set. The mesh surface elements in each mesh surface part set are then internally ordered to match pairwise between the two mesh surface part sets.
4. SET. For the SET option, the same number of segments must be in each set. The segments in each set are then internally ordered to match pairwise between the two sets.

## *CESE_BOUNDARY_FSI_OPTION

Available options are:
MSURF
MSURF_SET
SET

## SEGMENT

Purpose: Define an FSI boundary condition for the moving mesh CESE compressible flow solver. This card must not be combined with the immersed-boundary method CESE solver; doing so will result in an error termination condition.

This boundary condition must be applied on a surface of the CESE computational domain that is co-located with surfaces of the outside boundary of the structural mechanics' mesh. The nodes of the two meshes will generally not be shared.

The MSURF and MSURF_SET options are used when the CESE mesh has been created using *MESH cards. The SET and SEGMENT cards are used when *ELEMENT_SOLID cards are used to define the CESE mesh.

## Card Summary:

Card 1a. This card is included if the keyword option is MSURF. Provide as many cards as necessary. This input ends at the next keyword ("*") card.

| MSURFID |  |  |  |  |  |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

Card 1b. This card is included if the keyword option is MSURF_SET. Provide as many cards as necessary. This input ends at the next keyword ("*") card.

| MSURF_S |  |  |  |  |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- |

Card 1c. This card is included if the keyword option is SET. Provide as many cards as necessary. This input ends at the next keyword ("*") card.

| SSID |  |  |  |  |  |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

Card 1d. This card is included if the keyword option is SEGMENT. Provide as many cards as necessary. This input ends at the next keyword ("*") card.

| N1 | N2 | N3 | N4 |  |  |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

## Data Card Definitions:

Surface Part Card. Card 1 format used when the MSURF keyword option is active. Provide as many cards as necessary. This input ends at the next keyword ("*") card.

| Card 1a | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | MSURFID |  |  |  |  |  |  |  |
| Type | 1 |  |  |  |  |  |  |  |
| Default | none |  |  |  |  |  |  |  |

## VARIABLE

MSURFID

## DESCRIPTION

Mesh surface part ID referenced in *MESH_SURFACE_ELEMENT cards

Surface Part Set Card. Card 1 format used when the MSURF_SET keyword option is active. Provide as many cards as necessary. This input ends at the next keyword ("*") card.

| Card 1b | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | MSURF_S |  |  |  |  |  |  |  |
| Type | 1 |  |  |  |  |  |  |  |
| Default | none |  |  |  |  |  |  |  |

VARIABLE
MSURF_S

## DESCRIPTION

Identifier of a set of mesh surface part IDs created with a *LSO_ID_SET card, where each mesh surface part ID in the set is referenced in *MESH_SURFACE_ELEMENT cards.

Set Card. Card 1 format used when the SET keyword option is active. Provide as many cards as necessary. This input ends at the next keyword ("*") card.

| Card 1c | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | SSID |  |  |  |  |  |  |  |
| Type | 1 |  |  |  |  |  |  |  |
| Default | none |  |  |  |  |  |  |  |

## VARIABLE

SSID

DESCRIPTION
Segment set ID

Segment Cards. Card 1 format used when SEGMENT keyword option is active. Include an additional card for each corresponding pair of segments. This input ends at the next keyword ("*") card.

| Card 1d | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | N1 | N2 | N3 | N4 |  |  |  |  |
| Type | I | 1 | 1 | 1 |  |  |  |  |
| Default | none | none | none | none |  |  |  |  |

## VARIABLE

## DESCRIPTION

N1, ...
Node IDs defining a segment

## Remarks:

1. Conjugate Heat Transfer Problems. This boundary condition card is also needed for conjugate heat transfer problems with the moving mesh CESE solver.

## *CESE_BOUNDARY_NON_REFLECTIVE_OPTION

Available options are:
MSURF
MSURF_SET
SET

## SEGMENT

Purpose: Define a passive boundary condition for CESE compressible flows. This nonreflective boundary condition (NBC) provides an artificial computational boundary for an open boundary that is passive.

Use the MSURF and MSURF_SET options when you use the *MESH cards to generate the CESE mesh. Use the SET and SEGMENT cards when *ELEMENT_SOLID cards define the CESE mesh.

## Card Summary:

Card 1a. Include this card for the MSURF keyword option. Include as many cards as necessary. This input ends at the next keyword ("*") card.

| MSURFID |  |  |  |  |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

Card 1b. Include this card for the MSURF_SET keyword option. Include as many cards as necessary. This input ends at the next keyword ("*") card.

| MSURF_S |  |  |  |  |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- |

Card 1c. Include this card for the SET keyword option. Include as many cards as necessary. This input ends at the next keyword ("*") card.

| SSID |  |  |  |  |  |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

Card 1d. Include this card for the SEGMENT keyword option. Include an additional card for each corresponding pair of segments. This input ends at the next keyword ("*") card.

| N1 | N2 | N3 | N4 |  |  |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

## Data Card Definitions:

Surface Part Card. Card 1 used when the MSURF keyword option is active. Include as many cards as necessary. This input ends at the next keyword ("*") card.

| Card 1a | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | MSURFID |  |  |  |  |  |  |  |
| Type | 1 |  |  |  |  |  |  |  |
| Default | none |  |  |  |  |  |  |  |

## VARIABLE

MSURFID

## DESCRIPTION

Mesh surface part ID referenced by *MESH_SURFACE_ELEMENT cards

Surface Part Set Card. Card 1 used when the MSURF_SET keyword option is active. Include as many cards as necessary. This input ends at the next keyword ("*") card.

| Card 1b | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | MSURF_S |  |  |  |  |  |  |  |
| Type | 1 |  |  |  |  |  |  |  |
| Default | none |  |  |  |  |  |  |  |

VARIABLE DESCRIPTION

MSURF_S ID of a set of mesh surface parts created with an *LSO_ID_SET card, where each mesh surface part ID in the set is referenced by *MESH_SURFACE_ELEMENT cards.

Set Card. Card 1 used when the SET keyword option is active. Include as many cards as necessary. This input ends at the next keyword ("**) card.

| Card 1c | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | SSID |  |  |  |  |  |  |  |
| Type | I |  |  |  |  |  |  |  |
| Default | none |  |  |  |  |  |  |  |

## VARIABLE

SSID

## DESCRIPTION

Segment set ID

Segment Cards. Card 1 used when the SEGMENT keyword option is active. Include an additional card for each corresponding pair of segments. This input ends at the next keyword ("*") card.

| Card 1d | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | N1 | N2 | N3 | N4 |  |  |  |  |
| Type | I | 1 | 1 | 1 |  |  |  |  |
| Default | none | none | none | none |  |  |  |  |

## VARIABLE

## DESCRIPTION

N1, N2, ... Node IDs defining a segment

## Remarks:

1. Normal application of this boundary condition. We recommend imposing this boundary condition on an open surface far from the main disturbed flow (the further away, the better), meaning an almost uniform flow on the boundary surface.
2. Default boundary condition for CESE. LS-DYNA automatically assigns the non-reflective boundary condition to any boundary segment not already assigned a boundary condition by any of the *CESE_BOUNDARY_... cards.

## *CESE_BOUNDARY_PRESCRIBED_OPTION

Available options include:
MSURF
MSURF_SET
SET

## SEGMENT

Purpose: For the CESE compressible flow solver, set boundary values for velocity, density, pressure, and temperature. Boundary values are applied at the centroid of elements connected with this boundary. Use keyword options SET or SEGMENT for user-defined meshes, meaning mesh specified with *ELEMENT_SOLID card. Use MSURF or MSURF_SET for meshes created with the automatic volume mesher, meaning meshes generated with the *MESH cards.

## Card Summary:

A set of data cards for this keyword consists of 3 of the following cards:

1. Card 1 specifies the object to which the boundary condition is applied. Its format depends on the keyword option.
2. Card 2 reads in load curve IDs.
3. Card 3 reads in scale factors.

For each boundary condition to be specified include one set of cards. This input ends at the next keyword ("*") card.
Card 1a. Include this card for the MSURF keyword option.

| MSURFID | IDCOMP |  |  |  |  |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

Card 1b. Include this card for the MSURF_SET keyword option.

| MSURF_S | IDCOMP |  |  |  |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- |

Card 1c. Include this card for the SET keyword option.

| SSID | IDCOMP |  |  |  |  |  |  |
| :---: | :---: | :--- | :--- | :--- | :--- | :--- | :--- |

Card 1d. Include this card for the SEGMENT keyword option.

| N1 | N1 | N3 | N4 | IDCOMP |  |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

Card 2. This card is required.

| LC_U | LC_V | LC_W | LC_RHO | LC_P | LC_T |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

Card 3. This card is required.

| SF_U | SF_V | SF_W | SF_RHO | SF_P | SF_T |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

## Data Card Definitions:

Surface Part Card. Card 1 format for the MSURF keyword option

| Card 1a | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | MSURFID | IDCOMP |  |  |  |  |  |  |
| Type | 1 | 1 |  |  |  |  |  |  |
| Default | none | none |  |  |  |  |  |  |

VARIABLE
MSURFID

IDCOMP

## DESCRIPTION

Mesh surface part ID referenced in *MESH_SURFACE_ELEMENT cards

For inflow boundaries in problems involving chemical reacting flows, this ID references a *CHEMISTRY_COMPOSITION card that gives the chemical mixture of the fluid entering the domain

Surface Part Set Card. Card 1 format for the MSURF_SET keyword option

| Card 1b | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | MSURF_S | IDCOMP |  |  |  |  |  |  |
| Type | I | 1 |  |  |  |  |  |  |
| Default | none | none |  |  |  |  |  |  |

## VARIABLE

MSURF_S

IDCOMP

## DESCRIPTION

ID of a set of mesh surface parts created with a *LSO_ID_SET card, where each mesh surface part ID in the set is referenced in *MESH_SURFACE_ELEMENT cards.

For inflow boundaries in problems involving chemical reacting flows, this ID references a *CHEMISTRY_COMPOSITION card that gives the chemical mixture of the fluid entering the domain

Set Card. Card 1 format for the SET keyword option

| Card 1c | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | SSID | IDCOMP |  |  |  |  |  |  |
| Type | 1 | 1 |  |  |  |  |  |  |
| Default | none | none |  |  |  |  |  |  |

## VARIABLE

SSID
IDCOMP

## DESCRIPTION

Segment set ID
For inflow boundaries in problems involving chemical reacting flows, this ID references a *CHEMISTRY_COMPOSITION card that gives the chemical mixture of the fluid entering the domain

Segment Card. Card 1 for the SEGMENT keyword option

| Card 1d | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | N1 | N1 | N3 | N4 | IDCOMP |  |  |  |
| Type | I | I | 1 | 1 | 1 |  |  |  |
| Default | none | none | none | none | none |  |  |  |

## VARIABLE

## DESCRIPTION

N1, N2, .. Node IDs defining a segment

VARIABLE
IDCOMP

## DESCRIPTION

For inflow boundaries in problems involving chemical reacting flows, this ID references a *CHEMISTRY_COMPOSITION card that gives the chemical mixture of the fluid entering the domain

Load Curve Card.

| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | LC_U | LC_V | LC_W | LC_RHO | LC_P | LC_T |  |  |
| Type | I | I | 1 | 1 | I | I |  |  |
| Remarks | 1 | 1 | 1 | 1 | 1 | 1 |  |  |

## VARIABLE

LC_U

LC_V Load curve ID to describe the $y$-component of the velocity as a function of time

GT.O: Load curve ID
EQ.O: Constant value of SF_V
EQ.-1: Solver computes the boundary value for $y$-component of velocity

LC_W Load curve ID to describe the $z$-component of the velocity as a function of time

GT.0: Load curve ID
EQ.O: Constant value of SF_W
EQ.-1: Solver computes the boundary value for $x$-component of velocity

## VARIABLE

LC_RHO

LC_P

LC_T

## DESCRIPTION

Load curve ID to describe the density as a function of time
GT.0: Load curve ID
EQ.0: Constant value of SF_RHO
EQ.-1: Solver computes the boundary value for density
Load curve ID to describe the pressure as a function of time
GT.0: Load curve ID
EQ.O: Constant value of SF_P
EQ.-1: Solver computes the boundary value for pressure
Load curve ID to describe the temperature as a function of time
GT.0: Load curve ID
EQ.0: Constant value of SF_T
EQ.-1: Solver computes the boundary value for temperature

Scale Factor Card.

| Card 3 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | SF_U | SF_V | SF_W | SF_RH0 | SF_P | SF_T |  |  |
| Type | F | F | F | F | F | F |  |  |
| Default | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 |  |  |

VARIABLE

## DESCRIPTION

## SF_U

SF_V Scale factor for LC_V
SF_W Scale factor for LC_W
SF_RHO Scale factor for LC_RHO
SF_P Scale factor for LC_P
SF_T Scale factor for LC_T

## Remarks:

1. Well-posed boundaries. On each centroid or set of centroids, the variables ( $v_{x}$, $\left.v_{y}, v_{z}, \rho, P, T\right)$ with provided values must be consistent and make the model wellposed (that is, be such that the solution of the model exists, is unique, and is physical).

## *CESE_BOUNDARY_PRESCRIBED_VN_OPTION

Available options include:
MSURF
MSURF_SET
SET

## SEGMENT

Purpose: For the CESE compressible flow solver, set boundary values for velocity, density, pressure, and temperature. Boundary values are applied at the centroid of elements connected with this boundary. Use keyword options SET or SEGMENT for user-defined meshes, that is, meshes specified with *ELEMENT_SOLID cards. Use MSURF or MSURF_SET for meshes generated with the automatic volume mesher, meaning created with *MESH cards.

## Card Summary:

A set of data cards for this keyword consists of 3 of the following cards:

1. Card 1 specifies the object to which the boundary condition is applied. Its format depends on the keyword option.
2. Card 2 reads in load curve IDs.
3. Card 3 reads in scale factors.

For each boundary condition, include one set of cards. This input ends at the next keyword ("*") card.
Card 1a. Include this card for the MSURF keyword option.

| MSURFID | IDCOMP |  |  |  |  |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

Card 1b. Include this card for the MSURF_SET keyword option.

| MSURF_S | IDCOMP |  |  |  |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- |

Card 1c. Include this card for the SET keyword option.

| SSID | IDCOMP |  |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :--- | :--- | :--- | :--- |

Card 1d. Include this card for the SEGMENT keyword option.

| N1 | N1 | N3 | N4 | IDCOMP |  |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

Card 2. This card is required.

| LC_VN |  |  | LC_RHO | LC_P | LC_T |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

Card 3. This card is required.

| SF_VN |  |  | SF_RHO | SF_P | SF_T |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

## Data Card Definitions:

Surface Part Card. Card 1 format for the MSURF keyword option

| Card 1a | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | MSURFID | IDCOMP |  |  |  |  |  |  |
| Type | 1 | 1 |  |  |  |  |  |  |
| Default | none | none |  |  |  |  |  |  |

VARIABLE
MSURFID

IDCOMP

## DESCRIPTION

A mesh surface part ID referenced in *MESH_SURFACE_ELEMENT cards

For inflow boundaries in problems involving chemical reacting flows, this ID references a *CHEMISTRY_COMPOSITION card that gives the chemical mixture of the fluid entering the domain.

Surface Part Set Card. Card 1 format for the MSURF_SET keyword option

| Card 1b | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | MSURF_S | IDCOMP |  |  |  |  |  |  |
| Type | I | 1 |  |  |  |  |  |  |
| Default | none | none |  |  |  |  |  |  |

## VARIABLE

MSURF_S

IDCOMP

## DESCRIPTION

ID of a set of mesh surface parts created with a *LSO_ID_SET card, where each mesh surface part ID in the set is referenced in *MESH_SURFACE_ELEMENT cards.

For inflow boundaries in problems involving chemical reacting flows, this ID references a *CHEMISTRY_COMPOSITION card that gives the chemical mixture of the fluid entering the domain.

Set Card. Card 1 format for the SET keyword option

| Card 1c | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | SSID | IDCOMP |  |  |  |  |  |  |
| Type | 1 | 1 |  |  |  |  |  |  |
| Default | none | none |  |  |  |  |  |  |

## VARIABLE

SSID
IDCOMP

## DESCRIPTION

Segment set ID
For inflow boundaries in problems involving chemical reacting flows, this ID references a *CHEMISTRY_COMPOSITION card that gives the chemical mixture of the fluid entering the domain.

Segment Card. Card 1 for the SEGMENT keyword option

| Card 1d | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | N1 | N1 | N3 | N4 | IDCOMP |  |  |  |
| Type | I | I | 1 | 1 | 1 |  |  |  |
| Default | none | none | none | none | none |  |  |  |

## VARIABLE

## DESCRIPTION

N1, N2, ... Node IDs defining a segment

VARIABLE
IDCOMP

DESCRIPTION
For inflow boundaries in problems involving chemical reacting flows, this ID references a *CHEMISTRY_COMPOSITION card that gives the chemical mixture of the fluid entering the domain.

Load Curve Card.

| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | LC_VN |  |  | LC_RHO | LC_P | LC_T |  |  |
| Type | I |  |  | 1 | I | I |  |  |
| Remarks | 1 |  |  | 1 | 1 | 1 |  |  |

## VARIABLE

LC_VN

LC_RHO Load curve ID to describe the density as a function of time
GT.0: Load curve ID
EQ.O: Constant value of SF_RHO
EQ.-1: Solver determines the boundary value for the density
LC_P
Load curve ID to describe the pressure as a function of time
GT.0: Load curve ID
EQ.0: Constant value of SF_P
EQ.-1: Solver determines the boundary value for the pressure
LC_T Load curve ID to describe the temperature as a function of time
GT.0: Load curve ID
EQ.0: Constant value of SF_T

## DESCRIPTION

EQ.-1: Solver determines the boundary value for the temperature

## Scale Factor Card.

| Card 3 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | SF_VN |  |  | SF_RHO | SF_P $_{2}$ | SF_T $^{\prime}$ |  |  |
| Type | F |  |  | F | F | F |  |  |
| Default | 1.0 |  |  | 1.0 | 1.0 | 1.0 |  |  |

## VARIABLE

SF_VN
SF_RHO Scale factor for LC_RHO
SF_P
SF_T

## DESCRIPTION

Scale factor for LC_VN

Scale factor for LC_P
Scale factor for LC_T

## Remarks:

1. Well-posed boundary conditions. On each centroid or set of centroids, the variables ( $V_{N}, \rho, P, T$ ) with provided values must be consistent and make the model well-posed (that is, be such that the solution of the model exists, is unique and physical).

## *CESE_BOUNDARY_REFLECTIVE_OPTION

Available options are:
MSURF
MSURF_SET
SET

## SEGMENT

Purpose: Define a reflective boundary condition for the CESE compressible flow solver. You can apply this boundary condition on a symmetrical surface or a solid wall of the computational domain. Use the MSURF or MSURF_SET keyword options when generating the CESE mesh with the *MESH cards. Use the SET or SEGMENT keyword options cards when *ELEMENT_SOLID cards specify CESE mesh.

## Card Summary:

Card 1a. Include this card for the MSURF keyword option. Provide as many cards as necessary. This input ends at the next keyword ("*") card.


Card 1b. Include this card for the MSURF_SET keyword option. Provide as many cards as necessary. This input ends at the next keyword ("**) card.

| MSURF_S |  |  |  |  |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- |

Card 1c. Include this card for the SET keyword option. Provide as many cards as necessary. This input ends at the next keyword ("*") card.

| SSID |  |  |  |  |  |  |  |
| :---: | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

Card 1d. Include this card for the SEGMENT keyword option. Provide as many cards as necessary. This input ends at the next keyword ("*") card.

| N1 | N2 | N3 | N4 |  |  |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

## Data Card Definitions:

Surface Part Card. Card 1 format for the MSURF keyword option. Provide as many cards as necessary. This input ends at the next keyword ("*") card.

| Card 1a | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | MSURFID |  |  |  |  |  |  |  |
| Type | 1 |  |  |  |  |  |  |  |
| Default | none |  |  |  |  |  |  |  |

## VARIABLE

MSURFID

## DESCRIPTION

Mesh surface part ID referenced in *MESH_SURFACE_ELEMENT cards.

Surface Part Set Card. Card 1 format for the MSURF_SET keyword option. Provide as many cards as necessary. This input ends at the next keyword ("*") card.

| Card 1b | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | MSURF_S |  |  |  |  |  |  |  |
| Type | 1 |  |  |  |  |  |  |  |
| Default | none |  |  |  |  |  |  |  |

VARIABLE
MSURF_S

## DESCRIPTION

ID of a set of mesh surface parts created with a *LSO_ID_SET card, where each mesh surface part ID in the set is referenced in *MESH_SURFACE_ELEMENT cards.

Set Card. Card 1 format for the SET keyword option. Provide as many cards as necessary. This input ends at the next keyword ("*") card.

| Card 1c | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | SSID |  |  |  |  |  |  |  |
| Type | 1 |  |  |  |  |  |  |  |
| Default | none |  |  |  |  |  |  |  |

## VARIABLE

SSID

DESCRIPTION
Segment set ID

Segment Cards. Card 1 format for the SEGMENT keyword option. Include an additional card for each corresponding pair of segments. This input ends at the next keyword ("*") card.

| Card 1d | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | N1 | N2 | N3 | N4 |  |  |  |  |
| Type | I | 1 | 1 | 1 |  |  |  |  |
| Default | none | none | none | none |  |  |  |  |

## VARIABLE

## DESCRIPTION

N1, N2, ...
Node IDs defining a segment

## Remarks:

1. Effect of the boundary condition. This boundary condition has the same effect as a solid-wall boundary condition for inviscid flows.

## *CESE_BOUNDARY_SLIDING_OPTION

Available options are:
MSURF
MSURF_SET
SET

## SEGMENT

Purpose: Allows the nodes of a fluid surface to translate in the main direction of the mesh movement. This feature is useful in piston-type applications.

Use the MSURF or MSURF_SET keyword options when generating the CESE mesh with *MESH cards. Use the SET and SEGMENT keywords when *ELEMENT_SOLID cards specify the CESE mesh.

## Card Summary:

Card 1a. Include this card for the MSURF keyword option. Provide as many cards as necessary. This input ends at the next keyword ("*") card.

| MSURFID |  |  |  |  |  |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

Card 1b. Include this card for the MSURF_SET keyword option. Provide as many cards as necessary. This input ends at the next keyword ("*") card.

| MSURF_S |  |  |  |  |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- |

Card 1c. Include this card for the SET keyword option. Provide as many cards as necessary. This input ends at the next keyword ("*") card.

| SSID |  |  |  |  |  |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

Card 1d. Include this card for the SEGMENT keyword option. Provide as many cards as necessary. This input ends at the next keyword ("*") card.

| N1 | N2 | N3 | N4 |  |  |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

## Data Card Definitions:

Surface Part Card. Card 1 format for the MSURF keyword option. Provide as many cards as necessary. This input ends at the next keyword ("*") card.

| Card 1a | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | MSURFID |  |  |  |  |  |  |  |
| Type | 1 |  |  |  |  |  |  |  |
| Default | none |  |  |  |  |  |  |  |

## VARIABLE

MSURFID

## DESCRIPTION

Mesh surface part ID referenced in *MESH_SURFACE_ELEMENT cards.

Surface Part Set Card. Card 1 format used when the MSURF_SET keyword option is active. Provide as many cards as necessary. This input ends at the next keyword ("*") card.

| Card 1b | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | MSURF_S |  |  |  |  |  |  |  |
| Type | I |  |  |  |  |  |  |  |
| Default | none |  |  |  |  |  |  |  |

VARIABLE
MSURF_S

## DESCRIPTION

ID of a set of mesh surface parts created with a *LSO_ID_SET card, where each mesh surface part ID in the set is referenced in *MESH_SURFACE_ELEMENT cards.

Set Card. Card 1 format used when the SET keyword option is active. Provide as many cards as necessary. This input ends at the next keyword ("*") card.

| Card 1c | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | SSID |  |  |  |  |  |  |  |
| Type | 1 |  |  |  |  |  |  |  |
| Default | none |  |  |  |  |  |  |  |

## VARIABLE

SSID

DESCRIPTION
Segment set ID

Segment Cards. Card 1 format used when SEGMENT keyword option is active. Include an additional card for each corresponding pair of segments. This input ends at the next keyword ("*") card.

| Card 1d | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | N1 | N2 | N3 | N4 |  |  |  |  |
| Type | I | 1 | 1 | 1 |  |  |  |  |
| Default | none | none | none | none |  |  |  |  |

## VARIABLE

## DESCRIPTION

N1, N2, ... Node IDs defining a segment
*CESE_BOUNDARY_SOLID_WALL_OPTION1_\{OPTION2\}
For OPTION1, the choices are:
MSURF
MSURF_SET
SET

## SEGMENT

For OPTION2, the choices are:
<BLANK>
ROTAT
Purpose: Define a solid wall boundary condition for this CESE compressible flow solver. This boundary condition models a solid boundary, that is, the physical boundary for the flow field. This boundary condition is a slip boundary condition for inviscid flows and a no-slip boundary condition for viscous flows.

Use the MSURF or MSURF_SET options when generating the CESE mesh through the *MESH cards. Use the SET or SEGMENT keyword options cards when *ELEMENT_SOLID cards define the CESE mesh. Use the ROTAT keyword option for a rotating solid boundary condition.

## Card Summary:

Card Sets. The following sequence of cards comprises a single set. LS-DYNA continues reading *CESE_BOUNDARY_SOLID_WALL card sets until the next keyword ("*") card.
Card 1a. Include this card when OPTION1 is MSURF and OPTION2 is not set (<BLANK>).

| MSURFID | LCID | VX | VY | VZ |  |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

Card 1b. Include this card when OPTION1 is MSURF and OPTION2 is ROTAT.

| MSURFID | LCID | VX | VY | VZ | NX | NY | NZ |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |

Card 1c. Include this card when OPTION1 is MSURF_SET and OPTION2 is not set (<BLANK>).

| MSURF_S | LCID | VX | VY | VZ |  |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

Card 1d. Include this card when OPTION1 is MSURF_SET and OPTION2 is ROTAT.

| MSURF_S | LCID | $V X$ | $V Y$ | $V Z$ | $N X$ | $N Y$ | $N Z$ |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |

Card 1e. Include this card when OPTION1 is SET and OPTION2 is not set (<BLANK>).

| SSID | LCID | VX | VY | VZ |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |

Card 1f. Include this card when OPTION1 is SET and OPTION2 is ROTAT.

| SSID | LCID | VX | VY | VZ | NX | NY | NZ |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |

Card 1g. Include this card when OPTION1 is SEGMENT and OPTION2 is not set (<BLANK>).

| N1 | N2 | N3 | N4 | LCID | VX | VY | VZ |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

Card 1h.1. Include this card when OPTION1 is SEGMENT and OPTION2 is ROTAT.

| N1 | N2 | N3 | N4 | LCID | VX | VY | VZ |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |

Card 1h.2. Include this card when OPTION1 is SEGMENT and OPTION2 is ROTAT.

| N1 | N2 | N3 |  |  |  |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

## Data Card Definitions:

Surface Part Card (Non-Rotating Boundary). Card 1 format used when the MSURF keyword option is active and the ROTAT keyword option is inactive.

| Card 1a | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | MSURFID | LCID | VX | VY | VZ |  |  |  |
| Type | I | I | F | F | F |  |  |  |
| Default | none | 0 | 0.0 | 0.0 | 0.0 |  |  |  |
| Remarks |  | 2 | 2 | 2 | 2 |  |  |  |



MSURFID

LCID
VX, VY, VZ

## DESCRIPTION

Mesh surface part ID referenced in *MESH_SURFACE_ELEMENT cards.

Load curve ID to define this solid wall boundary movement
Velocity vector of the solid wall:
LCID.EQ.0: Vector defined by (VX, VY, VZ).
LCID.NE.0: Vector defined by both the load curve and (VX, VY, VZ).

Surface Part Card (Rotating Boundary). Card 1 format used when the MSURF keyword option is active and the ROTAT keyword option is active.

| Card 1b | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | MSURFID | LCID | VX | VY | VZ | NX | NY | NZ |
| Type | I | I | F | F | F | F | F | F |
| Default | none | none | 0.0 | 0.0 | 0.0 | 0.0 | 0.0 | 0.0 |


| VARIABLE | DESCRIPTION |
| :---: | :--- |
| MSURFID | Mesh surface part ID referenced in *MESH_SURFACE_EL <br> cards. |
| LCID | Load curve ID giving the rotating speed frequency (HZ) <br> must be specified. |
| NX, NY, NZ | $x, y$, and $z$-coordinates of a point on the rotating axis |

Surface Part Set Card (Non-Rotating Boundary). Card 1 format used when the MSURF_SET keyword option is active and the ROTAT keyword option is not active.

| Card 1c | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | MSURF_S | LCID | VX | VY | VZ |  |  |  |
| Type | I | I | F | F | F |  |  |  |
| Default | none | 0 | 0.0 | 0.0 | 0.0 |  |  |  |
| Remarks |  | 2 | 2 | 2 | 2 |  |  |  |

## VARIABLE

MSURF_S

LCID Load curve ID to define this solid wall boundary movement
VX, VY, VZ Velocity vector of the solid wall:
LCID.EQ.O: Vector defined by (VX, VY, VZ).
LCID.NE.O: Vector defined by both the load curve and (VX, VY, VZ).

Surface Part Set Card (Rotating Boundary). Card 1 format used when the MSURF_SET keyword option is active and the ROTAT keyword option is active.

| Card 1d | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | MSURF_S | LCID | VX | VY | VZ | NX | NY | NZ |
| Type | I | I | F | F | F | $F$ | $F$ | $F$ |
| Default | none | 0 | 0.0 | 0.0 | 0.0 | 0.0 | 0.0 | 0.0 |

## VARIABLE

MSURF_S

LCID

NX, NY, NZ

VX, VY, VZ $\quad x, y$, and $z$-coordinates of a point on the rotating axis

## DESCRIPTION

ID of a set of mesh surface part IDs created with a *LSO_ID_SET card, where each mesh surface part ID in the set is referenced in *MESH_SURFACE_ELEMENT cards.

Load curve ID giving the rotating speed frequency (HZ). LCID must be specified.

Unit vector of the rotating axis (not used for the 2D case)

Segment Set Card (Non-Rotating Boundary). Card 1 format used when the SET keyword option is active and the ROTAT keyword option is not active.

| Card 1e | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | SSID | LCID | VX | VY | VZ |  |  |  |
| Type | I | I | F | F | F |  |  |  |
| Default | none | none | 0.0 | 0.0 | 0.0 |  |  |  |
| Remarks |  | 2 | 2 | 2 | 2 |  |  |  |

VARIABLE
DESCRIPTION
Segment set ID
Load curve ID to define this solid wall boundary movement
Velocity vector of the solid wall:
LCID.EQ.0: Vector defined by (VX, VY, VZ).
LCID.NE.O: Vector defined by both the load curve and (VX, VY, VZ).

Segment Set Card (Rotating Boundary). Card 1 format used when the SET keyword option is active and the ROTAT keyword option is active.

| Card 1f | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | SSID | LCID | VX | VY | VZ | NX | NY | NZ |
| Type | I | I | $F$ | $F$ | $F$ | $F$ | $F$ | $F$ |
| Default | none | none | 0.0 | 0.0 | 0.0 | 0.0 | 0.0 | 0.0 |

## VARIABLE

SSID Segment set ID
LCID Load curve ID giving the rotating speed frequency (HZ). LCID must be specified.
$\mathrm{VX}, \mathrm{VY}, \mathrm{VZ} \quad x, y$, and $z$-coordinates of a point on the rotating axis
$\mathrm{NX}, \mathrm{NY}, \mathrm{NZ} \quad$ Unit vector of the rotating axis (not used for the 2D case)
Segment Card (Non-Rotating Boundary). Card 1 format used when the SEGMENT keyword option is active and the ROTAT keyword option is not active.

| Card 1g | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | N1 | N2 | N3 | N4 | LCID | VX | VY | VZ |
| Type | I | I | I | I | I | F | F | F |
| Default | none | none | none | none | 0 | 0.0 | 0.0 | 0.0 |
| Remarks |  |  |  |  | 2 | 2 | 2 | 2 |

## VARIABLE

## DESCRIPTION

N1, N2, Node IDs defining a segment
N3, N4
LCID Load curve ID to define this solid wall boundary movement

## VARIABLE

VX, VY, VZ Velocity vector of the solid wall:
LCID.EQ.0: Vector defined by (VX, VY, VZ).
LCID.NE.0: Vector defined by both the load curve and (VX, VY, VZ).

Segment Card (Rotating Boundary). Card 1 format used when the SEGMENT keyword option is active and the ROTAT keyword option is active.

| Card 1h.1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | N 1 | N 2 | N 3 | N 4 | LCID | VX | VY | VZ |
| Type | I | I | I | I | I | F | F | F |
| Default | none | none | none | none | 0 | 0.0 | 0.0 | 0.0 |

Rotating Axis Card for Segment. Additional card for the "Segment Card (Rotating Boundary)" case that is read when the ROTAT keyword option is used.

| Card 1h.2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | NX | NY | NZ |  |  |  |  |  |
| Type | F | F | F |  |  |  |  |  |
| Default | 0.0 | 0.0 | 0.0 |  |  |  |  |  |

VARIABLE

## DESCRIPTION

N1, N2, $\quad$ Node IDs defining a segment
N3, N4
LCID Load curve ID giving the rotating speed frequency (HZ). LCID must be specified.

VX, VY, VZ $\quad x, y$, and $z$-coordinates of a point on the rotating axis
NX, NY, NZ Unit vector of the rotating axis (not used for the 2D case)

## Remarks:

1. Boundary movement. In this solid-wall condition, the boundary movement can only be in the tangential direction of the wall and should not affect the fluid domain size and mesh during the calculation. Otherwise, you should use an FSI or moving mesh solver. Also, this moving solid boundary condition only affects viscous flows (no-slip BC).
2. Regular solid wall boundary condition. If $L C I D=0$ and $V X=V Y=V Z=0.0$ (default), this boundary condition is a regular solid wall boundary condition.

Purpose: Cause mass fractions of the listed chemical species to be added to the CESE d3plot output. This is only used when chemistry is being solved with the CESE solver.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | MODELID |  |  |  |  |  |  |  |
| Type | I |  |  |  |  |  |  |  |
| Default | none |  |  |  |  |  |  |  |

Species Cards. Include one card for each species to be included in the d3plot database. This input ends at the next keyword ("*") card.

| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | SPECIES |  |  |  |  |  |  |  |
| Type |  |  |  |  |  |  |  |  |

VARIABLE
MODELID
SPECIES Name of a chemical species that is defined in the chemistry model identified by MODELID (see *CHEMISTRY_MODEL).

## *CESE_CONTROL_LIMITER

Purpose: Sets some stability parameters used in the CESE scheme for this CESE compressible flow solver.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | IDLMT | ALFA | BETA | EPSR |  |  |  |  |
| Type | I | F | F | F |  |  |  |  |
| Default | 0 | 0.0 | 0.0 | 0.0 |  |  |  |  |
| Remarks |  | 1 | 2 | 3 |  |  |  |  |

VARIABLE
IDLMT

ALFA Re-weighting coefficient (See CESE theory manual)
BETA Numerical viscosity control coefficient (See CESE theory manual)
EPSR Stability control coefficient (See CESE theory manual)

## Remarks:

1. $\alpha \geq 0$; larger values give more stability, but less accuracy. Usually $\alpha=2.0$ or 4.0 will be enough for normal shock problems.
2. $0 \leq \beta \leq 1$; larger values give more stability. For problems with shock waves, $\beta=1.0$ is recommended.
3. $\varepsilon \geq 0$; larger values give more stability, but less accuracy.

Purpose: For moving mesh CESE, this keyword is used to choose the type of algorithm to be used for calculating mesh movement.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | MMSH | LIM_ITER | RELTOL |  |  |  |  |  |
| Type | 1 | 1 | F |  |  |  |  |  |
| Default | 1 | 100 | $1.0 e-3$ |  |  |  |  |  |

VARIABLE
MMSH

LIM_ITER

RELTOL

## DESCRIPTION

Mesh motion selector:
EQ.1: mesh moves using an implicit ball-vertex spring method.
EQ.9: the IDW scheme is used to move the mesh.

Maximum number of linear solver iterations for the ball-vertex linear system.

Relative tolerance to use as a stopping criterion for the iterative linear solver (conjugate gradient solver with diagonal scaling preconditioner).

## *CESE_CONTROL_SOLVER

Purpose: Set general purpose control variables for the CESE compressible flow solver.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | ICESE | IFLOW | IGEOM | IFRAME | MIXID | IDC | ISNAN |  |
| Type | I | I | 1 | I | I | F | 1 |  |
| Default | 0 | 0 | none | 0 | none | 0.25 | 0 |  |
| Remarks |  |  | 1 |  |  | 2 |  |  |

## VARIABLE

ICESE

IFLOW

IGEOM Sets the geometric dimension:
EQ.2: Two-dimensional (2D) problem
EQ.3: Three-dimensional (3D) problem
EQ.101: 2D axisymmetric

IFRAME Choose the frame of reference:
EQ.O: Usual non-moving reference frame (default)
EQ.1000: Non-inertial rotating reference frame
MIXID Chemistry model ID that defines the chemical species to include in the mixing model (see *CHEMISTRY_MODEL). The species information is given through the model's card specifying the Chemkincompatible input.

VARIABLE
IDC

ISNAN Flag to check for a NaN in the CESE solver solution arrays at the completion of each time step. This option can be useful for debugging purposes. There is a cost overhead when this option is active.

EQ.0: No checking
EQ.1: Checking is active.

## Remarks:

1. 2D Solvers and Mesh Geometry. For the 2D $(\mathrm{IGEOM}=2)$ or 2 D axisymmetric (IGEOM = 101) solver, the mesh should only be distributed in the $x-y$ plane with the boundary conditions given only at the $x-y$ domain boundaries. Otherwise, a warning message will be given, and the 3D solver will be triggered instead.

The 2D axisymmetric case will work only if the 2D mesh and corresponding boundary conditions are properly defined, with the $x$ and $y$ coordinates corresponding to the radial and axial directions, respectively.
2. Contact Interaction Detection Coefficient. IDC is the same type of variable that is input on the *ICFD_CONTROL_FSI card. For an explanation, see Remark 1 for the *ICFD_CONTROL_FSI card.
3. CESE Solver and Restarts. Restarts are supported for the CESE solver. However, restarts do not currently work for simulations with conjugate heat transfer.

## *CESE_CONTROL_TIMESTEP

Purpose: Sets the time-step control parameters for the CESE compressible flow solver.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | IDDT | CFL | DTINT |  |  |  |  |  |
| Type | I | F | F |  |  |  |  |  |
| Default | 0 | 0.9 | $10^{-3}$ |  |  |  |  |  |

CFL CFL number (Courant-Friedrichs-Lewy condition)

## VARIABLE

IDDT

DTINT Initial time step size
Sets the time step option: time step size) step.
( $0.0<\mathrm{CFL} \leq 1.0$ )

## DESCRIPTION

EQ.0: fixed time step size (DTINT, meaning the given initial

NE.O: the time step size will be calculated based on the given CFL-number and the flow solution at the previous time

## *CESE_DATABASE_ELOUT

Purpose: Enable the output of CESE data on elements. If more than one element set is defined, then several output files will be generated.

Output Options Card.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | OUTLV | DTOUT |  |  |  |  |  |  |
| Type | 1 | F |  |  |  |  |  |  |
| Default | 0 | 0. |  |  |  |  |  |  |

Element Sets Card. Include as many cards as needed. This input ends at the next keyword ("*") card.

| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | ELSID |  |  |  |  |  |  |  |
| Type | 1 |  |  |  |  |  |  |  |
| Default | none |  |  |  |  |  |  |  |

VARIABLE
OUTLV

## DESCRIPTION

Determines if the output file should be created:
EQ.O: No output file is generated.
EQ.1: The output file is generated.
DTOUT Time interval to print the output. If DTOUT is equal to 0.0 , then the CESE timestep will be used.

ELSID Solid elements set ID

## Remarks:

1. Database File Name. The file name for this database is cese_elout.dat.
2. Restrictions. The *CESE_DATABASE... cards (pointout/elout) are restricted to CFD-only problems. They do not give correct results for chemically reacting flow where *CESE couples with *CHEMISTRY.

## *CESE_DATABASE_FLUXAVG

Purpose: Enable the output of CESE data on segment sets. If more than one segment set is defined, then several output files will be generated.

Output Options Card.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | OUTLV | DTOUT |  |  |  |  |  |  |
| Type | 1 | F |  |  |  |  |  |  |
| Default | 0 | 0. |  |  |  |  |  |  |

Segment Sets Card. Include as many cards as needed. This input ends at the next keyword ("*") card.

| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | SSID |  |  |  |  |  |  |  |
| Type | 1 |  |  |  |  |  |  |  |
| Default | none |  |  |  |  |  |  |  |

VARIABLE
OUTLV

DTOUT

SSID

DESCRIPTION
Determines if the output file should be created:
EQ.O: No output file is generated.
EQ.1: The output file giving the average fluxes is generated.

Time interval to print the output. If DTOUT is equal to 0.0 , then the CESE time step will be used.

Segment set ID

## Remarks:

1. Database File Name. The file name for this database is cese_fluxavg.dat.
2. Restrictions. The ${ }^{*}$ CESE_DATABASE... cards (pointout/elout/...) are restricted to CFD-only problems. They do not give correct results for chemically reacting flow where *CESE couples with *CHEMISTRY.

## *CESE_DATABASE_FSIDRAG

Purpose: Enable the output of the total fluid pressure force applied to solid parts in FSI problems at every time step.

## Output Options Card.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | OUTLV |  |  |  |  |  |  |  |
| Type | 1 |  |  |  |  |  |  |  |
| Default | 0 |  |  |  |  |  |  |  |

VARIABLE
OUTLV

## DESCRIPTION

Determines if the output file should be created:
EQ.0: No output file is generated.
EQ.1: The output file giving the pressure forces is generated.

## Remarks:

1. Database File Names. The file names for this database are cese_dragsol.dat, cese_dragshell.dat, cese_dragsol2D.dat and cese_dragbeam.dat, depending on what kind of solid is used.

## *CESE_DATABASE_POINTOUT

Purpose: Enable the output of CESE data at points.
Output Options Card.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | PSID | DTOUT | PSTYPE | VX | VY | VZ |  |  |
| Type | I | F | I | F | F | F |  |  |
| Default | 0 | 0. | 0 | 0. | 0. | 0. |  |  |

Point Specification Card. Include as many cards as needed. This input ends at the next keyword ("*") card.

| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | PID | X | Y | Z |  |  |  |  |
| Type | 1 | F | F | F |  |  |  |  |
| Default | none | none | none | none |  |  |  |  |
| VARIABLE |  | DESCRIPTION |  |  |  |  |  |  |
| PSID |  | Point set ID |  |  |  |  |  |  |
| DTOUT |  | Time interval to print the output. If DTOUT is equal to 0.0 , then the CESE timestep will be used. |  |  |  |  |  |  |
| PSTYPE |  | Point set type: |  |  |  |  |  |  |
|  |  | EQ.0: Fixed points |  |  |  |  |  |  |
|  |  | EQ.1: Tracer points using prescribed velocity |  |  |  |  |  |  |
|  |  | EQ.2: Tracer points using fluid velocity |  |  |  |  |  |  |
| VX, VY, VZ |  | Constant velocities to be used when PSTYPE $=1$ |  |  |  |  |  |  |
| PID |  | Point ID |  |  |  |  |  |  |

## VARIABLE

$X, Y, Z \quad$ Initial coordinates of the point

## Remarks:

1. Database File Name. The file name for this database is cese_pointout.dat.
2. Restrictions. The *CESE_DATABASE... cards (pointout/elout) are restricted to CFD-only problems. They do not give correct results for chemically reacting flow where *CESE couples with *CHEMISTRY.

## *CESE_DATABASE_SSETDRAG

Purpose: Enable the output of CESE drag forces on segment sets. If more than one segment set is specified, then several output files will be generated.

## Output Options Card.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | OUTLV | DTOUT |  |  |  |  |  |  |
| Type | 1 | F |  |  |  |  |  |  |
| Default | 0 | 0. |  |  |  |  |  |  |

Segment Sets Card. Include as many cards as needed. This input ends at the next keyword ("*") card.

| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | SSID |  |  |  |  |  |  |  |
| Type | 1 |  |  |  |  |  |  |  |
| Default | none |  |  |  |  |  |  |  |

VARIABLE
OUTLV

DTOUT

SSID

DESCRIPTION
Determines if the output file should be created:
EQ.O: No output file is generated.
EQ.1: The output file giving the average fluxes is generated.

Time interval to print the output. If DTOUT is equal to 0.0 , then the CESE timestep will be used.

Segment set ID

## Remarks:

1. Database File Name. The file name for this database is cese_ssetdrag.dat.
2. Friction Drag Results and Mesh Size. For the friction drag to give consistent results, special care must be given to the mesh close to the solid wall boundary (good capturing of the boundary layer behavior). We recommend a very fine structured mesh.

## *CESE_DEFINE_NONINERTIAL

Purpose: Define the CESE problem domain as a non-inertial rotating frame that rotates at a constant rate. This is used in rotating problems such as spinning cylinders, wind turbines and turbo machinery.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | FREQ | LCID | PID | Nx | Ny | Nz |  |  |
| Type | F | I | I | F | F | F |  |  |
| Default | none | 0 | none | none | none | none |  |  |


| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | L | R | RELV |  |  |  |  |  |
| Type | F | F | I |  |  |  |  |  |
| Default | none | none | 0 |  |  |  |  |  |

VARIABLE
FREQ
LCID Load curve ID for scaling factor of FREQ.
PID Starting point ID for the reference frame (See *CESE_DEFINE_POINT).
$\mathrm{Nx}, \mathrm{Ny}, \mathrm{Nz} \quad$ Rotating axis direction.
L Length of rotating frame.
$\mathrm{R} \quad$ Radius of rotating frame.

RELV

## DESCRIPTION

Velocity display mode:
EQ.0: Relative velocity, only the non-rotating components of the velocity are output.

EQ.1: Absolute velocity is output.

## *CESE_DEFINE_POINT

Purpose: Define points to be used by the CESE solver.
Point Cards. Include one card for each point. This input ends at the next keyword ("*") card.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | NID | X | Y | Z |  |  |  |  |
| Type | I | F | F | F |  |  |  |  |
| Default | none | none | none | none |  |  |  |  |


| VARIABLE |  |
| :---: | :--- |
| NID |  |
| Identifier for this point. |  |
| Coordinates of the point. |  |

*CESE_DRAG
Purpose: Provide the far-field (or free-stream) fluid pressure.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | PRESS |  |  |  |  |  |  |  |
| Type | F |  |  |  |  |  |  |  |
| VARIABLE |  |  |  |  |  |  |  |  |
| DESCRIPTION |  |  |  |  |  |  |  |  |

PRESS
Value of the free-stream fluid pressure (in units used by the current problem).

## *CESE_EOS_CAV_HOMOG_EQUILIB

Purpose: Define the coefficients in the equation of state (EOS) for the homogeneous equilibrium cavitation model.

Include as many cards as needed. This input ends at the next keyword ("*") card.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | EOSID | $\rho_{\text {vap }}$ | $\rho_{\text {liq }}$ | $a_{\text {vap }}$ | $a_{\text {liq }}$ | $\mu_{\text {vap }}$ | $\mu_{\text {liq }}$ | $P_{\text {Satrap }}$ |
| Type | I | F | F | F | F | F | F | F |
| Default | none | 0.8 | 880.0 | 334.0 | 1386.0 | $1.435 \mathrm{e}-$ <br> 5 | $1.586 \mathrm{e}-$ <br> 4 | $1.2 \mathrm{e}+4$ |

## VARIABLE

EOSID
$\rho_{\text {vap }} \quad$ density of the saturated vapor
$\rho_{\text {liq }} \quad$ density of the saturated liquid
$a_{\text {vap }} \quad$ sound speed of the saturated vapor
$a_{\text {liq }} \quad$ sound speed of the saturated liquid
$\mu_{\text {vap }} \quad$ dynamic viscosity of the vapor
$\mu_{\text {liq }}$
$P_{\text {SatVap }}$

## DESCRIPTION

Equation of state identifier
dynamic viscosity of the liquid
pressure of the saturated vapor

## Remarks:

1. Once a cavitation EOS is used, the cavitation flow solver will be triggered.
2. In this homogeneous equilibrium cavitation model, a barotropic equation of state is used. This model can be used in small scale \& high speed cavitation flows, and it is not good for large-scale, low-speed cavitation calculations.

## *CESE_EOS_IDEAL_GAS

Purpose: Define the coefficients Cv and Cp in the equation of state for an ideal gas in the CESE fluid solver.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | EOSID | Cv | Cp |  |  |  |  |  |
| Type | I | F | F |  |  |  |  |  |
| Default | none | 717.5 | 1004.5 |  |  |  |  |  |

VARIABLE
EOSID
$\mathrm{Cv} \quad$ Specific heat at constant volume
Cp Specific heat at constant pressure

## Remarks:

1. Units. As with other solvers in LS-DYNA, the user is responsible for unit consistency. For example, if a user wants to use dimensionless variables, Cv and Cp should also be replaced by the corresponding dimensionless ones.

## *CESE_EOS_INFLATOR1

Purpose: To define an EOS using Cp and Cv thermodynamic expansions for an inflator gas mixture with a single temperature range.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | EOSID |  |  |  |  |  |  |  |
| Type | 1 |  |  |  |  |  |  |  |
| Default | none |  |  |  |  |  |  |  |


| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | Cp0 | Cp1 | Cp2 | Cp3 | Cp4 |  |  |  |
| Type | F | F | F | F | F |  |  |  |
| Default | 0. | 0. | 0. | 0. | 0. |  |  |  |


| Card 3 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | Cv0 | Cv1 | Cv2 | Cv3 | Cv4 |  |  |  |
| Type | F | F | F | F | F |  |  |  |
| Default | 0. | 0. | 0. | 0. | 0. |  |  |  |

## VARIABLE

EOSID
Cp0, ..., Cp4

## DESCRIPTION

Equation of state identifier for the CESE solver.
Coefficients of temperature-dependent specific heat at constant pressure

$$
C_{p}(T)=C_{p 0}+C_{p 1} T+C_{p 2} T^{2}+C_{p 3} T^{3}+C_{p 4} T^{4}
$$

## VARIABLE

$\mathrm{Cv} 0, \ldots, \mathrm{Cv} 4 \quad$ Coefficients of temperature-dependent specific heat at constant volume

$$
\mathrm{C}_{\mathrm{v}}(\mathrm{~T})=\mathrm{C}_{\mathrm{v} 0}+\mathrm{C}_{\mathrm{v} 1} \mathrm{~T}+\mathrm{C}_{\mathrm{v} 2} \mathrm{~T}^{2}+\mathrm{C}_{\mathrm{v} 3} \mathrm{~T}^{3}+\mathrm{C}_{\mathrm{v} 4} \mathrm{~T}^{4}
$$

## Remark:

1.These coefficient expansions for the specific heats over the entire temperature range are generated by the 0-D inflator model solver. See *CHEMISTRY_CONTROL_INFLATOR and *CHEMISTRY_INFLATOR_PROPERTIES for details related to running that solver.

## *CESE_EOS_INFLATOR2

Purpose: To define an EOS using Cp and Cv thermodynamic expansions for an inflator gas mixture with two temperature ranges, one below 1000 degrees Kelvin, and the other above 1000 degrees Kelvin.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | EOSID |  |  |  |  |  |  |  |
| Type | I |  |  |  |  |  |  |  |
| Default | none |  |  |  |  |  |  |  |

Card for the expansion of Specific Heat at Constant Pressure. Valid for $\mathrm{T}<1000{ }^{0} \mathrm{~K}$

| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | Cp1_0 | Cp1_1 | Cp1_2 | Cp1_3 | Cp1_4 |  |  |  |
| Type | F | F | F | F | F |  |  |  |
| Default | 0. | 0. | 0. | 0. | 0. |  |  |  |

Card for the expansion of Specific Heat at Constant Pressure. Valid for T > $1000{ }^{\circ} \mathrm{K}$.

| Card 3 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | Cp2_0 | Cp2_1 | Cp2_2 | Cp2_3 | Cp2_4 |  |  |  |
| Type | F | F | F | F | F |  |  |  |
| Default | 0. | 0. | 0. | 0. | 0. |  |  |  |

Card for the expansion of Specific Heat at Constant Volume. Valid for $\mathrm{T}<1000{ }^{0} \mathrm{~K}$

| Card 4 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | Cv1_0 | Cv1_1 | Cv1_2 | Cv1_3 | Cv1_4 |  |  |  |
| Type | F | F | F | F | F |  |  |  |
| Default | 0. | 0. | 0. | 0. | 0. |  |  |  |

Card for the expansion of Specific Heat at Constant Volume. Valid for $\mathrm{T}>1000{ }^{0} \mathrm{~K}$.

| Card 5 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | CV2_0 | Cv2_1 | Cv2_2 | Cv2_3 | Cv2_4 |  |  |  |
| Type | F | F | F | F | F |  |  |  |
| Default | 0. | 0. | 0. | 0. | 0. |  |  |  |

VARIABLE
EOSID

Cp1_0,...,
Cp1_4

Cp2_0,...,
Cp2_4

## DESCRIPTION

Equation of state identifier for the CESE solver.
Coefficients of temperature-dependent specific heat at constant pressure valid for $\mathrm{T}<1000{ }^{0} \mathrm{~K}$.
$\mathrm{C}_{\mathrm{p} 1}(\mathrm{~T})=\mathrm{C}_{\mathrm{p} 1 \_0}+\mathrm{C}_{\mathrm{p} 1 \_1} \mathrm{~T}+\mathrm{C}_{\mathrm{p} 1 \_2} \mathrm{~T}^{2}+\mathrm{C}_{\mathrm{p} 1 \_3} \mathrm{~T}^{3}+\mathrm{C}_{\mathrm{p} 1 \_4} \mathrm{~T}^{4}$
Coefficients of temperature-dependent specific heat at constant pressure valid for $\mathrm{T}>1000^{0} \mathrm{~K}$.
$\mathrm{C}_{\mathrm{p} 2}(\mathrm{~T})=\mathrm{C}_{\mathrm{p} 2 \_0}+\mathrm{C}_{\mathrm{p} 2_{-} 1} \mathrm{~T}+\mathrm{C}_{\mathrm{p} 2_{2} 2} \mathrm{~T}^{2}+\mathrm{C}_{\mathrm{p} 2^{2} 3} \mathrm{~T}^{3}+\mathrm{C}_{\mathrm{p} 2^{2} 4} \mathrm{~T}^{4}$
Cv1_0, ...,
Cv1_4
Coefficients of temperature-dependent specific heat at constant volume valid for $\mathrm{T}<1000{ }^{0} \mathrm{~K}$.

$$
\mathrm{C}_{\mathrm{v} 1}(\mathrm{~T})=\mathrm{C}_{\mathrm{v} 1 \_0}+\mathrm{C}_{\mathrm{v} 1 \_1} \mathrm{~T}+\mathrm{C}_{\mathrm{v} 1 \_2} \mathrm{~T}^{2}+\mathrm{C}_{\mathrm{v} 1 \_3} \mathrm{~T}^{3}+\mathrm{C}_{\mathrm{v} 1 \_4} \mathrm{~T}^{4}
$$

Cv2_0,..., Cv2_4

Coefficients of temperature-dependent specific heat at constant volume valid for $\mathrm{T}>1000{ }^{\circ} \mathrm{K}$.

$$
\mathrm{C}_{\mathrm{v} 2}(\mathrm{~T})=\mathrm{C}_{\mathrm{v} 2^{\prime} 0}+\mathrm{C}_{\mathrm{v} 2 \_1} \mathrm{~T}+\mathrm{C}_{\mathrm{v} 2 \_2} \mathrm{~T}^{2}+\mathrm{C}_{\mathrm{v} 2 \_3} \mathrm{~T}^{3}+\mathrm{C}_{\mathrm{v} 2^{4}} \mathrm{~T}^{4}
$$

## Remark:

2.These coefficient expansions for the specific heats over two temperature ranges are generated by the 0-D inflator model solver. See *CHEMISTRY_CONTROL_INFLATOR and *CHEMISTRY_INFLATOR_PROPERTIES for details related to running that solver.

## *CESE_FSI_EXCLUDE

Purpose: Provide a list of mechanics solver parts that are not involve in the CESE FSI calculation. This is intended to be used as an efficiency measure for parts that will not involve significant FSI interactions with the CESE compressible fluid solver..

Include as many cards as needed. This input ends at the next keyword ("*") card.

| Card1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | PID1 | PID2 | PID3 | PID4 | PID5 | PID6 | PID7 | PID8 |
| Type | I | I | 1 | 1 | 1 | 1 | 1 | 1 |
| Default | none | none | none | none | none | none | none | none |

VARIABLE
PIDn

## DESCRIPTION

IDs of mechanics parts that will be excluded from the FSI interaction calculation with the CESE solver.

## *CESE_INITIAL

Purpose: Specify constant initial conditions (ICs) for flow variables at the centroid of each fluid element.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | U | V | W | RH | P | T |  |  |
| Type | F | F | F | F | F | F |  |  |
| Default | 0 | 0.0 | 0.0 | 1.225 | 0.0 | 0.0 |  |  |

## VARIABLE

U, V, W
RHO
P Pressure, $P$
T
Temperature, $T$

## Remarks:

1. Required Input. Usually, only two of $\rho, P$, and $T$ need to be specified (besides the velocity). If all three are given, only $\rho$ and $P$ will be used.
2. Applicable Elements. These initial conditions will be applied only in those elements that have not been assigned a value by *CESE_INITIAL_OPTION cards for individual elements or sets of elements.

## *CESE_INITIAL_OPTION

Available options include:
SET

## ELEMENT

Purpose: Specify initial conditions for the flow variables at the centroid of each element in a set of elements or at the centroid of a single element.

Include as many cards as needed. This input ends at the next keyword ("*") card.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | EID/ESID | U | V | W | RHO | P | T |  |
| Type | I | F | F | F | F | F | F |  |
| Default | none | 0.0 | 0.0 | 0.0 | 1.225 | 0.0 | 0.0 |  |
| Remarks |  |  |  |  | 1 | 1 | 1 |  |

VARIABLE
EID/ESID Solid element ID (EID) or solid element set ID (ESID)
U, V, W $\quad x-, y-, z$-velocity components, respectively
RHO Density, $\rho$
P Pressure, $P$
T Temperature, $T$

## Remarks:

1. Required Input. Usually, only two of $\rho, P$, and $T$ need to be specified (along with the velocity). If all three are given, only $\rho$ and $P$ will be used.
2. Initial Condition Specification Priority. The priority of this card is higher than *CESE_INITIAL, meaning that if an element is assigned an initial value by this card, *CESE_INITIAL will no longer apply to that element.

## *CESE_INITIAL_CHEMISTRY

Purpose: Initializes the chemistry and fluid state in every element of the CESE mesh that has not already been initialized by one of the other *CESE_INITIAL_CHEMISTRY cards. This is only used when chemistry is being solved with the CESE solver.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | CHEMID | COMPID |  |  |  |  |  |  |
| Type | 1 | 1 |  |  |  |  |  |  |
| Default | none | none |  |  |  |  |  |  |


| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | UIC | VIC | WIC | RHOIC | PIC | TIC | HIC |  |
| Type | F | F | F | F | F | F | F |  |
| Default | none | none | none | none | none | none | none |  |

## VARIABLE

## DESCRIPTION

CHEMID Identifier of chemistry control card to use
COMPID Identifier of chemical composition to use
UIC $\quad X$-component of the fluid velocity
VIC $\quad Y$-component of the fluid velocity
WIC Z-component of the fluid velocity
RHOIC Initial fluid density
PIC Initial fluid pressure
TIC Initial fluid temperature

HIC

## DESCRIPTION

Initial fluid enthalpy. However, when CHEMID refers to a ZND 1-step reaction card, this is the progressive variable (degree of combustion).

## *CESE_INITIAL_CHEMISTRY_ELEMENT

Purpose: Initializes the chemistry and fluid state in every element of the list of CESE elements. This is only used when chemistry is being solved with the CESE solver.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | CHEMID | COMPID |  |  |  |  |  |  |
| Type | 1 | 1 |  |  |  |  |  |  |
| Default | none | none |  |  |  |  |  |  |


| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | UIC | VIC | WIC | RHOIC | PIC | TIC | HIC |  |
| Type | F | F | F | F | F | F | F |  |
| Default | none | none | none | none | none | none | none |  |

Element List Card. Include as many cards as necessary. This input ends at the next keyword ("*") card.

| Card 3 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | ELE1 | ELE2 | ELE3 | ELE4 | ELE5 | ELE6 | ELE7 | ELE8 |
| Type | 1 | 1 | 1 | 1 | 1 | 1 | 1 | 1 |

VARIABLE
CHEMID Identifier of chemistry control card to use
COMPID Identifier of chemical composition to use
UIC $\quad X$-component of the fluid velocity
VIC $\quad Y$-component of the fluid velocity

## VARIABLE

WIC
RHOIC Initial fluid density
PIC Initial fluid pressure
TIC Initial fluid temperature
HIC Initial fluid enthalpy. However, when CHEMID refers to a ZND 1-step reaction card, this is the progressive variable (degree of combustion).

ELE $i \quad$ User element numbers to initialize

## *CESE_INITIAL_CHEMISTRY_PART

Purpose: Initializes the chemistry and fluid state in every element of the specified CESE part that has not already been initialized by *CESE_INITIAL_CHEMISTRY_ELEMENT or *CESE_INITIAL_CHEMISTRY_SET cards. This is only used when chemistry is being solved with the CESE solver.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | PARTID | CHEMID | COMPID |  |  |  |  |  |
| Type | 1 | 1 | 1 |  |  |  |  |  |
| Default | none | none | none |  |  |  |  |  |


| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | UIC | VIC | WIC | RHOIC | PIC | TIC | HIC |  |
| Type | F | F | F | F | F | F | F |  |
| Default | none | none | none | none | none | none | none |  |

## VARIABLE

## DESCRIPTION

PARTID Identifier of the CESE part on which to initialize.
CHEMID Identifier of chemistry control card to use.
COMPID Identifier of chemical composition to use.
UIC X-component of the fluid velocity.
VIC Y-component of the fluid velocity.
WIC Z-component of the fluid velocity.
RHOIC Initial fluid density.
PIC Initial fluid pressure.
TIC Initial fluid temperature.

VARIABLE
HIC

## DESCRIPTION

Initial fluid enthalpy. However, when CHEMID refers to a ZND 1-step reaction card, this is the progressive variable (degree of combustion).

## *CESE_INITIAL_CHEMISTRY_SET

Purpose: Initializes the chemistry and fluid state in every element of the specified element set in the CESE mesh that has not already been initialized by *CESE_INITIAL_CHEMISTRY_ELEMENT cards. This is only used when chemistry is being solved with the CESE solver.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | SETID | CHEMID | COMPID |  |  |  |  |  |
| Type | 1 | 1 | 1 |  |  |  |  |  |
| Default | none | none | none |  |  |  |  |  |


| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | UIC | VIC | WIC | RHOIC | PIC | TIC | HIC |  |
| Type | F | F | F | F | F | F | F |  |
| Default | none | none | none | none | none | none | none |  |

## VARIABLE

## DESCRIPTION

SETID Identifier of the CESE element set to initialize.
CHEMID Identifier of chemistry control card to use.
COMPID Identifier of chemical composition to use.
UIC X-component of the fluid velocity.
VIC Y-component of the fluid velocity.
WIC Z-component of the fluid velocity.
RHOIC Initial fluid density.
PIC Initial fluid pressure.
TIC Initial fluid temperature.

VARIABLE
HIC

## DESCRIPTION

Initial fluid enthalpy. However, when CHEMID refers to a ZND 1-step reaction card, this is the progressive variable (degree of combustion).
*CESE_MAT_000
Purpose: Define the fluid (gas) properties in a viscous flow for the CESE solver.
Material Definition Cards. Include one card for each instance of this material type. This input ends at the next keyword ("**) card.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | MID | MU | K |  |  |  |  |  |
| Type | I | F | F |  |  |  |  |  |
| Default | none | none | none |  |  |  |  |  |

## VARIABLE

MID
MU Fluid dynamic viscosity. For Air at $15{ }^{\circ} \mathrm{C}, \mathrm{MU}=1.81 \times$ $10^{-5} \mathrm{~kg} / \mathrm{ms}$

K
Thermal conductivity of the fluid

## Remarks:

1. The viscosity is only used viscous flows, so for inviscid flows, it is not necessary to define it. The thermal conductivity is only used to calculate the heat transfer between the structure and the thermal solver when coupling is activated.
2. As with other solvers in LS-DYNA, the user is responsible for unit consistency. For example, if dimensionless variables are used, MU should be replaced by the corresponding dimensionless one.

## *CESE_MAT_001(_GAS)

Purpose: Define the fluid (gas) properties in a viscous flow for the CESE solver.
Include as many cards as needed. This input ends at the next keyword ("*") card.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | MID | C1 | C2 | PRND |  |  |  |  |
| Type | I | F | F | F |  |  |  |  |
| Default | none | $1.458 \mathrm{E}-$ <br> 6 | 110.4 | 0.72 |  |  |  |  |

VARIABLE
MID
C1, C2
Two coefficients in the Sutherland's formula for viscosity, i.e.,

$$
\mu=\frac{C_{1} T^{\frac{3}{2}}}{T+C_{2}}
$$

where $C_{1}$ and $C_{2}$ are constants for a given gas. For example, for air at moderate temperatures,

$$
C_{1}=1.458 \times 10^{-6} \mathrm{~kg} / \mathrm{msK}^{1 / 2}, \quad C_{2}=110.4 \mathrm{~K}
$$

PRND
The Prandtl Number (used to determine the coefficient of thermal conductivity). It is approximately constant for most gases. For air at standard conditions PRND $=0.72$.

## Remarks:

1. C 1 and C 2 are only used to calculate the viscosity in viscous flows, so for inviscid flows, this material card is not needed. The Prandtl number is used to extract the thermal conductivity, which is used when thermal coupling with the structure is activated.
2. As with other solvers in LS-DYNA, the user is responsible for unit consistency. For example, if dimensionless variables are used, $\mathrm{C}_{1}$ and $\mathrm{C}_{2}$ should be replaced by the corresponding dimensionless ones.

## *CESE_MAT_002

Purpose: Define the fluid (gas) properties in a viscous flow for the CESE solver.
Material Definition Cards. Include one card for each instance of this material type. This input ends at the next keyword ("**") card.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | MID | MU0 | SMU | K0 | SK | T0 |  |  |
| Type | I | F | F | F | F | F |  |  |
| Default | none | $1.716 \mathrm{E}-5$ | 111. | 0.0241 | 194.0 | 273.0 |  |  |

## VARIABLE

MID Material identifier
MU0 / SMU

## DESCRIPTION

Two coefficients appearing in the equation derived by combining Sutherland's formula with the power law for dilute gases:

$$
\frac{\mu}{\mu_{0}}=\left(\frac{T}{T_{0}}\right)^{3 / 2} \frac{T_{0}+S_{\mu}}{T+S_{\mu}} .
$$

$\mu_{0}$ is a reference value, and $S_{\mu}$ is an effective temperature called the Sutherland constant, which is characteristic of the gas. For air at moderate temperatures,

$$
\mu_{0}=1.716 \times 10^{-5} \mathrm{Ns} / \mathrm{m}^{2}, \quad S_{\mu}=111 \mathrm{~K}
$$

K0/SK Two coefficients appearing in the equation derived by combining Sutherland's formula with the power law for dilute gases:

$$
\frac{k}{k_{0}}=\left(\frac{T}{T_{0}}\right)^{3 / 2} \frac{T_{0}+S_{k}}{T+S_{k}}
$$

Here $k$ is the thermal conductivity, $k_{0}$ is a reference value, and $S_{k}$ is the Sutherland constant, which is characteristic of the gas. For air at moderate temperatures,

$$
k_{0}=0.0241 \mathrm{~W} / \mathrm{m}, \quad S_{k}=194 \mathrm{~K}
$$

Reference temperature, $T_{0}$. The default value (273.0) is for air, in degrees K.

## Remarks:

1. Fields that Depend on Problem Physics. The viscosity is only used for viscous flow. Therefore, for inviscid flows, it is not necessary to define it. The thermal conductivity is only used to calculate the heat transfer between the structure and the thermal solver when coupling is activated.
2. Unit Consistency. As with other solvers in LS-DYNA, the user is responsible for unit consistency. For example, if dimensionless variables are used, MU should be replaced by the corresponding dimensionless one.

## *CESE_PART

Purpose: Define CESE solver parts, i.e., connect CESE material and EOS information.
Part Cards. Include one card for each CESE part. This input ends at the next keyword ("*") card.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | PID | MID | EOSID |  |  |  |  |  |
| Type | 1 | 1 | 1 |  |  |  |  |  |
| Default | none | none | none |  |  |  |  |  |

## VARIABLE

PID
MID
EOSID

## DESCRIPTION

Part identifier (must be different from any PID on a *PART card)
Material identifier defined by a *CESE_MAT_... card
Equation of state identifier defined by a *CESE_EOS_... card

## Remarks:

1. Since material coefficients are only used in viscous flows, the MID can be left blank for inviscid flows.

## *CESE_SURFACE_MECHSSID_D3PLOT

Purpose: Identify the surfaces to be used in generating surface D3PLOT output for the CESE solver. These surfaces must be on the outside of volume element parts that are in contact with the CESE fluid mesh. The variables in question are part of the CESE FSI solution process or of the CESE conjugate heat transfer solver.

Include as many cards as needed. This input ends at the next keyword ("*") card.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | SSID | SurfaceLabel |  |  |  |  |  |
| Type | I | A |  |  |  |  |  |
| Default | none | none |  |  |  |  |  |

VARIABLE
SSID

SurfaceLabel Name to use in d3plot output to identify the SSID for the LSPP user.

## *CESE_SURFACE_MECHVARS_D3PLOT

Purpose: List of variables to output on the surfaces designated by the segment set IDs given in the *CESE_SURFACE_MECHSSID_D3PLOT cards. Most of the allowed variables are defined only on the fluid-structure interface, and so the segment set IDs defining a portion of the fluid-structure interface must involve only segments (element faces) that are on the outside of volume element parts that are in contact with the CESE fluid mesh.

Include as many cards as needed. This input ends at the next keyword ("*") card.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |
| Variable | Output Quantity |  |  |  |  |  |  |  |
| Type | A |  |  |  |  |  |  |  |
| Default | none |  |  |  |  |  |  |  |

## VARIABLE

Output
Quantity

## DESCRIPTION

Descriptive phrase for the mechanics surface variable to output for the LSPP user. Output will be done on all SSIDs selected by the *CESE_SURFACE_MECHSSID_D3PLOT cards in the problem.
Supported variables include:
FLUID FSI FORCE
FLUID FSI PRESSURE
INTERFACE TEMPERATURE
SOLID INTERFACE HEAT FLUX
FLUID INTERFACE HEAT FLUX
INTERFACE HEAT FLUX RATE
SOLID INTERFACE DISPLACEMENT
SOLID INTERFACE VELOCITY
SOLID INTERFACE ACCELERATION

Force, displacement, velocity, and acceleration are output as vector quantities. The rest of the variables are scalar quantities. The fluxes are in the normal direction to the fluid/structure interface, with the heat fluxes relative to the normal pointing into the structure.

## *CHEMISTRY

The keyword *CHEMISTRY is used to access chemistry databases that include Chemkinbased descriptions of a chemical model, as well as to select a method of solving the model. The keyword cards in this section are defined in alphabetical order:

*CHEMISTRY_BATTERY ${ }^{\dagger}$<br>*CHEMISTRY_COMPOSITION<br>*CHEMISTRY_CONTROL_OD<br>*CHEMISTRY_CONTROL_1D ${ }^{\dagger}$<br>*CHEMISTRY_CONTROL_CSP<br>*CHEMISTRY_CONTROL_FULL<br>*CHEMISTRY_CONTROL_INFLATOR ${ }^{\dagger}$<br>*CHEMISTRY_CONTROL_TBX<br>*CHEMISTRY_CONTROL_ZND ${ }^{\dagger}$<br>*CHEMISTRY_DET_INITIATION ${ }^{\dagger}$<br>*CHEMISTRY_INFLATOR_PROPERTIES ${ }^{\dagger}$<br>*CHEMISTRY_MODEL<br>*CHEMISTRY_PATH

$\dagger$ : Card may be used only once in a given model
An additional option "_TITLE" may be appended to all *CHEMISTRY keywords. If this option is used, then an 80 character string is read as a title from the first card of that keyword's input. At present, LS-DYNA does not make use of the title. Inclusion of titles gives greater clarity to input decks.

The *CHEMISTRY_BATTERY card is only intended to be used in battery electrochemistry models that involve *BATTERY_... cards. Note that none of the comments below apply for this particular case.

## *CHEMISTRY

In order to use one of the chemistry solvers, the input must include at least one * CHEM ISTRY_MODEL card. For each spatial region containing a different chemical composition, at least one *CHEMISTRY_COMPOSITION card is required.

The *CHEMISTRY_CONTROL_0D card is intended to be used in a standalone fashion to verify the validity of a given chemistry model. This model includes the total number of species and all elementary reactions with their Arrhenius rate parameters. For instance, this solver could be used to check the induction time of the model.

The *CHEMISTRY_BLAST_INITIATION, *CHEMISTRY_CONTROL_1D, *CHEMISTRY_DET_INITIATION, and *CHEMISTRY_CONTROL_ZND cards are intended to provide a one-dimensional initialization to a 2 D or 3D chemically-reacting flow.

In order to perform a full, general purpose chemistry calculation in 2D or 3D, the * CHEM ISTRY_CONTROL_FULL card should be used.

The *CHEMISTRY_CONTROL_CSP card is an option for reducing the number of species and reactions that are used in a general purpose chemistry calculation. Other reduction mechanisms are planned for the future.

An airbag inflator model is available with *CHEMISTRY_CONTROL_INFLATOR along with *CHEMISTRY_INFLATOR_PROPERTIES and a chemistry model that is referenced via three chemical compositions. This involves zero-dimensional modeling, with pyrotechnic inflator, and cold and hot flow hybrid inflator options.

The *CHEMISTRY_CONTROL_TBX card is intended for use only in a stochastic particle model, where the *STOCHASTIC_TBX_PARTICLES card is used.

## *CHEMISTRY_BATTERY

Purpose: For the battery electrochemistry solver (*BATTERY), this card identifies the files that define a Chemkin chemistry model for use with that solver.

Battery Reaction and Species Input File Card (Chemkin-compatible).

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |
| Variable | FILE1 |  |  |  |  |  |  |  |
| Type | A |  |  |  |  |  |  |  |

Thermodynamics Database File Card.

| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |
| Variable | FILE2 |  |  |  |  |  |  |  |
| Type | A |  |  |  |  |  |  |  |

Transport Properties Database File Card.

| Card 3 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |
| Variable | FILE3 |  |  |  |  |  |  |  |
| Type | A |  |  |  |  |  |  |  |

VARIABLE
FILE1
FILE2 Name of the file containing the chemistry thermodynamics database

FILE3 Name of the file containing the chemistry transport properties database

## *CHEMISTRY_COMPOSITION

Purpose: Provides a general way to specify a chemical composition via a list of species mole numbers in the context of a Chemkin database model.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | ID | MODELID |  |  |  |  |  |  |
| Type | I | 1 |  |  |  |  |  |  |
| Default | none | none |  |  |  |  |  |  |

Species List Card. Provide as many cards as necessary. This input ends at the next keyword ("*") card.

| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | MOLFR | SPECIES |  |  |  |  |  |  |
| Type | F |  |  |  |  |  |  |  |
| Default | none | A |  |  |  |  |  |  |
|  |  |  |  |  |  |  |  |  |

VARIABLE
ID
MODELID

MOLFR

SPECIES The Chemkin-compatible name of a chemical species that is defined in the chemistry model identified by MODELID (see *CHEMISTRY_MODEL).
*CHEMISTRY_CONTROL_OD
Purpose: Performs a zero-dimensional isotropic chemistry calculation that operates standalone (does not call the CESE solver). This is for ISOBARIC or ISOCHORIC cases.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | ID | COMPID | SOLTYP | PLOTDT | CSP_SEL |  |  |  |
| Type | I | 1 | 1 | F | 1 |  |  |  |
| Default | none | none | none | $1.0 \mathrm{e}-6$ | 0 |  |  |  |
| Remarks |  |  |  |  | 1 |  |  |  |


| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | DT | TLIMIT | TIC | PIC | RIC | EIC |  |  |
| Type | F | F | F | F | F | F |  |  |
| Default | none | none | none | none | none | none |  |  |

CSP Parameters Card. Include cards for each chemical species in the following format when CSP_SEL.GT.0. This input ends at the next keyword ("*") card.

| Card 3 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | AMPL | YCUT |  |  |  |  |  |  |
| Type | F | F |  |  |  |  |  |  |
| Default | none | none |  |  |  |  |  |  |

## VARIABLE

ID

## DESCRIPTION

Identifier for this 0D computation.

VARIABLE
COMPID Chemical composition identifier of composition to use.
SOLTYP Type of 0D calculation:
EQ.1: Isochoric
EQ.2: Isobaric
PLOTDT Simulation time interval for output both to the screen and to the isocom.csv file. This file can be loaded into LS-PREPOST for curve plotting using the $x-y$ plot facility.

CSP_SEL CSP solver option:
EQ.0: Do not use the CSP solver, and ignore the AMPL and YCUT parameters (default).
GT.0: Use the CSP solver, with the AMPL and YCUT parameters.

DT Initial time step
TLIMIT Time limit for the simulation
TIC Initial temperature
PIC Initial pressure
RIC Initial density
EIC Initial internal energy
AMPL Relative accuracy for the mass fraction of a chemical species in the Chemkin input file.

YCUT Absolute accuracy for the mass fraction of a chemical species in the Chemkin input file.

## Remarks:

1. If CSP_SEL.GT.0, then instead of using the full chemistry solver, the computational singular perturbation (CSP) method solver is used.

## *CHEMISTRY_CONTROL_1D

Purpose: Loads a previously-computed one-dimensional detonation. It is then available for use in the CESE solver for initializing a computation. In the product regions, this card overrides the initialization of the *CESE_INITIAL_CHEMISTRY_... cards.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | ID | XYZD | DETDIR | CSP_SEL |  |  |  |  |
| Type | I | F | 1 | 1 |  |  |  |  |
| Default | none | none | none | 0 |  |  |  |  |
| Remarks |  |  |  | 1 |  |  |  |  |

One-Dimensional Solution LSDA Input File Card.

| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | FILE |  |  |  |  |  |  |  |
| Type | A |  |  |  |  |  |  |  |

CSP Parameters Card Include cards for each chemical species in the following format when CSP_SEL > 0. This input ends at the next keyword ("*") card.

| Card 3 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | AMPL | YCUT |  |  |  |  |  |  |
| Type | F | F |  |  |  |  |  |  |
| Default | none | none |  |  |  |  |  |  |

VARIABLE
ID
XYZD

## DESCRIPTION

Identifier for this one-dimensional detonation solution.
Position of the detonation front in the DETDIR direction.
*CHEMISTRY_CONTROL_1D

VARIABLE
DETDIR

CSP_SEL

FILE
AMPL

YCUT

## DESCRIPTION

Detonation propagation direction
EQ.1: $x$
EQ.2: $y$
EQ.3: z
CSP solver option:
EQ.0: Do not use the CSP solver, and ignore the AMPL and YCUT parameters (default).
GT.0: Use the CSP solver, with the AMPL and YCUT parameters.

Name of the LSDA file containing the one-dimensional solution.
Relative accuracy for the mass fraction of a chemical species in the chemkin input file.

## Remarks:

1. If CSP_SEL $>0$, then instead of using the full chemistry solver, the computational singular perturbation (CSP) method solver is used.

## *CHEMISTRY_CONTROL_CSP

Purpose: Computes reduced chemistry for a specified Chemkin chemistry model using the Computational Singular Perturbation (CSP) method. This card can be used for gen-eral-purpose chemical reaction calculations.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | ID | IERROPT |  |  |  |  |  |  |
| Type | I | 1 |  |  |  |  |  |  |
| Default | none | none |  |  |  |  |  |  |

CSP Parameters Card. Include cards for each chemical species in the following format as indicated by the value of IERROPT. This input ends at the next keyword ("*") card.

| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | AMPL | YCUT |  |  |  |  |  |  |
| Type | F | F |  |  |  |  |  |  |
| Default | none | none |  |  |  |  |  |  |

## VARIABLE

ID
IERROPT

## DESCRIPTION

Identifier for this computational singular perturbation solver.
Selector:
EQ.0: AMPL and YCUT values for all chemical species are required.

EQ.1: One CSP Parameter Card should be provided, and it will be used for all species.

AMPL Relative accuracy for the mass fraction of a chemical species in the Chemkin input file.

YCUT

Absolute accuracy for the mass fraction of a chemical species in the Chemkin input file.

## *CHEMISTRY_CONTROL_FULL

Purpose: Computes the full chemistry specified by a Chemkin chemistry model. This card can be used for general-purpose chemical reaction calculations.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | ID | ERRLIM | RHOMIN | TMIN |  |  |  |  |
| Type | I | F | F | F |  |  |  |  |
| Default | none | none | 0.0 | 0.0 |  |  |  |  |

## VARIABLE

ID
ERRLIM
RHOMIN Minimum fluid density above which chemical reactions are computed.

TMIN Minimum temperature above which chemical reactions are computed.

## *CHEMISTRY_CONTROL_INFLATOR

Purpose: Provide the required properties of an inflator model for airbag inflation.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | MODEL | OUT_TYPE | TRUNTIM | DELT | PTIME |  |  |  |
| Type | I | I | F | F | F |  |  |  |
| Remarks | 1 | 2,4 |  |  |  |  |  |  |

Inflator Output Database File (an ASCII file) Card.

| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |
| Variable | FILE |  |  |  |  |  |  |  |
| Type | A |  |  |  |  |  |  |  |

Densities for Condensed Species. Include as many cards as needed. This input ends at the next keyword ("*") card.

| Card 3 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :--- | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | DENSITY |  | Species Name |  |  |  |  |  |
| Type | F |  | A |  |  |  |  |  |
| Default | none | none |  |  |  |  |  |  |
| Remark |  | 3 |  |  |  |  |  |  |

Name

VARIABLE
MODEL

OUT_TYPE Selects the output file format that will be used in an airbag simulation.

EQ.0: $\quad$ Screen output calibration output (see Remark 4)
EQ.1: CESE compressible flow solver (default)
EQ.2: ALE solver
EQ.3: $\quad$ CPM solver (with $2^{\text {nd }}$-order expansion of $C_{p}$ )
EQ.4: $\quad$ CPM solver (with $4^{\text {th }}$-order expansion of $C_{p}$ )
TRUNTIM Total run time.
DELT $\quad \operatorname{Delta}(\mathrm{t})$ to use in the model calculation.
PTIME Time interval for output of time history data to FILE.
FILE $\quad$ Name of the ASCII file in which to write the time history data and other data output by the inflator simulation.

DENSITY
Species

## DESCRIPTION

Type of inflator model to compute.
EQ.1: Pyrotechnic model
EQ.2: Hybrid model with cold flow option in the gas chamber

EQ.3: Hybrid model with heat flow in the gas chamber
EQ.4: $\quad$ Hybrid model with heat flow in one additional gas chamber

EQ.5: $\quad$ Hybrid model with heat flow in two additional gas chambers

Density of a condensed-phase species present in the inflator. Chemkin-compatible name of a condensed-phase species.

## Remarks:

1. If MODEL $=3$, the solution of an elementary reaction system is required for the finite-rate chemistry in the gas chamber. For MODEL $=4$ and 5 , the condensed phase is computed only in the combustion chamber.
2. Output file includes all of the necessary thermodynamics variables and load curves for the species mass flow rate, temperature, and density curve. This will make it possible to generate the velocity curve which is required by each solver that carries out an airbag simulation.
3. At least one of these cards will be input if condensed-phase species are present during the propellant combustion. In this case, the user must specify each con-densed-phase density. This density is then used to compute the volume fractions in both the combustion and gas chamber, where the energy equations are needed.
4. If OUT_TYPE $=0$, the propellant information will be displayed on the screen, including total mass, remaining mass percentage, and mass burning rate versus time, and the calibration data will be saved in the output file, including the time versus pressure, temperature, total mass flow rate, and individual species mass fractions for all chambers. With this option, the user can quickly see the effect of changing the parameters on the first three *CHEMISTRY_INFLATOR_PROPERTIES cards.

## *CHEMISTRY_CONTROL_TBX

Purpose: Specify a chemistry solver for use in conjunction with stochastic TBX particles. This is intended only for modeling the second phase of an explosion where the explosive has embedded metal (aluminum) particles that are too large to have burned in the first phase of the explosion.

This chemistry card points to a *CHEMISTRY_MODEL card (via IDCHEM) with its associated *CHEMISTRY_COMPOSITION cards to set up the initial conditions. That is, it establishes the spatial distribution of the species in the model.

It is assumed that there is no chemical reaction rate information in the chemistry model files. This is done since a special chemical reaction mechanism is implemented for TBX modeling. If particles other than solid aluminum particles are embedded in the explosive, then another burn model has to be implemented.

Surface Part Card. Card 1 format used when the PART keyword option is active.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | IDCHEM | USEPAR |  |  |  |  |  |  |
| Type | 1 | 1 |  |  |  |  |  |  |
| Default | none | 1 |  |  |  |  |  |  |

VARIABLE
IDCHEM
USEPAR Coupling flag indicating if a *STOCHASTIC_TBX_PARTICLES card is provided for this model:

EQ.1: uses a *STOCHASTIC_TBX_PARTICLES card (default).
EQ.O: does not use such a card.

## *CHEMISTRY_CONTROL_ZND

Purpose: Computes the one-dimensional reduced chemistry of a ZND model. It is then used in the initialization of the chemistry part of the CESE solver. When this card is used, the *CESE_INITIAL_CHEMISTRY... cards must specify the progressive variable (degree of combustion) in the HIC field.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | ID |  |  |  |  |  |  |  |
| Type | I |  |  |  |  |  |  |  |
| Default | none |  |  |  |  |  |  |  |


| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | F | EPLUS | Q0 | GAM | XYZD | DETDIR |  |  |
| Type | F | F | F | F | F | I |  |  |
| Default | none | none | none | none | none | none |  |  |

## VARIABLE

## DESCRIPTION

ID
Identifier for this full chemistry calculation.
F Overdriven factor
EPLUS EPLUS parameter of the ZND model.
Q0 Q0 parameter of the ZND model.
GAM GAM parameter of the ZND model.
XYZD Position of the detonation front in the DETDIR direction.

DETDIR Detonation propagation direction (1 => X; $2=>Y ; 3=>Z)$

## *CHEMISTRY_DET_INITIATION

Purpose: Performs a one-dimensional detonation calculation based upon a chemical composition and initial conditions. It is then available for use immediately in the CESE solver for initializing a computation, or it can be subsequently used by the *CHEMISTRY_CONTROL_1D card in a later run. In the product regions, this card overrides the initialization of the *CESE_INITIAL_CHEMISTRY... cards.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | ID | COMPID | NMESH | DLEN | CFL | TLIMIT | XYZD | DETDIR |
| Type | I | I | I | F | F | F | F | I |
| Default | none | none | none | none | none | none | none | none |

## LSDA Output File Card.

| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | FILE |  |  |  |  |  |  |  |
| Type |  | A |  |  |  |  |  |  |

## VARIABLE

## DESCRIPTION

ID
COMPID
Chemical composition identifier of composition to use.
NMESH Number of equal-width elements in the one-dimensional domain.
DLEN Length of the one-dimensional domain.
CFL Time-step limiting factor.
TLIMIT Time limit for the simulation
XYZD Position of the detonation front in the DETDIR direction.
DETDIR Detonation propagation direction ( $1=>X ; 2=>Y ; 3=>Z)$

VARIABLE
FILE

## DESCRIPTION

Name of the LSDA file in which to write the one-dimensional solution.

## *CHEMISTRY

## *CHEMISTRY_INFLATOR_PROPERTIES

Purpose: Provide the required properties of an inflator model.

| Card1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | COMP_ID | PDIA | PHEIGHT | PMASS | TOTMASS |  |  |  |
| Type | I | F | F | F | F |  |  |  |
| Remarks | 1 | 2 | 2 |  |  |  |  |  |


| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :--- | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | TFLAME | PINDEX | A0 | TDELAY | RISETIME |  |  |  |
| Type | F | F | F | F | F |  |  |  |
| Default | none | none | none | none | None |  |  |  |

Combustion Chamber Parameter Card.

| Card 3 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | COMP1ID | V0L1 | AREA1 | CD1 | P1 | T1 | DELP1 | DELTI |
| Type | I | F | F | F | $F$ | $F$ | $F$ | $F$ |
| Default | none | none | none | none | none | none | none | none |

Gas Plenum Parameter Card.

| Card 4 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :--- | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | COMP2ID | VOL2 | AREA2 | CD2 | P2 | T2 | DELP2 | DELT2 |
| Type | I | F | F | F | F | $F$ | $F$ | $F$ |
| Default | none | none | none | none | none | none | none | none |

Tank Parameter Card.

| Card 5 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :--- | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | COMP3ID | VOL3 | P3 | T3 |  |  |  |  |
| Type | I | F | F | F |  |  |  |  |
| Default | none | none | none | none |  |  |  |  |

Gas Chamber 1 (Optional, see Remark 3) Card.

| Card 6 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :--- | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | COMP4ID | VOL4 | AREA4 | CD4 | P4 | T4 | DELP4 | DELT4 |
| Type | I | F | F | F | F | $F$ | $F$ | $F$ |
| Default | none | none | none | none | none | none | none | none |

Gas Chamber 2 (Optional, see Remark 3) Card.

| Card 7 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | COMP5ID | V0L5 | AREA5 | CD5 | P5 | T5 | DELP5 | DELT5 |
| Type | I | F | F | F | F | $F$ | $F$ | $F$ |
| Default | none | none | none | none | none | none | none | none |

## VARIABLE

COMP_ID

PDIA Propellant diameter (see Remark 2).
PHEIGHT
PMASS
TOTMASS
TFLAME
PINDEX
A0
TDELAY
RISETIME
COMP1ID Chemical composition identifier of composition to use in the combustion chamber.

VOL1 Volume of the combustion chamber.
AREA1 Area of the combustion chamber.
CD1 Discharge coefficient of the combustion chamber.
P1 Pressure in the combustion chamber.

DELP1
DELT1

COMP2ID

VOL2
AREA2
CD2
P2
T2
DELP2
DELT2
COMP3ID
VOL3 Volume of the tank.
P3
T3
COMP4ID

VOL4
P4
T4
DELP4
DELT4

COMP5ID plenum.

Volume of the gas plenum.
Area of the gas plenum.

Pressure in the gas plenum.

Pressure in the tank.
Temperature in the tank. tional (second) gas chamber. ond gas chambers

## DESCRIPTION

Temperature in the combustion chamber.
Rupture pressure in the combustion chamber.
Elapsed time for breaking the burst disk between the chambers
Chemical composition identifier of composition to use in the gas

Discharge coefficient of the gas plenum.

Temperature in the gas plenum.
Rupture pressure in the gas plenum.
Elapsed time for breaking the burst disk between the chambers
Chemical composition identifier of composition to use in the tank.

Chemical composition identifier of composition to use in the addi-

Volume of the second gas chamber.
Pressure in the second gas chamber.
Temperature in the second gas chamber.
Rupture pressure in the second gas chamber.
Elapsed time for breaking the burst disk between the first and sec-

Chemical composition identifier of composition to use in the additional (third) gas chamber.

VOL5 Volume of the third gas chamber.
P5 Pressure in the third gas chamber.
T5 Temperature in the third gas chamber.
DELP5 Rupture pressure in the third gas chamber.
DELT5 Elapsed time for breaking the burst disk between the second and third gas chambers

## Remarks:

1. The propellant composition can be obtained by running a chemical equilibrium program such as NASA CEA, the CHEETAH code, or the PEP code. LSTC provides a modified version of the PEP code along with documentation for users; it is available upon request.
2. A spherical shape for the propellant particles can be chosen if an identical value for the diameter and height is given.
3. To simulate a 4 or 5 chamber inflator, an additional chamber card can be used. In these cases of the inflator models, the condensed phase species are limited to the combustion chamber only if involved in the propellant combustion.

## *CHEMISTRY_MODEL

Purpose: Identifies the files that define a Chemkin chemistry model.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | MODELID | JACSEL | ERRLIM |  |  |  |  |  |
| Type | I | 1 | F |  |  |  |  |  |
| Default | none | 1 | $1.0 \mathrm{e}-3$ |  |  |  |  |  |

Chemkin Input File Card.

| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |
| Variable | FILE1 |  |  |  |  |  |  |  |
| Type | A |  |  |  |  |  |  |  |

Thermodynamics Database File Card.

| Card 3 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |
| Variable | FILE2 |  |  |  |  |  |  |  |
| Type | A |  |  |  |  |  |  |  |

## Transport Properties Database File Card.

| Card 4 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |
| Variable | FILE3 |  |  |  |  |  |  |  |
| Type | A |  |  |  |  |  |  |  |

## VARIABLE

## DESCRIPTION

MODELID Identifier for this Chemkin-based chemistry model..

## VARIABLE

JACSEL

ERRLIM
FILE1

FILE2

FILE3

## DESCRIPTION

Selects the form of the Jacobian matrix for use in the source term.
EQ.1: Fully implicit (default)
EQ.2: Simplified implicit
Allowed error in element balance in a chemical reaction.
Name of the file containing the Chemkin-compatible input.
Name of the file containing the chemistry thermodynamics database.

Name of the file containing the chemistry transport properties database.

## *CHEMISTRY_PATH

Purpose: To specify one or more search paths to look for chemistry database files.
Include as many cards as needed. This input ends at the next keyword ("*") card.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | DIR |  |  |  |  |  |  |  |
| Type | A |  |  |  |  |  |  |  |

## VARIABLE

## DESCRIPTION

DIR
Directory path to add to the search set.

## *DUALCESE

The keyword *DUALCESE provides input data for the dual Conservation Element/Solution Element (dual CESE) compressible fluid solver:
*DUALCESE_BOUNDARY_AXISYMMETRIC_\{OPTION\}
*DUALCESE_BOUNDARY_CYCLIC_\{OPTION\}
*DUALCESE_BOUNDARY_FSI_\{OPTION\}
*DUALCESE_BOUNDARY_NON_REFLECTIVE_\{OPTION\}
*DUALCESE_BOUNDARY_PRESCRIBED_\{OPTION\}
*DUALCESE_BOUNDARY_PRESCRIBED_HYBRID_\{OPTION\}
*DUALCESE_BOUNDARY_PRESCRIBED_TWO-PHASE_\{OPTION\}
*DUALCESE_BOUNDARY_REFLECTIVE_\{OPTION\}
*DUALCESE_BOUNDARY_SLIDING_\{OPTION\}
*DUALCESE_BOUNDARY_SOLID_WALL_\{OPTION1\}_\{OPTION2\}
*DUALCESE_CONTROL_LIMITER
*DUALCESE_CONTROL_MESH_MOV
*DUALCESE_CONTROL_SOLVER (deprecated as of R14)
*DUALCESE_CONTROL_TIMESTEP
*DUALCESE_DATABASE_HISTORY_ELEMENT_SET
*DUALCESE_DATABASE_HISTORY_GLOBALS
*DUALCESE_DATABASE_HISTORY_NODE_SET
*DUALCESE_DATABASE_HISTORY_POINT_SET
*DUALCESE_DATABASE_HISTORY_SEGMENT_SET
*DUALCESE_D3PLOT
*DUALCESE_D3PLOT_FLUID_SSID
*DUALCESE_ELE2D

## *DUALCESE

*DUALCESE_ELE3D
*DUALCESE_ELEMENTSET
*DUALCESE_EOS_CAV_HOMOG_EQUILIB
*DUALCESE_EOS_COCHRAN_CHAN
*DUALCESE_EOS_COOLPROP
*DUALCESE_EOS_IDEAL_GAS
*DUALCESE_EOS_INFLATOR1
*DUALCESE_EOS_INFLATOR2
*DUALCESE_EOS_JWL
*DUALCESE_EOS_REFPROP
*DUALCESE_EOS_REFPROP_PATH
*DUALCESE_EOS_SET (deprecated as of R14)
*DUALCESE_EOS_STIFFENED_GAS
*DUALCESE_EOS_VAN_DER_WAALS_GENERALIZED
*DUALCESE_FSI_EXCLUDE
*DUALCESE_INCLUDE_MODEL
*DUALCESE_INITIAL
*DUALCESE_INITIAL_\{OPTION\}
*DUALCESE_INITIAL_HYBRID
*DUALCESE_INITIAL_HYBRID_SET
*DUALCESE_INITIAL_TWO-PHASE
*DUALCESE_INITIAL_TWO-PHASE_SET
*DUALCESE_MAT_GAS
*DUALCESE_MAT_GAS_0
*DUALCESE_MAT_GAS_2
*DUALCESE_MESH_GEOMETRY

*DUALCESE_MESH_PART<br>*DUALCESE_MODEL<br>*DUALCESE_NODE2D<br>*DUALCESE_NODE3D<br>*DUALCESE_NODESET<br>*DUALCESE_PART (deprecated as of R14)<br>*DUALCESE_PART_MULTIPHASE (deprecated as of R14)<br>*DUALCESE_POINTSET<br>*DUALCESE_REACTION_RATE_IG<br>*DUALCESE_REACTION_RATE_IG_REDUCED<br>*DUALCESE_REACTION_RATE_P_DEPEND<br>*DUALCESE_SEGMENTSET<br>*DUALCESE_SOLVER_SELECTION<br>*DUALCESE_SOLVER_CAV_EQNS<br>*DUALCESE_SOLVER_EULER_EQNS<br>*DUALCESE_SOLVER_HYBRID_MULTIPHASE<br>*DUALCESE_SOLVER_NAVIER_STOKES<br>*DUALCESE_SOLVER_TWO-PHASE_MULTIPHASE

An additional keyword option TITLE may be appended to the *DUALCESE keywords. If this option is used, then an addition line is read for the DUALCESE card in 80a format which can be used to describe that particular DUALCESE card. At present, the title serves no purpose other than to perhaps lend clarity to input decks.

## Dual CESE Keyword Deck Structure:

The structure of the keyword setup for the *DUALCESE solvers is different from the way most keyword input is handled in the LS-DYNA input phase. This stems from the fact that there can be several *DUALCESE models in the same problem. In order to deal with this, each such model is restricted to be specified with one file hierarchy that starts with the keyword file designated with the *DUALCESE_MODEL card.

## *DUALCESE

That keyword file can include any number of other keyword files with the *DUALCESE_INCLUDE_MODEL card, and each of those files can in turn include other keyword files, again with the *DUALCESE_INCLUDE_MODEL card. Standard *INCLUDE cards are not allowed. In fact, in each file in the file hierarchy of a *DUALCESE_MODEL card, only *DUALCESE cards may be used. The only exception to this is when the fluid mesh is defined using *MESH cards; a fatal error will be encountered when using other non-*DUALCESE keywords. Any required non-*DUALCESE keyword cards should be defined in some other place in the keyword input (outside the scope of the *DUALCESE_MODEL keyword card).

The mesh for each dual CESE model must be defined within the keyword input file hierarchy for that model. If a mesh created with ${ }^{*}$ MESH cards is used, the ${ }^{*}$ MESH cards for that mesh must all be defined within the scope of the *DUALCESE_MODEL card.

With R14, we introduce an updated, preferred way of specifying the dual CESE problem. The required or recommended keywords to use are:

```
*DUALCESE_MESH_GEOMETRY
*DUALCESE_MESH_PART
*DUALCESE_SOLVER_SELECTION
```

along with one of the following specific solver cards:
*DUALCESE_SOLVER_CAV_EQNS
*DUALCESE_SOLVER_EULER_EQNS
*DUALCESE_SOLVER_HYBRID_MULTIPHASE
*DUALCESE_SOLVER_NAVIER_STOKES
*DUALCESE_SOLVER_TWO-PHASE_MULTIPHASE
The following keywords are deprecated as of R14:
*DUALCESE_CONTROL_SOLVER
*DUALCESE_EOS_SET
*DUALCESE_PART
*DUALCESE_PART_MULTIPHASE
We also added time history cards with R14 that cause certain output to the binout file:
*DUALCESE_DATABASE_HISTORY_ELEMENT_SET

# *DUALCESE_DATABASE_HISTORY_GLOBALS <br> *DUALCESE_DATABASE_HISTORY_NODE_SET <br> *DUALCESE_DATABASE_HISTORY_POINT_SET <br> *DUALCESE_DATABASE_HISTORY_SEGMENT_SET 

We support two external equation-of-state (EOS) libraries, REFPROP and COOLPROP, for dual CESE. Including these libraries is complex, so we will clarify how to include them here. Each of them is accessed through a shared library that has to be loaded into LSDYNA at runtime with *MODULE_LOAD like in the follwing:

```
*MODULE_LOAD
UserA DUALCESE REFPROP
<path to the installed REFPROP shared library>
```

Note that this *MODULE_LOAD card must not be given inside a keyword file in the file hierarchy of a *DUALCESE_MODEL card. As discussed above, this is the case for all non-*DUALCESE keyword cards. Note also that since *MODULE is not available in the Windows version of LS-DYNA, this capability cannot be used in that version.

## REFPROP and COOLPROP Libraries:

While the REFPROP v10.0 version library and its directory of data sets is provided by ANSYS, COOLPROP libraries are not provided by ANSYS.

For the COOLPROP shared library, you can find the current production version here:
https:/ / sourceforge.net/projects/coolprop/files/CoolProp/6.3.0/shared_library/Linux/64bit/

## Multiphase Capabilities:

The multiphase capabilities added for R13 with modifications in R14 all have FSI capabilities, including with structural element failure. This includes the 'hybrid' multiphase, 'two-phase' multiphase, and the cavitation solvers. These keywords are involved with these capabilities:

*DUALCESE_BOUNDARY_PRESCRIBED_HYBRID<br>*DUALCESE_BOUNDARY_PRESCRIBED_TWO-PHASE<br>*DUALCESE_EOS_CAV_HOMOG_EQUILIB<br>*DUALCESE_EOS_COCHRAN_CHAN

## *DUALCESE

*DUALCESE_EOS_JWL
*DUALCESE_EOS_SET (deprecated as of R14; use one of the *DUALCESE_SOLVER_... cards instead)
*DUALCESE_EOS_STIFFENED_GAS
*DUALCESE_EOS_VAN_DER_WAALS_GENERALIZED
*DUALCESE_INITIAL_HYBRID
*DUALCESE_INITIAL_HYBRID_SET
*DUALCESE_INITIAL_TWO-PHASE
*DUALCESE_INITIAL_TWO-PHASE_SET
*DUALCESE_PART_MULTIPHASE (deprecated as of R14; use *DUALCESE_MESH_PART instead)
*DUALCESE_REACTION_RATE_IG
*DUALCESE_REACTION_RATE_IG_REDUCED
*DUALCESE_REACTION_RATE_P_DEPEND
*DUALCESE_SOLVER_HYBRID_MULTIPHASE (added in R14)
*DUALCESE_SOLVER_TWO-PHASE_MULTIPHASE (added in R14)

## Comparison to CESE:

The capabilities implemented in the dual CESE solvers are only a part of what is available in the *CESE solvers, especially those that involve couplings with the *CHEMISTRY and *STOCHASTIC_PARTICLE solvers. We plan to port some of those capabilities to the *DUALCESE solvers as well.

## *DUALCESE_BOUNDARY_AXISYMMETRIC_OPTION

Available options are:
MSURF
SEGMENT_SET
Purpose: Define an axisymmetric boundary condition on the axisymmetric axis for the 2D axisymmetric dual CESE compressible flow solver.

The MSURF option should be used when the dual CESE mesh has been created using *MESH cards. The SEGMENT_SET option should be used when *DUALCESE_ELE2D or *DUALCESE_ELE3D cards specify the dual CESE mesh.

## Card Summary:

Card 1a. This card is included for the MSURF keyword option. Provide as many cards as necessary. This input ends at the next keyword ("*") card.

| MSPID |  |  |  |  |  |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

Card 1b. This card is included for the SEGMENT_SET keyword option. Provide as many cards as necessary. This input ends at the next keyword ("*") card.

| SSID |  |  |  |  |  |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

## Data Card Definitions:

Surface Part Card. Card 1 used when the MSURF keyword option is active. Include as many cards as necessary. This input ends at the next keyword ("*") card.

| Card 1a | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | MSPID |  |  |  |  |  |  |  |
| Type | 1 |  |  |  |  |  |  |  |
| Default | none |  |  |  |  |  |  |  |

## VARIABLE

MSPID Mesh surface part ID that is referenced by *MESH_SURFACE_ELEMENT cards

Set Card. Card 1 used when the SEGMENT_SET keyword option is active. Include as many cards as necessary. This input ends at the next keyword ("*") card.

| Card 1b | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | SSID |  |  |  |  |  |  |  |
| Type | 1 |  |  |  |  |  |  |  |
| Default | none |  |  |  |  |  |  |  |

VARIABLE
SSID

## DESCRIPTION

Segment set ID for the segment set created with *DUALCESE_SEGMENTSET

## Remarks:

This boundary condition can only be used on the axisymmetric axis for the 2D axisymmetric dual CESE fluid solver.

## *DUALCESE_BOUNDARY_CYCLIC_OPTION

Available options are:

## MSURF

SEGMENT_SET
Purpose: Define a cyclic (periodic) boundary condition for dual CESE compressible flows. This cyclic boundary condition can be used on periodic boundary surfaces.

The MSURF option should be used when the dual CESE mesh has been created using *MESH cards. The SEGMENT_SET option should be used when *DUALCESE_ELE2D or *DUALCESE_ELE3D cards specify the dual CESE mesh.

## Card Summary:

Card Sets. The following sequence of cards comprises a single set. LS-DYNA will continue reading these cards sets until the next keyword ("*") card is encountered.
Card 1a. This card is included if the MSURF keyword option is used.

| MSPID1 | MSPID2 | CYCTYP |  |  |  |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

Card 1b. This card is included if the SEGMENT_SET keyword option is used.

| SSID1 | SSID2 | CYCTYP |  |  |  |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

Card 2a. This card is included when CYCTYP $=1$.

| AXISX1 | AXISY1 | AXISZ1 | DIRX | DIRY | DIRZ | ROTANG |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

Card $\mathbf{2 b}$. This card is included when CYCTYP $=2$.

| TRANSX | TRANSY | TRANSZ |  |  |  |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

## Data Card Definitions:

Surface Part Card. Card 1 format used when the MSURF keyword option is active.

| Card 1a | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | MSPID1 | MSPID2 | CYCTYP |  |  |  |  |  |
| Type | 1 | 1 | 1 |  |  |  |  |  |
| Default | none | none | 0 |  |  |  |  |  |
| Remarks |  |  | 1,2 |  |  |  |  |  |

## VARIABLE

MSPID1,
MSPID2
CYCTYP Relationship between the two cyclic boundary condition surfaces:

EQ.O: Relationship determined by LS-DYNA (default)
EQ.1: The first surface is rotated about an axis to match the second surface.

EQ.2: The faces of the first surface are translated in a given direction to obtain the corresponding faces on the second surface.

Set Card. Card 1 format used when the SEGMENT_SET keyword option is active.

| Card 1b | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | SSID1 | SSID2 | CYCTYP |  |  |  |  |  |
| Type | 1 | 1 | 1 |  |  |  |  |  |
| Default | none | none | 0 |  |  |  |  |  |
| Remarks |  |  | 1,3 |  |  |  |  |  |

## VARIABLE

SSID1, SSID2

CYCTYP

## DESCRIPTION

Segment set IDs for the segment sets created with *DUALCESE_SEGMENTSET

Relationship between the two cyclic boundary condition surfaces:

EQ.O: Relationship determined by LS-DYNA (default)
EQ.1: The first surface is rotated about an axis to match the second surface.

EQ.2: The faces of the first surface are translated in a given direction to obtain the corresponding faces on the second surface.

Rotation Case Card. Additional card when CYCTYP $=1$.

| Card 2a | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | AXISX1 | AXISY1 | AXISZ1 | DIRX | DIRY | DIRZ | ROTANG |  |
| Type | F | F | F | F | F | $F$ | $F$ |  |
| Default | 0.0 | 0.0 | 0.0 | none | none | none | none |  |

## VARIABLE

AXIS[X,Y,Z]1
$\operatorname{DIR}[X, Y, Z] \quad$ The direction which together with AXIS[X,Y,Z]1 defines the axis of rotation for the transformation between the surfaces

ROTANG The angle of rotation (in degrees) that transforms the centroid of each face on the first surface to the centroid of the corresponding face on the second surface

Translation Case Card. Additional card when CYCTYP $=2$.

| Card 2b | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | TRANSX | TRANSY | TRANSZ |  |  |  |  |  |
| Type | F | F | F |  |  |  |  |  |
| Default | none | none | none |  |  |  |  |  |

VARIABLE
TRANS[X,Y,Z]

## DESCRIPTION

The translation direction that enables the identification of the segment in the second surface that matches a segment in the first surface

## Remarks:

1. Unspecified Relationship between Surfaces. For the case CYCTYP $=0$, LSDYNA examines the geometry of two faces of the two surfaces in order to determine if the surfaces are approximately parallel $($ CYCTYP $=2)$ or related through a rotation $(C Y C T Y P=1)$. The geometric parameters required are then computed.
2. MSURF. For the MSURF option, each mesh surface part must contain the same number of mesh surface elements. The mesh surface elements in each mesh surface part are internally ordered for pairwise matching between the two mesh surface parts.
3. SEGMENT_SET. For the SEGMENT_SET option, each segment set must contain the same number segments. The segments in each set are internally ordered for pairwise matching between the two sets.

## *DUALCESE_BOUNDARY_FSI_OPTION

Available options are:
MSURF
SEGMENT_SET
Purpose: Define an FSI boundary condition for the moving mesh dual CESE compressible flow solver. This keyword must not be combined with the dual CESE immersed-boundary method FSI solver in the same dual CESE part on the same dual CESE mesh. Doing so will result in an error termination condition.

This boundary condition must be applied on a surface of the dual CESE computational domain that is co-located with surfaces of the outside boundary of the structural mesh. The nodes of the two meshes will generally not be shared.

The MSURF option should be used when the dual CESE mesh has been created using *MESH cards. The SEGMENT_SET option should be used when *DUALCESE_ELE2D or *DUALCESE_ELE3D cards specify the dual CESE mesh.

## Card Summary:

Card 1a. This card is included for the MSURF keyword option. Provide as many cards as necessary. This input ends at the next keyword ("*") card.

| MSPID | REF_P |  |  |  |  |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

Card 1b. This card is included for the SEGMENT_SET keyword option. Provide as many cards as necessary. This input ends at the next keyword ("*") card.

| SSID | REF_P |  |  |  |  |  |
| :---: | :--- | :--- | :--- | :--- | :--- | :--- |

## Data Card Definitions:

Surface Part Card. Card 1 used when the MSURF keyword option is active. Include as many cards as necessary. This input ends at the next keyword ("*") card.

| Card 1a | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | MSPID | REF_P |  |  |  |  |  |  |
| Type | I | F |  |  |  |  |  |  |
| Default | none | 0.0 |  |  |  |  |  |  |

## VARIABLE

MSPID

## DESCRIPTION

Mesh surface part ID that is referenced by *MESH_SURFACE_ELEMENT cards

REF_P Ambient/reference pressure of the fluid domain on the side opposite this structural interface to the fluid simulation domain. This ambient pressure only needs to be specified in the case where the FSI structural part(s) connected with this FSI interface are not immersed in the dual CESE mesh. This reference pressure defaults to 0.0 since moving mesh FSI calculations most often involve structures surrounded by the dual CESE mesh, and there is no need for a reference pressure in that case.

Set Card. Card 1 used when the SEGMENT_SET keyword option is active. Include as many cards as necessary. This input ends at the next keyword ("*") card.

| Card 1b | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | SSID | REF_P |  |  |  |  |  |  |
| Type | I | F |  |  |  |  |  |  |
| Default | none | 0.0 |  |  |  |  |  |  |

## VARIABLE

SSID

## DESCRIPTION

Segment set ID for the segment set created with *DUALCESE_SEGMENTSET

VARIABLE
REF_P

## DESCRIPTION

Ambient/reference pressure of the fluid domain on the side opposite this structural interface to the fluid simulation domain. This ambient pressure only needs to be specified in the case where the FSI structural part(s) connected with this FSI interface are not immersed in the dual CESE mesh. This reference pressure defaults to 0.0 since moving mesh FSI calculations most often involve structures surrounded by the dual CESE mesh, and there is no need for a reference pressure in that case..

## Remarks:

This boundary condition card is also needed for conjugate heat transfer problems with the moving mesh dual CESE solver. But the conjugate heat transfer capability is not yet implemented in the dual CESE solver.

## *DUALCESE_BOUNDARY_NON_REFLECTIVE_OPTION

Available options are:
MSURF
SEGMENT_SET
Purpose: Define a passive boundary condition for dual CESE compressible flows. This non-reflective boundary condition provides an artificial computational boundary for an open boundary that is passive.

The MSURF option should be used when the dual CESE mesh has been created using *MESH cards. The SEGMENT_SET option should be used when *DUALCESE_ELE2D or *DUALCESE_ELE3D cards are used to specify the dual CESE mesh.

## Card Summary:

Card 1a. This card is included when the MSURF keyword option is used. Include as many cards as necessary. This input ends at the next keyword ("*") card.

| MSPID | DIRX | DIRY | DIRZ |  |  |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

Card 1b. This card is included when the SEGMENT_SET keyword option is used. Include as many cards as necessary. This input ends at the next keyword ("*") card.

| SSID | DIRX | DIRY | DIRZ |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |

## Data Card Definitions:

Surface Part Card. Card 1 used when the MSURF keyword option is active. Include as many cards as necessary. This input ends at the next keyword ("*") card.

| Card 1a | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | MSPID | DIRX | DIRY | DIRZ |  |  |  |  |
| Type | I | F | F | F |  |  |  |  |
| Default | none | 0.0 | 0.0 | 0.0 |  |  |  |  |

## VARIABLE

MSPID

## DESCRIPTION

Mesh surface part ID that is referenced by *MESH_SURFACE_ELEMENT cards

DIRX, DIRY, If this vector is nonzero, then it is used as the prescribed flow diDIRZ rection.

Set Card. Card 1 used when the SET keyword option is active. Include as many cards as necessary. This input ends at the next keyword ("*") card.

| Card 1b | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | SSID | DIRX | DIRY | DIRZ |  |  |  |  |
| Type | I | F | F | F |  |  |  |  |
| Default | none | 0.0 | 0.0 | 0.0 |  |  |  |  |

## VARIABLE

SSID

DIRX, DIRY, DIRZ

## DESCRIPTION

Segment set ID for the segment set created with *DUALCESE_SEGMENTSET

If this vector is nonzero, then it is used as the prescribed flow direction.

## Remarks:

1. Boundary Surface Flow. This boundary condition is usually imposed on an open surface that is far from the main disturbed flow (the further away, the better), meaning the flow on the boundary surface should be almost uniform.
2. Default Boundary Condition. If any boundary segment has not been assigned a boundary condition by any of the *DUALCESE_BOUNDARY $\qquad$ cards, then it will automatically be assigned this non-reflective boundary condition.

## *DUALCESE_BOUNDARY_PRESCRIBED_OPTION

Available options include:

## MSURF

SEGMENT_SET
Purpose: For the dual CESE compressible flow solver, set boundary values for velocity, density, pressure and temperature. Boundary values are applied at the centroid of elements connected with this boundary.

The MSURF option should be used when the dual CESE mesh has been created using *MESH cards. The SEGMENT_SET card should be used when *DUALCESE_ELE2D or *DUALCESE_ELE3D cards specify the dual CESE mesh.

## Card Sets:

A set of data cards for this keyword consists of 3 of the following cards:

1. Card 1 specifies the object to which the boundary condition is applied. Its format depends on the keyword option.
2. Card 2 reads in load curve IDs.
3. Card 3 reads in scale factors.

For each boundary condition to be specified include one set of cards. This input ends at the next keyword ("*") card.

## Card Summary:

Card 1a. This card is included if the MSURF keyword option is used.

| MSPID | IDCOMP | DIRX | DIRY | DIRZ |  |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

Card 1b. This card is included if the SEGMENT_SET keyword option is used.

| SSID | IDCOMP | DIRX | DIRY | DIRZ |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |

Card 2. This card is required.

| LC_U | LC_V | LC_W | LC_RH0 | LC_P | LC_T |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

Card 3. This card is required.

| SF_U | SF_V | SF_W | SF_RHO | SF_P | SF_T |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

## Data Card Definitions:

Surface Part Card. Card 1 format used when the MSURF keyword option is active.

| Card 1a | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | MSPID | IDCOMP | DIRX | DIRY | DIRZ |  |  |  |
| Type | I | I | F | F | F |  |  |  |
| Default | none | none | 0.0 | 0.0 | 0.0 |  |  |  |

## VARIABLE

MSPID

IDCOMP

DIRX, DIRY, DIRZ

## DESCRIPTION

Mesh surface part ID that is referenced by *MESH_SURFACE_ELEMENT cards

For inflow boundaries in problems involving chemical reacting flows, the chemical mixture of the fluid entering the domain is defined with a *CHEMISTRY_COMPOSITION card with this ID [Not yet available].

If this vector is non-zero, then it is used as the prescribed flow direction.

Set Card. Card 1 format used when the SEGMENT_SET keyword option is active.

| Card 1b | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | SSID | IDCOMP | DIRX | DIRY | DIRZ |  |  |  |
| Type | I | I | F | F | F |  |  |  |
| Default | none | none | 0.0 | 0.0 | 0.0 |  |  |  |

## VARIABLE

SSID
IDCOMP

DIRX, DIRY, If this vector is non-zero, then it is used as the prescribed flow diDIRZ

## DESCRIPTION

ID for the segment set created with *DUALCESE_SEGMENTSET
For inflow boundaries in problems involving chemical reacting flows, the chemical mixture of the fluid entering the domain is defined with a *CHEMISTRY_COMPOSITION card with this ID [Not yet available]. rection.

## Load Curve Card.

| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | LC_U | LC_V | LC_W | LC_RHO | LC_P | LC_T |  |  |
| Type | I | 1 | 1 | 1 | 1 | 1 |  |  |
| Remarks | 1 | 1 | 1 | 1 | 1 | 1 |  |  |

VARIABLE
LC_U

LC_V

LC_W

Load curve ID to describe the $z$-component of the velocity as a function of time or function ID to give the $z$-component of the

## VARIABLE

LC_RHO

LC_P

LC_T Load curve ID to describe the temperature as a function of time or function ID to give the temperature as a function of position, velocity, temperature, pressure, and time, $f(x, y, z, v x, v y, v z$, temp, pres, time).

EQ.O: Temperature is a constant with value SF_T.
EQ.-1: Temperature is computed by the solver.

## Scale Factor Card.

| Card 3 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | SF_U | SF_V | SF_W | SF_RHO | SF_P | SF_T |  |  |
| Type | F | F | F | F | F | F |  |  |
| Default | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 |  |  |

Scale factor for LC_U

## VARIABLE

## DESCRIPTION

SF_V Scale factor for LC_V
SF_W Scale factor for LC_W
SF_RHO Scale factor for LC_RHO
SF_P Scale factor for LC_P
SF_T Scale factor for LC_T

## Remarks:

1. Consistent Boundary Values. On each centroid or set of centroids, the variables $\left(v_{x}, v_{y}, v_{z}, \rho, P, T\right)$ that are given values must be consistent and make the model well-posed (meaning be such that the solution of the model exists, is unique, and is physical).

## *DUALCESE_BOUNDARY_PRESCRIBED_HYBRID_OPTION

MSURF

## SEGMENT_SET

Purpose: For the dual CESE compressible flow solver, set boundary values for velocity, density, pressure, temperature, and other values in the hybrid multiphase model. Boundary values are applied at the centroid of elements connected with this boundary.

The MSURF option should be used when the dual CESE mesh has been created using *MESH cards. The SEGMENT_SET card should be used when *DUALCESE_ELE2D or *DUALCESE_ELE3D cards specify the dual CESE mesh.

## Card Sets:

A set of data cards for this keyword consists of 3 of the following cards:

1. Card 1 specifies the object to which the boundary condition is applied. Its format depends on the keyword option.
2. Cards 2 and 3 provide load curve IDs.
3. Cards 4 and 5 provide scale factors.

For each boundary condition to be specified include one set of cards. This input ends at the next keyword ("*") card.

## Card Summary:

Card 1a. This card is included if the keyword option is set to MSURF.

| MSPID | IDCOMP | DIRX | DIRY | DIRZ |  |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

Card 1b. This card is included if the keyword option is set to SSID.

| SSID | IDCOMP | DIRX | DIRY | DIRZ |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |

Card 2. This card is required.

| LC_Z1 | LC_RA | LC_U | LC_V | LC_W | LC_D1 | LC_DA | LC_DB |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |

Card 3. This card is required.

| LC_P | LC_T |  |  |  |  |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

Card 4. This card is required.

| SF_Z1 | SF_RA | SF_U | SF_V | SF_W | SF_D1 | SF_DA | SF_DB |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |

Card 5. This card is required.

| SF_P | SF_T $_{-}$ |  |  |  |  |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

## Data Card Definitions:

Surface Part Set Card. Card 1 format used when the MSURF keyword option is active.

| Card 1a | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | MSPID | IDCOMP | DIRX | DIRY | DIRZ |  |  |  |
| Type | I | I | F | F | F |  |  |  |
| Default | none | none | 0.0 | 0.0 | 0.0 |  |  |  |

## VARIABLE

MSPID

IDCOMP

DIRX, DIRY, DIRZ

## DESCRIPTION

Mesh surface part ID that is referenced by *MESH_SURFACE_ELEMENT cards

For inflow boundaries in problems involving chemical reacting flows, the chemical mixture of the fluid entering the domain is defined with a *CHEMISTRY_COMPOSITION card with this ID [Not yet available].

If this vector is nonzero, then it is used as the prescribed flow direction.

Segment Set Card. Card 1 format used when the SEGMENT_SET keyword option is active.

| Card 1b | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | SSID | IDCOMP | DIRX | DIRY | DIRZ |  |  |  |
| Type | I | 1 | F | F | F |  |  |  |
| Default | none | none | 0.0 | 0.0 | 0.0 |  |  |  |

## VARIABLE

SSID
IDCOMP

## DESCRIPTION

ID for the segment set created with *DUALCESE_SEGMENTSET
For inflow boundaries in problems involving chemical reacting flows, the chemical mixture of the fluid entering the domain is defined with a *CHEMISTRY_COMPOSITION card with this ID [Not yet available].

DIRX, DIRY, If this vector is nonzero, then it is used as the prescribed flow diDIRZ rection.

Load Curve Card. See Remark 1.

| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | LC_Z1 | LC_RA | LC_U | LC_V | LC_W | LC_D1 | LC_DA | LC_DB |
| Type | I | I | I | I | I | I |  |  |

## VARIABLE

LC_Z1

## DESCRIPTION

Load curve ID or function ID to describe the volume fraction of material 1 as a function of time or a function of position, velocity, temperature, pressure, and time, $f(x, y, z, v x, v y, v z$, temp, pres, time), respectively.

EQ.O: The volume fraction is a constant with value SF_Z1.
EQ.-1: The volume fraction is computed by the solver.
LC_RA Load curve or function ID to describe the mass fraction of reactant (material $\alpha$ ) with respect to the explosive mixture (material 2 ) as a
function of time or a function of position, velocity, temperature, pressure, and time, $f(x, y, z, v x, v y, v z$, temp, pres, time), respectively.

EQ.0: The mass fraction is a constant with value SF_RA.
EQ.-1: The mass fraction is computed by the solver.
LC_U Load curve or defined function ID to describe the $x$-component of the velocity as a function of time or a function of position, velocity, temperature, pressure, and time, $f(x, y, z, v x, v y, v z$, temp, pres, time), respectively.

EQ.0: The $x$-component of velocity is a constant with value SF_- $^{-}$ U.

EQ.-1: The $x$-component of velocity is computed by the solver.
LC_V Load curve or defined function ID to describe the $y$-component of the velocity as a function of time or a function of position, velocity, temperature, pressure, and time, $f(x, y, z, v x, v y, v z$, temp, pres, time), respectively.

EQ.0: The $y$-component of velocity is a constant with value SF_- $^{-}$ V.

EQ.-1: The $y$-component of velocity is computed by the solver.
LC_W Load curve or defined function ID to describe the $z$-component of the velocity as a function of time or a function of position, velocity, temperature, pressure, and time, $f(x, y, z, v x, v y, v z$, temp, pres, time), respectively.

EQ.0: The $z$-component of velocity is a constant with value SF_- $^{-}$ W.

EQ.-1: The $z$-component of velocity is computed by the solver.
LC_D1 Load curve or defined function ID to describe the density of the first multiphase material as a function of time or a function of position, velocity, temperature, pressure, and time, $f(x, y, z, v x, v y, v z$, temp, pres, time), respectively.

EQ.O: The density of the first multiphase material is a constant with value SF_D1.
EQ.-1: The density of the first multiphase material is computed by the solver.

## VARIABLE

LC_DA

LC_DB

## DESCRIPTION

Load curve or defined function ID to describe the density of the reactant (material $\alpha$ ) as a function of time or a function of position, velocity, temperature, pressure, and time, $f(x, y, z, v x, v y, v z$, temp, pres, time), respectively.

EQ.0: The density of the reactant is a constant with value SF_- $^{-}$ DA.

EQ.-1: The density of the reactant is computed by the solver.
Load curve or defined function ID to describe the density of the product (material $\beta$ ) as a function of time or a function of position, velocity, temperature, pressure, and time, $f(x, y, z, v x, v y, v z$, temp, pres, time), respectively.

EQ.0: The density of the product is a constant with value SF_- $_{-}$ DB.
EQ.-1: The density of the product is computed by the solver.

Load Curve Card 2. See Remark 1.

| Card 3 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | LC_P | LC_T |  |  |  |  |  |  |
| Type | 1 | 1 |  |  |  |  |  |  |

## VARIABLE

LC_P

LC_T Load curve or defined function ID to describe the temperature as a function of time or a function of position, velocity, temperature, pressure, and time, $f(x, y, z, v x, v y, v z$, temp, pres, time $)$, respectively.

EQ.O: The temperature is a constant with value SF_T.

## DESCRIPTION

EQ.-1: The temperature is computed by the solver.
Scale Factor Card.

| Card 4 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | SF_Z1 | SF_RA | SF_U | SF_V | SF_W | SF_D1 | SF_DA | SF_DB |
| Type | F | F | F | F | F | F | F | F |
| Default | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 |

## VARIABLE

## DESCRIPTION

SF_Z
SF_RA Scale factor for LC_RA
SF_U Scale factor for LC_U
SF_V Scale factor for LC_V
SF_W Scale factor for LC_W
SF_D1 Scale factor for LC_D1
SF_DA Scale factor for LC_DA
SF_DB Scale factor for LC_DB
Scale Factor Card 2.

| Card 5 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | SF_P $^{\prime}$ | SF_T $^{\prime}$ |  |  |  |  |  |  |
| Type | F | F |  |  |  |  |  |  |
| Default | 1.0 | 1.0 |  |  |  |  |  |  |

## VARIABLE

SF_P Scale factor for LC_P
SF_T Scale factor for LC_T

## Remark:

1. Consistent Boundary Values. On each centroid or set of element centroids, the variables ( $v_{x}, v_{y}, v_{z}, \rho, P, T, \ldots$ ) that are given values must be consistent and make the model well-posed (meaning be such that the solution of the model exists, is unique, and is physical).

## *DUALCESE_BOUNDARY_PRESCRIBED_TWO-PHASE_OPTION

Available options include:

## MSURF

SEGMENT_SET
Purpose: For the dual CESE compressible flow solver, set boundary values for velocity, density, pressure, temperature, and other values in the two-phase multiphase model. Boundary values are applied at the centroid of elements connected with this boundary. OPTION = SEGMENT_SET is for user defined meshes whereas OPTION = MSURF is associated with the automatic volume mesher (See *MESH keywords).

That is, the MSURF option is used when the dual CESE mesh has been created using *MESH cards. The SEGMENT_SET card is used when *DUALCESE_ELE2D or *DUALCESE_ELE3D cards are used to define the dual CESE mesh.

## Card Sets:

A set of data cards for this keyword consists of 3 of the following cards:

1. Card 1 specifies the object to which the boundary condition is applied. Its format depends on the keyword option.
2. Card 2 provides load curve IDs.
3. Card 3 provides scale factors.

For each boundary condition to be specified include one set of cards. This input ends at the next keyword ("*") card.

## Card Summary:

Card 1a. This card is included if the keyword option is set to MSURF.

| MSPID | IDCOMP | DIRX | DIRY | DIRZ |  |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

Card 1 b . This card is included if the keyword option is set to SSID.

| SSID | IDCOMP | DIRX | DIRY | DIRZ |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |

Card 2. This card is required.

| LC_Z1 | LC_U | LC_V | LC_W | LC_D1 | LC_D2 | LC_P | LC_T |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

Card 3. This card is required.

| SF_Z1 | SF_U | SF_V | SF_W | SF_D1 | SF_D2 | SF_P | SF_T |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |

## Data Card Definitions:

Surface Part Set Card. Card 1 format used when the MSURF keyword option is active.

| Card 1a | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | MSPID | IDCOMP | DIRX | DIRY | DIRZ |  |  |  |
| Type | I | I | F | F | F |  |  |  |
| Default | none | none | 0.0 | 0.0 | 0.0 |  |  |  |

## VARIABLE

MSPID

IDCOMP

DIRX, DIRY,
DIRZ

## DESCRIPTION

Mesh surface part ID that is referenced by *MESH_SURFACE_ELEMENT cards

For inflow boundaries in problems involving chemical reacting flows, the chemical mixture of the fluid entering the domain is defined with a *CHEMISTRY_COMPOSITION card with this ID [Not yet available].

If this vector is nonzero, then it is used as the prescribed flow direction.

Segment Set Card. Card 1 format used when the SEGMENT_SET keyword option is active.

| Card 1b | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | SSID | IDCOMP | DIRX | DIRY | DIRZ |  |  |  |
| Type | I | I | F | F | F |  |  |  |
| Default | none | none | 0.0 | 0.0 | 0.0 |  |  |  |

## VARIABLE

SSID
IDCOMP

## DESCRIPTION

ID for the segment set created with *DUALCESE_SEGMENTSET
For inflow boundaries in problems involving chemical reacting flows, the chemical mixture of the fluid entering the domain is defined with a *CHEMISTRY_COMPOSITION card with this ID [Not yet available].

DIRX, DIRY, If this vector is nonzero, then it is used as the prescribed flow diDIRZ rection.

Load Curve Card. See Remark 1.

| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | LC_Z1 | LC_U | LC_V | LC_W | LC_D1 | LC_D2 | LC_P | LC_T |
| Type | I | I | I | I | । | I | । | । |

## VARIABLE

LC_Z1

## DESCRIPTION

Load curve or defined function ID to describe the volume fraction of material 1 as a function of time or a function of position, velocity, temperature, pressure, and time, $f(x, y, z, v x, v y, v z$, temp, pres, time), respectively.

EQ.0: The volume fraction is a constant with value SF_Z1.
EQ.-1: The volume fraction is computed by the solver.
LC_U Load curve or defined function ID to describe the $x$-component of the velocity as a function of time or a function of position, velocity, temperature, pressure, and time, $f(x, y, z, v x, v y, v z$, temp, pres, time), respectively.

EQ.0: The $x$-component of velocity is a constant with value SF_- $_{-}$ U.

EQ.-1: The $x$-component of velocity is computed by the solver.
LC_V Load curve or defined function ID to describe the $y$-component of the velocity as a function of time or a function of position, velocity, temperature, pressure, and time, $f(x, y, z, v x, v y, v z$, temp, pres, time), respectively.

VARIABLE

LC_W Load curve or defined function ID to describe the $z$-component of the velocity as a function of time or a function of position, velocity, temperature, pressure, and time, $f(x, y, z, v x, v y, v z$, temp, pres, time), respectively.

EQ.0: The $z$-component of velocity is a constant with value SF_- $^{-}$ W.

EQ.-1: The $z$-component of velocity is computed by the solver.
Load curve or defined function ID to describe the density of material 1 as a function of time or a function of position, velocity, temperature, pressure, and time, $f(x, y, z, v x, v y, v z$, temp, pres, time), respectively.

EQ.0: The density is a constant with value SF_D1.
EQ.-1: The density is computed by the solver.
Load curve or defined function ID to describe the density of material 2 as a function of time or a function of position, velocity, temperature, pressure, and time, $f(x, y, z, v x, v y, v z$, temp, pres, time), respectively.

EQ.0: The density is a constant with value SF_D2.
EQ.-1: The density is computed by the solver.
LC_P Load curve or defined function ID to describe the pressure as a function of time or a function of position, velocity, temperature, pressure, and time, $f(x, y, z, v x, v y, v z$, temp, pres, time), respectively.

EQ.O: The pressure is a constant with value SF_P.
EQ.-1: The pressure is computed by the solver.
LC_T Load curve or defined function ID to describe the temperature as a function of time or a function of position, velocity, temperature, pressure, and time, $f(x, y, z, v x, v y, v z$, temp, pres, time $)$, respectively.

EQ.0: The temperature is a constant with value SF_T.

EQ.-1: The temperature is computed by the solver.
Scale Factor Card.

| Card 3 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | SF_Z1 | SF_U | SF_V | SF_W | SF_D1 | SF_D2 | SF_P | SF_T |
| Type | F | F | F | F | F | F | F | F |
| Default | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 | 1.0 |

## VARIABLE

## DESCRIPTION

SF_Z
SF_U Scale factor for LC_U
SF_V Scale factor for LC_V
SF_W Scale factor for LC_W
SF_D1 Scale factor for LC_D1
SF_D2 Scale factor for LC_D2
SF_P Scale factor for LC_P
SF_T Scale factor for LC_T

## Remark:

1. Consistent Boundary Values. On each centroid or set of element centroids, the variables ( $v_{x}, v_{y}, v_{z}, \rho, P, T, \ldots$ ) that are given values must be consistent and make the model well-posed (meaning be such that the solution of the model exists, is unique, and is physical).

## *DUALCESE_BOUNDARY_PRESCRIBED_VN_OPTION

Available options include:
MSURF
SEGMENT_SET
Purpose: For the dual CESE compressible flow solver, set boundary values for the normal velocity, density, pressure and temperature. Boundary values are applied at the centroid of elements connected with this boundary. This boundary condition differs from *DUALCESE_BOUNDARY_PRESCRIBED_VN in that the normal velocity is prescribed instead of each velocity component.

The MSURF option should be used when the dual CESE mesh has been created using *MESH cards. The SEGMENT_SET option should be used when *DUALCESE_ELE2D or *DUALCESE_ELE3D cards specify the dual CESE mesh.

## Card Sets:

A set of data cards for this keyword consists of 3 of the following cards:

1. Card 1 specifies the object to which the boundary condition is applied. Its format depends on the keyword option.
2. Card 2 reads in load curve IDs.
3. Card 3 reads in scale factors.

For each boundary condition to be specified include one set of cards. This input ends at the next keyword ("*") card.

## Card Summary:

Card 1a. This card is included if the MSURF keyword option is used.

| MSPID | IDCOMP |  |  |  |  |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

Card 1b. This card is included if the SEGMENT_SET keyword option is used.

| SSID | IDCOMP |  |  |  |  |  |  |
| :---: | :---: | :--- | :--- | :--- | :--- | :--- | :--- |

Card 2. This card is required.

| LC_VN |  |  | LC_RHO | LC_P | LC_T |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

Card 3. This card is required.

| SF_VN |  |  | SF_RH0 | SF_P | SF_T |  |  |
| :---: | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

## Data Card Definitions:

Surface Part Card. Card 1 format used when the MSURF keyword option is active.

| Card 1a | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | MSPID | IDCOMP |  |  |  |  |  |  |
| Type | 1 | 1 |  |  |  |  |  |  |
| Default | none | none |  |  |  |  |  |  |

## VARIABLE

MSPID

IDCOMP For inflow boundaries in problems involving chemical reacting flows, the chemical mixture of the fluid entering the domain is defined with a *CHEMISTRY_COMPOSITION card with this ID (not yet available).

Set Card. Card 1 format used when the SEGMENT_SET keyword option is active.

| Card 1b | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | SSID | IDCOMP |  |  |  |  |  |  |
| Type | I | I |  |  |  |  |  |  |
| Default | none | none |  |  |  |  |  |  |

## VARIABLE <br> SSID

## DESCRIPTION

ID for the segment set created with *DUALCESE_SEGMENTSET

VARIABLE
IDCOMP

## DESCRIPTION

For inflow boundaries in problems involving chemical reacting flows, the chemical mixture of the fluid entering the domain is defined with a *CHEMISTRY_COMPOSITION card with this ID.

Load Curve Card. See Remark 1.

| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | LC_VN |  |  | LC_RH0 | LC_P | LC_T |  |  |
| Type | I |  |  | 1 | 1 | 1 |  |  |

## VARIABLE

LC_VN

LC_RHO

LC_P

LC_T Load curve ID to describe the temperature as a function of time or a function of position, velocity, temperature, pressure, and time, $f(x, y, z, v x, v y, v z$, temp, pres, time), respectively.

EQ.O: The temperature is a constant with value SF_T.
EQ.-1: The temperature is computed by the solver.

Scale Factor Card.

| Card 3 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | SF_VN |  |  | SF_RH0 | SF_P | SF_T |  |  |
| Type | F |  |  | F | F | F |  |  |
| Default | 1.0 |  |  | 1.0 | 1.0 | 1.0 |  |  |

## VARIABLE

SF_VN
Scale factor for LC_VN
SF_RHO
Scale factor for LC_RHO
SF_P Scale factor for LC_P
SF_T Scale factor for LC_T

## Remarks:

1. Consistent Boundary Values. On each centroid or set of centroids, the variables ( $V_{N}, \rho, P, T$ ) that are given values must be consistent and make the model well-posed (meaning be such that the solution of the model exists, is unique, and is physical).

## *DUALCESE_BOUNDARY_REFLECTIVE_OPTION

Available options are:
MSURF
SEGMENT_SET
Purpose: Define a reflective boundary condition for the dual CESE compressible flow solver. This boundary condition can be applied on a symmetrical surface or a solid wall of the computational domain.

The MSURF option should be used when the dual CESE mesh has been created using *MESH cards. The SEGMENT_SET option should be used when *DUALCESE_ELE2D or *DUALCESE_ELE3D cards specify the dual CESE mesh.

## Card Summary:

Card 1a. This card is included for the MSURF keyword option. Provide as many cards as necessary. This input ends at the next keyword ("*") card.

| MSPID |  |  |  |  |  |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

Card 1b. This card is included for the SEGMENT_SET keyword option. Provide as many cards as necessary. This input ends at the next keyword ("*") card.

| SSID |  |  |  |  |  |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

## Data Card Definitions:

Surface Part Card. Card 1 format used when the MSURF keyword option is active. Provide as many cards as necessary. This input ends at the next keyword ("*") card.

| Card 1a | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | MSPID |  |  |  |  |  |  |  |
| Type | 1 |  |  |  |  |  |  |  |
| Default | none |  |  |  |  |  |  |  |

## VARIABLE

MSPID Mesh surface part ID that is referenced by *MESH_SURFACE_ELEMENT cards

## DESCRIPTION

Set Card. Card 1 format used when the SEGMENT_SET keyword option is active. Provide as many cards as necessary. This input ends at the next keyword ("*") card.

| Card 1b | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | SSID |  |  |  |  |  |  |  |
| Type | 1 |  |  |  |  |  |  |  |
| Default | none |  |  |  |  |  |  |  |

## VARIABLE

SSID

## DESCRIPTION

Segment set ID for the segment set created with *DUALCESE_SEGMENTSET

## Remarks:

This boundary condition has the same effect as a solid wall boundary condition for inviscid flows.

## *DUALCESE_BOUNDARY_SOLID_WALL_OPTION1_\{OPTION2\}

For OPTION1 the choices are:
MSURF
SEGMENT_SET
For OPTION2 the choices are:

```
<BLANK>
```

ROTATE
Purpose: Define a solid wall boundary condition for the dual CESE compressible flow solver. This boundary condition can be applied at a solid boundary that is the physical boundary for the flow field. For inviscid flow, this will be a slip boundary condition; while for viscous flows, it is a no-slip boundary condition.

The MSURF option should be used when the dual CESE mesh has been created using *MESH cards. The SEGMENT_SET option should be used when *DUALCESE_ELE2D or *DUALCESE_ELE3D cards specify the dual CESE mesh. The ROTATE keyword option allows the boundary condition to rotate around an axis with a variable speed given by a load curve.

## Card Summary:

Card Sets. The following sequence of cards comprises a single set. LS-DYNA will continue reading data card sets until the next keyword ("*") card is encountered.

Card 1a. This card is included for the MSURF keyword option without the ROTATE keyword option.

| MSPID | LCID | VX | VY | VZ |  |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

Card 1b. This card is included for the MSURF keyword option with the ROTATE keyword option.

| MSPID | LCID | XP | YP | ZP | NX | NY | NZ |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

Card 1c. This card is included for the SEGMENT_SET keyword option without the ROTATE keyword option.

| SSID | LCID | VX | VY | VZ |  |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

Card 1d. This card is included for the SEGMENT_SET keyword option with the ROTATE keyword option.

| SSID | LCID | XP | YP | ZP | NX | NY | NZ |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |

## Data Card Definitions:

Surface Part Card without Rotation. Card 1 format used when the MSURF keyword option is active without the ROTATE keyword option

| Card 1a | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | MSPID | LCID | VX | VY | VZ |  |  |  |
| Type | I | I | F | F | F |  |  |  |
| Default | none | 0 | 0.0 | 0.0 | 0.0 |  |  |  |
| Remarks |  | 2 | 2 | 2 | 2 |  |  |  |

## VARIABLE

MSPID

LCID

VX, VY, VZ Velocity vector of the solid wall boundary condition:
LCID.EQ.0: Constant velocity vector specified with VX, VY, and VZ.
LCID.NE.0: VX, VY, and VZ give the velocity vector that is scaled by LCID.

Surface Part Card with Rotation. Card 1 format used when the MSURF keyword option is active with the ROTATE keyword option.

| Card 1b | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | MSPID | LCID | XP | YP | ZP | NX | NY | NZ |
| Type | I | I | F | F | $F$ | $F$ | $F$ | $F$ |
| Default | none | 0 | 0.0 | 0.0 | 0.0 | 0.0 | 0.0 | 0.0 |

## VARIABLE

MSPID

LCID

## DESCRIPTION

Mesh surface part ID that is referenced by *MESH_SURFACE_ELEMENT cards

Load curve ID for specifying the rotating speed frequency in Hz . This input is required.

XP, YP, ZP Coordinates for a point on the axis of rotation
NX, NY, NZ Unit vector for specifying the direction of the axis of rotation. This is not used for the 2D case.

Segment Set Card without Rotation. Card 1 format used when the SEGMENT_SET keyword option is active without the ROTATE keyword option

| Card 1c | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | SSID | LCID | VX | VY | VZ |  |  |  |
| Type | I | I | F | F | F |  |  |  |
| Default | none | 0 | 0.0 | 0.0 | 0.0 |  |  |  |
| Remarks |  | 2 | 2 | 2 | 2 |  |  |  |

## VARIABLE

## DESCRIPTION

ID of the segment set created with *DUALCESE_SEGMENTSET
$\frac{\text { VARIABLE }}{\text { LCID }}$

VX, VY, VZ Velocity vector of the solid wall boundary condition:
LCID.EQ.0: Constant velocity vector specified with VX, VY, and VZ.

LCID.NE.0: VX, VY, and VZ give the velocity vector that is scaled by LCID.

## DESCRIPTION

Load curve ID scales the velocity vector specified with (VX, VY, VZ) to give the solid wall boundary movement. If not defined, the solid wall boundary moves with a constant velocity vector specified by (VX, VY, VZ).

Segment Set Card with Rotation. Card 1 format used when the SEGMENT_SET keyword option is active with the ROTATE keyword option.

| Card 1d | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | SSID | LCID | XP | YP | ZP | NX | NY | NZ |
| Type | I | I | F | F | F | F | F | F |
| Default | none | 0 | 0.0 | 0.0 | 0.0 | 0.0 | 0.0 | 0.0 |

## VARIABLE

SSID
LCID

XP, YP, ZP
NX, NY, NZ
or moving mesh solver should be used. Also, this moving boundary condition only affects viscous flows (no-slip boundary condition).
2. Fixed Solid Wall Boundary Condition. If LCID $=0$ and $V x=V y=V z=0.0$ (default), this will be a regular solid wall boundary condition.

## *DUALCESE_CONTROL_LIMITER

Purpose: Sets some stability parameters used in the dual CESE compressible flow solver on the current dual CESE model.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | IDLMT | ALPHA | BETA | EPSR |  |  |  |  |
| Type | I | F | F | F |  |  |  |  |
| Default | 0 | 0.0 | 0.0 | 0.0 |  |  |  |  |
| Remarks |  | 1 | 2 | 3 |  |  |  |  |

## VARIABLE

IDLMT

ALPHA

BETA Numerical viscosity control coefficient, $\beta$ (see dual CESE theory manual). $0 \leq \beta \leq 1$.

EPSR Stability control coefficient, $\varepsilon$ (see dual CESE theory manual). Must be $\geq 0$.

## Remarks:

1. Re-weighting Coefficient. Larger values of $\alpha$ give more stability, but less accuracy. Usually $\alpha=2.0$ or 4.0 will be enough for normal shock problems.
2. Numerical Viscosity Control Coefficient. Larger values of $\beta$ give more stability. For problems with shock waves, $\beta=1.0$ is recommended.
3. Stability Control Coefficient. Larger values of $\varepsilon$ give more stability, but less accuracy.

## *DUALCESE_CONTROL_MESH_MOV

Purpose: Specify the algorithm for calculating the mesh movement (morphing) of a given DUALCESE part in an FSI problem. This keyword is for the moving mesh version of dual CESE.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | ID | IALG | NINTER | RELERR | MXDISPR |  |  |  |
| Type | I | 1 | 1 | F | F |  |  |  |
| Default | none | 9 | 100 | $10^{-3}$ | $10^{-2}$ |  |  |  |

## VARIABLE

ID
IALG Mesh motion algorithm:
EQ.9: IDW scheme (default)
NITER Number of linear solver iterations (when using a linear solver specified in IALG). No linear solvers have been implemented at this time, so this field is ignored.

RELERR

MXDISPR

## DESCRIPTION

ID for this mesh motion algorithm

Relative error for determining convergence when using a linear solver specified in IALG. No linear solvers have been implemented at this time, so this field is ignored.

Maximum displacement relative to element size to use as a crite- rion for avoiding the full calculation of the motion of the DUALCESE part on a given time step. If the full calculation can be avoided, the elements touching an FSI interface are still morphed, but it is assumed that this approximation will not lead to elements that are overly distorted.

## *DUALCESE_CONTROL_SOLVER

Purpose: Set general purpose control variables for the dual CESE compressible flow solver.

WARNING: As of version R14, this keyword is deprecated. Instead, in order to select the appropriate equations to solve, you should use *DUALCESE_SOLVER_SELECTION along with the corresponding *DUALCESE_SOLVER_... card. Also, instead of this card, use *DUALCESE_MESH_GEOM for the geometry-related details.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | EQNS | IGEOM | IFRAME | MIXTYPE | IDC | ISNAN |  |  |
| Type | A | A | A | A | F | 1 |  |  |
| Default | EULER | none | FIXED | optional | 0.25 | 0 |  |  |
| Remarks |  | 1 |  |  | 2 |  |  |  |

## VARIABLE

EQNS

IGEOM
Sets the geometric dimension:
EQ.2D: Two-dimensional (2D) problem
EQ.3D: Three-dimensional (3D) problem
EQ.AXI: 2D axisymmetric
IFRAME Choose the frame of reference:
EQ.FIXED: Usual non-moving reference frame (default).
EQ.ROT: Non-inertial rotating reference frame. $\quad$ IFRAME $=$ ROTATING may also be used.

## VARIABLE

MIXTYPE

IDC

ISNAN

## DESCRIPTION

Select the mix or multiphase model solver (if any):
EQ.<blank>: No mix or multiphase model (default)
EQ.HYBRID: Hybrid multiphase model solver
EQ.TWO-PHASE: Two-phase multiphase solver
Contact interaction detection coefficient (for FSI and conjugate heat transfer problems)

Flag to check for NaN in the dual CESE solver solution arrays at the completion of each time step. This option can be useful for debugging purposes. There is a cost overhead when this option is active.

EQ.0: No checking.
EQ.1: Checking is active.

## Remarks:

1. Mesh and Boundary Conditions for 2D Problems. If you want to use the 2D (IGEOM $=2 \mathrm{D}$ ) or 2D axisymmetric (IGEOM=AXI) solver, the mesh should only be distributed in the $x y$-plane with the boundary conditions given only at the $x y$ domain boundaries. Otherwise, a warning message will be given, and the 3D solver will be triggered instead.

The 2D axisymmetric case will work only if the 2D mesh and corresponding boundary conditions are properly defined with the $x$ and $y$ coordinates corresponding to the radial and axial directions, respectively.
2. Contact Interaction Detection Coefficient. IDC is the same type of variable that is input on the *ICFD_CONTROL_FSI card. For an explanation, see Remark 1 for the *ICFD_CONTROL_FSI card.
3. Dual CESE Solver and Restarts. The dual CESE solver is not currently supported for restarts.

## *DUALCESE_CONTROL_TIMESTEP

Purpose: Sets the time step control parameters for the CESE compressible flow solver.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | IDDT | CFL | DTINT |  |  |  |  |  |
| Type | I | F | F |  |  |  |  |  |
| Default | 0 | 0.9 | $10^{-3}$ |  |  |  |  |  |

## VARIABLE

IDDT

## DESCRIPTION

Sets the time step option:
EQ.0: Fixed time step size of DTINT, the given initial time step size

NE.O: The time step size will be calculated based on the given CFL-number and the flow solution at the previous time step.

CFL $\quad$ CFL number (Courant-Friedrichs-Lewy condition). $0.0<\mathrm{CFL} \leq$ 1.0

DTINT Initial time step size

## *DUALCESE_DATABASE_HISTORY_ELEMENT_SET

Purpose: Enable output of dual CESE solver data at the element centroids of the selected dual CESE mesh elements. This element output goes to binary database binout. At most, one instance of this keyword may appear in the input deck.

Output Options Card.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | ESID | DT | LCUR | IOOPT |  |  |  |  |
| Type | I | F | 1 | 1 |  |  |  |  |
| Default | 0 | 0. | 0 | 0 |  |  |  |  |

## VARIABLE

ESID
DT

LCUR Optional curve ID specifying the time interval between outputs. Use *DEFINE_CURVE to define the curve. The abscissa is time, and the ordinate is time interval between outputs.

IOOPT Flag to govern behavior of the output frequency load curve defined by LCUR:

EQ.1: When output is generated at time $t_{n}$, the next output time $t_{n+1}$ is computed as

$$
t_{n+1}=t_{n}+\operatorname{LCUR}\left(t_{n}\right)
$$

This is the default behavior.
EQ.2: When output is generated at time $t_{n}$, the next output time $t_{n+1}$ is computed as

$$
t_{n+1}=t_{n}+\operatorname{LCUR}\left(t_{n+1}\right) .
$$

EQ.3: Output is generated for each abscissa point in the load curve definition. The actual value of the load curve is ignored.

## *DUALCESE_DATABASE_HISTORY_GLOBALS

Purpose: Enable output of global dual CESE solver data. The output goes to binary database binout. At most, one instance of this keyword may appear in the input deck.

## Output Options Card.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | DT | LCUR | IO0PT |  |  |  |  |  |
| Type | F | 1 | 1 |  |  |  |  |  |
| Default | 0.0 | 0 | 0 |  |  |  |  |  |

## VARIABLE

DT

LCUR

IOOPT

## DESCRIPTION

Time interval between outputs. If DT is zero, no output is generated.

Optional curve ID specifying the time interval between outputs. Use *DEFINE_CURVE to define the curve; the abscissa is time, and the ordinate is time interval between outputs.

Flag to govern behavior of the output frequency load curve defined by LCUR:

EQ.1: When output is generated at time $t_{n}$, the next output time $t_{n+1}$ is computed as

$$
t_{n+1}=t_{n}+\operatorname{LCUR}\left(t_{n}\right)
$$

This is the default behavior.
EQ.2: When output is generated at time $t_{n}$, the next output time $t_{n+1}$ is computed as

$$
t_{n+1}=t_{n}+\operatorname{LCUR}\left(t_{n+1}\right)
$$

EQ.3: Output is generated for each abscissa point in the load curve definition. The actual value of the load curve is ignored.

## *DUALCESE_DATABASE_HISTORY_NODE_SET

Purpose: Enable output of dual CESE solver data at the selected dual CESE mesh nodes. This nodal output goes to binary database binout. At most, one instance of this keyword may appear in the input deck.

Output Options Card.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | NSID | DT | LCUR | IOOPT |  |  |  |  |
| Type | I | F | 1 | 1 |  |  |  |  |
| Default | 0 | 0. | 0 | 0. |  |  |  |  |

## VARIABLE

NSID
DT

LCUR Optional curve ID specifying the time interval between outputs. Use *DEFINE_CURVE to define the curve; the abscissa is time, and the ordinate is time interval between outputs.

IOOPT Flag to govern behavior of the output frequency load curve defined by LCUR:

EQ.1: When output is generated at time $t_{n}$, the next output time $t_{n+1}$ is computed as

$$
t_{n+1}=t_{n}+\operatorname{LCUR}\left(t_{n}\right)
$$

This is the default behavior.
EQ.2: When output is generated at time $t_{n}$, the next output time $t_{n+1}$ is computed as

$$
t_{n+1}=t_{n}+\operatorname{LCUR}\left(t_{n+1}\right) .
$$

EQ.3: Output is generated for each abscissa point in the load curve definition. The actual value of the load curve is ignored.

## *DUALCESE_DATABASE_HISTORY_POINT_SET

Purpose: Enable output of dual CESE solver data at the selected points inside the dual CESE mesh. This point-based output goes to binary database binout. At most, one instance of this keyword may appear in the input deck.

## Output Options Card.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | PNTSID | DT | LCUR | IOOPT |  |  |  |  |
| Type | I | F | 1 | 1 |  |  |  |  |
| Default | 0 | 0. | 0 | 0. |  |  |  |  |

## VARIABLE

PNTSID
DT Time interval between outputs. If DT is zero, no output is generated.

LCUR Optional curve ID specifying the time interval between outputs. Use *DEFINE_CURVE to define the curve; the abscissa is time, and the ordinate is time interval between outputs.

IOOPT Flag to govern behavior of the output frequency load curve defined by LCUR:

EQ.1: When output is generated at time $t_{n}$, the next output time $t_{n+1}$ is computed as

$$
t_{n+1}=t_{n}+\operatorname{LCUR}\left(t_{n}\right)
$$

This is the default behavior.
EQ.2: When output is generated at time $t_{n}$, the next output time $t_{n+1}$ is computed as

$$
t_{n+1}=t_{n}+\operatorname{LCUR}\left(t_{n+1}\right)
$$

EQ.3: Output is generated for each abscissa point in the load curve definition. The actual value of the load curve is ignored.

## *DUALCESE_DATABASE_HISTORY_SEGMENT_SET

Purpose: Enable output of dual CESE solver data averaged on each of the specified dual CESE segment sets. This segment set-based output goes to binary database binout. At most, one instance of this keyword may appear in the input deck.

## Output Options Card.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | DT | LCUR | IOOPT |  |  |  |  |  |
| Type | F | 1 | 1 |  |  |  |  |  |
| Default | 0.0 | 0 | 0 |  |  |  |  |  |

Segment Sets Card. Define as many cards as necessary. Input ends at the next keyword ("*") card.

| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | SSID1 | SSID2 | SSID3 | SSID4 | SSID5 | SSID6 | SSID7 | SSID8 |
| Type | 1 | 1 | 1 | 1 | 1 | 1 | 1 | 1 |
| Default | none | none | none | none | none | none | none | none |

## VARIABLE

DT

LCUR Optional curve ID specifying the time interval between outputs. Use *DEFINE_CURVE to define the curve; the abscissa is time, and the ordinate is time interval between outputs.

IOOPT Flag to govern behavior of the output frequency load curve defined by LCUR:

## DESCRIPTION

EQ.1: When output is generated at time $t_{n}$, the next output time $t_{n+1}$ is computed as

$$
t_{n+1}=t_{n}+\operatorname{LCUR}\left(t_{n}\right)
$$

This is the default behavior.
EQ.2: When output is generated at time $t_{n}$, the next output time $t_{n+1}$ is computed as

$$
t_{n+1}=t_{n}+\operatorname{LCUR}\left(t_{n+1}\right)
$$

EQ.3: Output is generated for each abscissa point in the load curve definition. The actual value of the load curve is ignored.

SSID $i \quad i^{\text {th }}$ dual CESE segment set ID (see *DUALCESE_SEGMENTSET). For each of these segment sets, an average value of each dual CESE output variable is output to the binout file at the times selected by the fields in Card 1.

## *DUALCESE_D3PLOT

Purpose: Specify the flow variables to be added to the dual CESE d3plot output.
Provide as many cards as necessary. This input ends at the next keyword ("*") card.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | FLOW_VAR |  |  |  |  |  |  |  |
| Type | A |  |  |  |  |  |  |  |

## VARIABLE

FLOW_VAR

## DESCRIPTION

Name of a flow variable to output to the d3plot file. The currently supported variables are listed in the table below.

Flow Variables:
This table lists the supported flow variables.

| FLOW_VAR |  | DESCRIPTION |
| :---: | :--- | :--- | :--- |
| DENSITY |  | Density |
| VELOCITY |  | Velocity |
| MOMENTUM |  | Momentum |
| VORTICITY |  | Vorticity |
| TOTAL_ENERGY |  | Total energy |
| INTERNAL_ENERGY |  | Internal energy |
| PRESSURE |  | Pressure |
| TEMPERATURE |  | Temperature |
| MACH_NUMBER |  | Flow Mach number |
| SCHLIEREN_NUMBER |  | Quantity for capturing or highlighting the shock |
|  |  | structure in a compressible flow |

## FLOW_VAR

VOID_FRACTION

VOLUME_FRACTION

REACTANT_MASS_FRACTION

## DESCRIPTION

Void fraction (only for the dual CESE cavitation solver)

Volume fraction of the different materials in a hybrid or two-phase multiphase model

Mass fraction of the reactant (material $\alpha$ ) with respect to the explosive material (material 2) in a hybrid multiphase model

## *DUALCESE_D3PLOT_FLUID_SSID

Purpose: Generate surface d3plot output for the dual CESE solver on a specified dual CESE mesh segment set. These surfaces may be on the outside of the dual CESE fluid mesh that is in contact with the structural volume element parts.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | SSID |  |  |  |  |  |  |  |
| Type |  |  |  |  |  |  |  |  |

## VARIABLE

## DESCRIPTION

SSID
ID of a segment set created with *DUALCESE_SEGMENTSET

Dual CESE variables to output. Provide as many cards as necessary. This input ends at the next keyword ("*") card.

| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | FLOW_VAR |  |  |  |  |  |  |  |
| Type | A |  |  |  |  |  |  |  |

## VARIABLE

FLOW_VAR

## DESCRIPTION

Name of a flow variable to output to the d3plot file. The currently supported variables are listed in the table below.

## Flow Variables:

This table lists the supported flow variables.

| FLOW_VAR |  |  |
| :---: | :---: | :--- | :--- |
| DENSITY |  | Density |
| VELOCITY |  | Velocity |
| MOMENTUM |  | Momentum |

FLOW_VAR
VORTICITY
TOTAL_ENERGY
INTERNAL_ENERGY
PRESSURE
TEMPERATURE
MACH_NUMBER
SCHLIEREN_NUMBER

VOID_FRACTION

VOLUME_FRACTION

REACTANT_MASS_FRACTION

## DESCRIPTION

Vorticity
Total energy
Internal energy
Pressure
Temperature
Flow Mach number
Quantity for capturing or highlighting the shock structure in a compressible flow

Void fraction (only for the dual CESE cavitation solver)

Volume fraction of the different materials in a hybrid or two-phase multiphase model

Mass fraction of the reactant (material $\alpha$ ) with respect to the explosive material (material 2) in a hybrid multiphase model

## *DUALCESE_ELE2D

Purpose: Define three and four node elements.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 | 9 | 10 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | EID | PID | N1 | N2 | N3 | N4 |  |  |  |  |
| Type | I | I | I | I | I | I |  |  |  |  |
| Default | none | none | none | none | none | none |  |  |  |  |

## VARIABLE

EID

PID

N1
N2

N3

N4

DESCRIPTION
Element ID. Choose a unique number with respect to other elements.

Part ID, see *DUALCESE_MESH_PART.
Nodal point 1
Nodal point 2
Nodal point 3
Nodal point 4

## *DUALCESE_ELE3D

Purpose: Define three-dimensional fluid volume elements. These can be 4 node tetrahedra, 5 node pyramids, 6 node wedges (prisms), and 8 node hexahedra.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 | 9 | 10 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | EID | PID | N1 | N2 | N3 | N4 | N5 | N6 | N7 | N8 |
| Type | 1 | 1 | 1 | 1 | 1 | 1 | 1 | 1 | 1 | 1 |
| Default | none | none | none | none | none | none | none | none | none | none |
| Remarks | 1 |  |  |  |  |  |  |  |  |  |
| VARIABLE |  | DESCRIPTION |  |  |  |  |  |  |  |  |
| EID |  | Element ID. A unique number must be chosen. |  |  |  |  |  |  |  |  |
| PID |  | Part ID, see *DUALCESE_MESH_PART. |  |  |  |  |  |  |  |  |
| N1 |  | Nodal point 1 |  |  |  |  |  |  |  |  |
| N2 |  | Nodal point 2 |  |  |  |  |  |  |  |  |
| N3 |  | Nodal point 3 |  |  |  |  |  |  |  |  |
| $\vdots \quad \vdots$ |  |  |  |  |  |  |  |  |  |  |
| N8 |  | Nodal point 8 |  |  |  |  |  |  |  |  |

## Remarks:

1. Node Numbering. Four, five, six, and eight node elements are allowed as numbered below. This ordering must be followed, or code termination will occur during the initialization phase with a negative volume message. In the case of a pyramid element, the base of the pyramid must follow the ordering used for the hexahedron. See *ELEMENT_SOLID for a figure showing the positions of the nodes in 4,6 , and 8 node elements.

| 4-noded tetrahedron | $\mathrm{N} 1, \mathrm{~N} 2, \mathrm{~N} 3, \mathrm{~N} 4, \mathrm{~N} 4, \mathrm{~N} 4, \mathrm{~N} 4, \mathrm{~N} 4$ |
| :--- | :--- |
| $\underline{\text { 5-noded pyramid }}$ | $\mathrm{N} 1, \mathrm{~N} 2, \mathrm{~N} 3, \mathrm{~N} 4, \mathrm{~N} 5, \mathrm{~N} 5, \mathrm{~N} 5, \mathrm{~N} 5$ |


| 6-noded pentahedron | N1, N2, N3, N4, N5, N5, N6, N6 |
| :--- | :--- |
| 8-noded hexahedron | N1, N2, N3, N4, N5, N6, N7, N8 |

## *DUALCESE_ELEMENTSET

Purpose: Define a set of dual CESE mesh elements.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | ESID |  |  |  |  |  |  |  |
| Type | 1 |  |  |  |  |  |  |  |
| Default | none |  |  |  |  |  |  |  |

Element ID Cards. List of elements in the set, where the element IDs are defined with *DUALCESE_ELE2D or *DUALCESE_ELE3D cards. Include as many cards as needed. This input ends at the next keyword ("*") card.

| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | EID1 | EID2 | EID3 | EID4 | EID5 | EID6 | EID7 | EID8 |
| Type | 1 | 1 | 1 | 1 | 1 | 1 | 1 | 1 |

## VARIABLE

ESID

EID $i$

## DESCRIPTION

Set ID. All dual CESE element sets should have a unique set ID.
Element ID $i$

## *DUALCESE_EOS_CAV_HOMOG_EQUILIB

Purpose: Define the coefficients in the equation of state (EOS) for the homogeneous equilibrium cavitation model.

Include as many cards as needed. This input ends at the next keyword ("*") card.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | EOSID | RHOVAP | RHOLIQ | AVAP | ALIQ | MUVAP | MULIQ | PSATVAP |
| Type | I | F | F | F | F | F | F | F |
| Default | none | 0.8 | 880.0 | 334.0 | 1386.0 | $1.435 e-5$ | $1.586 \mathrm{e}-4$ | $1.2 \mathrm{e}+4$ |


| VARIABLE | DESCRIPTION |
| :---: | :--- |
| EOSID |  |
| RHOVAP | Density of the saturated vapor, $\rho_{\mathrm{vap}}$ |
| RHOLIQ | Density of the saturated liquid, $\rho_{\mathrm{liq}}$ |
| AVAP | Sound speed of the saturated vapor, $a_{\mathrm{vap}}$ |
| ALIQ | Sound speed of the saturated liquid, $a_{\mathrm{liq}}$ |
| MUVAP | Dynamic viscosity of the vapor, $\mu_{\mathrm{vap}}$ |
| MULIQ | Dynamic viscosity of the liquid, $\mu_{\mathrm{liq}}$ |
| PSATVAP | Pressure of the saturated vapor, $P_{\text {SatVap }}$ |

## Remarks:

1. EOS Validity. In this homogeneous equilibrium cavitation model, a barotropic equation of state is used. This model can be used in small scale and high-speed cavitation flows, but it is not good for large-scale, low-speed cavitation calculations.

## *DUALCESE_EOS_COCHRAN_CHAN

Purpose: Define a Cochran-Chan type of EOS that provides a means to represent a condensed phase explosive in a dual CESE multiphase model.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | EOSID | A | B | EPS1 | EPS2 | GAMMAO | RH00 | E0 |
| Type | I | F | F | F | F | F | F | F |
| Default | none | none | none | none | none | none | none | 0.0 |


| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | CV |  |  |  |  |  |  |  |
| Type | F |  |  |  |  |  |  |  |
| Default | none |  |  |  |  |  |  |  |

## VARIABLE

EOSID
A

B
EPS1
EPS2
GAMMA0
RHOO
E0 Represents the heat of detonation released during the reactions, or the constant rate of afterburn energy added ( $\mathrm{E} 0=0.0$ is the default), $e_{0}$

CV Heat capacity, $C_{v}$

## Remarks:

The Cochran-Chan EOS like the JWL EOS (*DUALCESE_EOS_JWL) is a type of MieGruneisen EOS. The equations of state of a Mie-Gruneisen form are given by:

$$
P(\rho, e)=P_{\mathrm{ref}}+\Gamma(\rho) \rho\left[e-e_{\mathrm{ref}}(\rho)\right]
$$

Here $\Gamma(\rho)$ is the Gruneisen coefficient. For the Cochran-Chan EOS reference pressure and energy are given by:

$$
\begin{aligned}
P_{r e f}(\rho) & =A\left(\frac{\rho_{0}}{\rho}\right)^{-\varepsilon_{1}}-B\left(\frac{\rho_{0}}{\rho}\right)^{-\varepsilon_{2}} \\
e_{\text {ref }}(\rho) & =\frac{A}{\rho_{0}\left(1-\varepsilon_{1}\right)}\left(\frac{\rho_{0}}{\rho}\right)^{1-\varepsilon_{1}}+\frac{B}{\rho_{0}\left(1-\varepsilon_{2}\right)}\left(\frac{\rho_{0}}{\rho}\right)^{1-\varepsilon_{2}}-e_{0}
\end{aligned}
$$

## *DUALCESE_EOS_COOLPROP

Purpose: Define an equation of state (EOS) to be evaluated using the COOLPROP EOS library

Note that the COOLPROP library is not provided by ANSYS. You need to download a 64-bit version of the shared library from a public repository, such as:
https:/ / sourceforge.net/ projects/coolprop/files/CoolProp/6.3.0/shared_library/Linux/64bit/

WARNING: Since the *MODULE capability is not yet working in the Windows build of LS-DYNA, do not attempt to use a Windows DLL version of the COOLPROP shared library.

To use the COOLPROP shared library with this keyword card, load this shared library into LS-DYNA using the *MODULE capability. The following *MODULE card needs to appear before a *DUALCESE_MODEL card (not inside the file hierarchy of any file specified with a *DUALCESE_MODEL card):

```
*MODULE_LOAD
UserA - DUALCESE COOLPROP
    < path to installed COOLPROP shared library >
```


## Card Summary:

Card 1. This card is required.

| EOSID | NCOMP | TYPE | PHASE | TABULAR |  |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

Card 2. Include as many cards as needed to specify mole fractions for the NCOMP components of the fluid.

| MOL_FR1 | MOL_FR2 | MOL_FR3 | MOL_FR4 | MOL_FR5 | MOL_FR6 | MOL_FR7 | MOL_FR8 |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

Card 3. Include this card when the TABULAR field is active on Card 1.

| N_T | N_DEN | DEN_LOW | DEN_HIGH | T_LOW | T_HIGH |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |

Card 4. This card is required.

## Data Card Definitions:

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | EOSID | NCOMP | TYPE | PHASE | TABULAR |  |  |  |
| Type | 1 | 1 | A | A | A |  |  |  |
| Default | none | none | none | GAS | optional |  |  |  |

VARIABLE
EOSID ID for this EOS
NCOMP Number of components in the fluid composition

EQ.PURE: A single component fluid (default)
EQ.PSEUDOPURE: A predefined fluid mixture
EQ.MIXTURE: A fluid mixture with NCOMP components
PHASE Phase of the fluid.
EQ.GAS: Gas phase
EQ.LIQUID: Liquid phase
TABULAR Type of lookup tables to build for this EOS:
EQ. $<B L A N K>:$ No table lookup (default)
EQ.P_EIN: Build tables of pressure and internal energy, both as a function of density and temperature.

COOLPROP Parameters by Fluid Component. Repeat this card as many times as needed to input mole fractions for the NCOMP components of the fluid.

| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | MOL_FR1 | MOL_FR2 | MOL_FR3 | MOL_FR4 | MOL_FR5 | MOL_FR6 | MOL_FR7 | MOL_FR8 |
| Type | F | F | F | F | $F$ | $F$ | $F$ | F |

## VARIABLE

MOL_FR $i$

## DESCRIPTION

Mole fraction of the $i^{\text {th }}$ component

COOLPROP EOS Table Density and Temperature Ranges. This card is included when the TABULAR option on Card 1 is active.

| Card 3 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | N_T | N_DEN | DEN_LOW | DEN_HIGH | T_LOW | T_HIGH |  |  |
| Type | I | I | F | F | F | F |  |  |
| Default | none | none | none | none | none | none |  |  |
| Remarks | 1 | 1 | 2 | 2 | 2 | 2 |  |  |


| VARIAB |  | DESCRIPTION |  |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| N_T |  | Number of temperature values in the tables |  |  |  |  |  |  |
| N_DEN |  | Number of density values (on a log scale) in the tables |  |  |  |  |  |  |
| DEN_LOW |  | Minimum density available in the tables (in model units) |  |  |  |  |  |  |
| DEN_HIGH |  | Maximum density available in the tables (in model units) |  |  |  |  |  |  |
| T_LOW |  | Minimum temperature available in the tables (in model units) |  |  |  |  |  |  |
| T_HIGH |  | Maximum temperature available in the tables (in model units) |  |  |  |  |  |  |
| Name of CoolProp fluid. This card is required. |  |  |  |  |  |  |  |  |
| Card 4 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| Variable |  |  |  |  |  |  |  |  |
| Type |  |  |  |  |  |  |  |  |

VARIABLE
FLUIDNAME

## DESCRIPTION

Name of a fluid that has an EOS in CoolProp. For a list of the supported pure and pseudo-pure fluids, see:
http:/ /www.coolprop.org/fluid_properties/PurePseudo-Pure.html\#list-of-fluids

Note that the predefined fluid mixtures are not supported at this time.

## Remarks:

1. Number of Values in the Lookup Tables. The number of density and temperature values in the tables should not be too few to give good resolution of the EOS. Note that the cost of building the EOS from these tables rises with these numbers, as well as the computer memory required. Nevertheless, if these numbers are too small $(<20)$, then the accuracy may suffer, while larger numbers of density and temperature points improves the accuracy.
2. Valid Value Ranges for the Lookup Tables. For many equations of state in the CoolProp library, there is a range of valid densities and temperatures. Thus, the low and high limits for the table densities and temperatures should not lie outside these ranges. Please refer to the CoolProp documentation for that information.

## *DUALCESE_EOS_IDEAL_GAS

Purpose: Define the coefficients $C_{v}$ and $C_{p}$ in the equation of state for an ideal gas in the dual CESE fluid solver.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | EOSID | CV | CP | E0 |  |  |  |  |
| Type | I | F | F | F |  |  |  |  |
| Default | none | 717.5 | 1004.5 | 0.0 |  |  |  |  |
| Remarks |  | 1 | 1 | 2 |  |  |  |  |

## VARIABLE

EOSID
CV Specific heat at constant volume, $C_{v}$
$\mathrm{CP} \quad$ Specific heat at constant pressure, $C_{p}$
E0

## DESCRIPTION

Equation of state ID

Represents the heat of detonation released during the reactions, or the constant rate of afterburn energy added ( $\mathrm{E} 0=0.0$ is the default), $e_{0}$

## Remarks:

1. Units. As with other solvers in LS-DYNA, you are responsible for unit consistency. For example, if you want to use dimensionless variables, CV and CP should also be replaced by the corresponding dimensionless ones. If the dual CESE model has a specified system of units either directly from the *DUALCESE_MODEL card or inherited from the overall problem input, then these values need to be given in that unit system.
2. E0. E0 is used only with the hybrid multiphase solver where the EOS of the reactant is specified by this ideal gas EOS in the *DUALCESE_EOS_SET card.

## *DUALCESE_EOS_INFLATOR1

Purpose: Define an EOS using $C_{p}$ and $C_{v}$ thermodynamic expansions for an inflator gas mixture with a single temperature range.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | EOSID |  |  |  |  |  |  |  |
| Type | 1 |  |  |  |  |  |  |  |
| Default | none |  |  |  |  |  |  |  |


| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | CP0 | CP1 | CP2 | CP3 | CP4 |  |  |  |
| Type | F | F | F | F | F |  |  |  |
| Default | 0. | 0. | 0. | 0. | 0. |  |  |  |


| Card 3 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | CV0 | CV1 | CV2 | CV3 | CV4 |  |  |  |
| Type | F | F | F | F | F |  |  |  |
| Default | 0. | 0. | 0. | 0. | 0. |  |  |  |

## VARIABLE

EOSID
CP0, ..., CP4

## DESCRIPTION

Equation of state ID for the dual CESE solver
Coefficients of temperature-dependent specific heat at constant pressure

$$
C_{p}(T)=C_{p_{0}}+C_{p_{1}} T+C_{p_{2}} T^{2}+C_{p_{3}} T^{3}+C_{p_{4}} T^{4}
$$

VARIABLE
CV0, ..., CV4

## DESCRIPTION

Coefficients of temperature-dependent specific heat at constant volume

$$
C_{v}(T)=C_{v_{0}}+C_{v_{1}} T+C_{v_{2}} T^{2}+C_{v_{3}} T^{3}+C_{v_{4}} T^{4}
$$

## Remarks:

These coefficient expansions for the specific heats over the entire temperature range are generated by the zero-dimensional inflator model solver. See *CHEMISTRY_CONTROL_INFLATOR and *CHEMISTRY_INFLATOR_PROPERTIES for details related to running that solver.

## *DUALCESE_EOS_INFLATOR2

Purpose: Define an EOS using $C_{p}$ and $C_{v}$ thermodynamic expansions for an inflator gas mixture with two temperature ranges, one below 1000 Kelvin, and the other above 1000 Kelvin.

## Card Summary:

Card 1. This card is required.

| EOSID |  |  |  |  |  |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

Card 2. This card is required. This card with Card 3 specifies $C_{p}$. This card gives the coefficients for $\mathrm{T}<1000 \mathrm{~K}$.

| CP10 | CP11 | CP12 | CP13 | CP14 |  |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

Card 3. This card is required. This card gives the coefficients for T > 1000 K .

| CP20 | CP21 | CP22 | CP23 | CP24 |  |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

Card 4. This card is required. This card with Card 5 defines $C_{v}$. This card gives the coefficients for $\mathrm{T}<1000 \mathrm{~K}$.

| CV10 | CV11 | CV12 | CV13 | CV14 |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |

Card 5. This card is required. This card gives the coefficients for T $>1000 \mathrm{~K}$.

| CV20 | CV21 | CV22 | CV23 | CV24 |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |

## Data Card Definitions:

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | EOSID |  |  |  |  |  |  |  |
| Type | 1 |  |  |  |  |  |  |  |
| Default | none |  |  |  |  |  |  |  |

## VARIABLE

EOSID

## DESCRIPTION

Equation of state ID for the dual CESE solver

Coefficients for the expansion to determine specific heat at constant pressure for $\mathrm{T}<1000$ K.

| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | CP 10 | CP 11 | CP 12 | CP 13 | CP 14 |  |  |  |
| Type | F | F | F | F | F |  |  |  |
| Default | 0. | 0. | 0. | 0. | 0. |  |  |  |

Coefficients for the expansion to determine specific heat at constant pressure for $\mathrm{T}>1000$ K.

| Card 3 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | CP20 | CP21 | CP22 | CP23 | CP24 |  |  |  |
| Type | F | F | F | F | F |  |  |  |
| Default | 0. | 0. | 0. | 0. | 0. |  |  |  |

## VARIABLE

CP10, ..., CP14

CP20, ...,
CP24

## DESCRIPTION

Coefficients of temperature-dependent specific heat at constant pressure valid for $\mathrm{T}<1000 \mathrm{~K}$

Coefficients of temperature-dependent specific heat at constant pressure valid for $\mathrm{T}>1000 \mathrm{~K}$

Cards 2 and 3 give $C_{p}$ over the two temperature ranges:

$$
\mathrm{C}_{p}(T)= \begin{cases}\mathrm{CP} 10+\mathrm{CP} 11 \times T+\mathrm{CP} 12 \times T^{2}+\mathrm{CP} 13 \times T^{3}+\mathrm{CP} 14 \times T^{4} & \text { for } T<1000 \mathrm{~K} \\ \mathrm{CP} 20+\mathrm{CP} 21 \times T+\mathrm{CP} 22 \times T^{2}+\mathrm{CP} 23 \times T^{3}+\mathrm{CP} 24 \times T^{4} & \text { for } T>1000 \mathrm{~K}\end{cases}
$$

Card for the Expansion of Specific Heat at Constant Volume. Valid for T < 1000 K

| Card 4 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | CV10 | CV11 | CV12 | CV13 | CV14 |  |  |  |
| Type | F | F | F | F | F |  |  |  |
| Default | 0. | 0. | 0. | 0. | 0. |  |  |  |

Card for the Expansion of Specific Heat at Constant Volume. Valid for T > 1000 K

| Card 5 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | CV20 | CV21 | CV22 | CV23 | CV24 |  |  |  |
| Type | F | F | F | F | F |  |  |  |
| Default | 0. | 0. | 0. | 0. | 0. |  |  |  |

## VARIABLE

CV10, ...,
CV14
CV20, ...,
CV24

## DESCRIPTION

Coefficients of temperature-dependent specific heat at constant volume valid for $\mathrm{T}<1000 \mathrm{~K}$

Coefficients of temperature-dependent specific heat at constant volume valid for $\mathrm{T}>1000 \mathrm{~K}$

Cards 4 and 5 give $C_{v}$ over the two temperature ranges:

$$
C_{v}(T)= \begin{cases}\mathrm{CV} 10+\mathrm{CV} 11 \times T+\mathrm{CV} 12 \times T^{2}+\mathrm{CV} 13 \times T^{3}+\mathrm{CV} 14 \times T^{4} & \text { for } T<1000 \mathrm{~K} \\ \mathrm{CV} 20+\mathrm{CV} 21 \times T+\mathrm{CV} 22 \times T^{2}+\mathrm{CV} 23 \times T^{3}+\mathrm{CV} 24 \times T^{4} & \text { for } T>1000 \mathrm{~K}\end{cases}
$$

## Remarks:

These coefficient expansions for the specific heats over two temperature ranges are generated by the zero-dimensional inflator model solver. See *CHEMISTRY_CONTROL_INFLATOR and *CHEMISTRY_INFLATOR_PROPERTIES for details related to running that solver.

## *DUALCESE_EOS_JWL

Purpose: Define a JWL-type EOS that provides a means to represent a condensed phase explosive in a dual CESE multiphase model.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | EOSID | A | B | R1 | R2 | GAMMAO | RH00 | E0 |
| Type | I | F | F | F | F | F | F | F |
| Default | none | none | none | none | none | none | none | 0.0 |


| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | CV |  |  |  |  |  |  |  |
| Type | F |  |  |  |  |  |  |  |
| Default | none |  |  |  |  |  |  |  |

## VARIABLE

EOSID
A

B
R1
R2
GAMMA0
RHOO
E0 Represents the heat of detonation released during the reactions or
the constant rate of afterburn energy added ( $\mathrm{E} 0=0.0$ for standard
Represents the heat of detonation released during the reactions or
the constant rate of afterburn energy added ( $\mathrm{E} 0=0.0$ for standard JWL EOS), $e_{0}$

## DESCRIPTION

Equation of state ID for the dual CESE solver.
Model parameter, $A$ (in pressure units)
Model parameter, $B$ (in pressure units)
Model constant, $R_{1}$ (dimensionless)
Model constant, $R_{2}$ (dimensionless)
Gruneisen coefficient
Initial or reference density, $\rho_{0}$

Heat capacity, $C_{v}$

## Remark:

The equations of state of a Mie-Gruneisen form are given by:

$$
P(\rho, e)=P_{\mathrm{ref}}+\Gamma(\rho) \rho\left[e-e_{\mathrm{ref}}(\rho)\right]
$$

Here $\Gamma(\rho)$ is the Gruneisen coefficient. Equations of state of this type are very popular in condensed phase explosive modeling. Depending on the form of the reference pressure and energy functions, different EOS types can be retrieved. The JWL EOS is one type with reference pressure and energy given by:

$$
\begin{aligned}
P_{\text {ref }}(\rho) & =A \exp \left(\frac{-R_{1} \rho_{0}}{\rho}\right)+B \exp \left(\frac{-R_{2} \rho_{0}}{\rho}\right) \\
e_{\text {ref }}(\rho) & =\frac{A}{\rho_{0} R_{1}} \exp \left(\frac{-R_{1} \rho_{0}}{\rho}\right)+\frac{B}{\rho_{0} R_{2}} \exp \left(\frac{-R_{2} \rho_{0}}{\rho}\right)-e_{0}
\end{aligned}
$$

## *DUALCESE_EOS_REFPROP

Purpose: Define an equation of state (EOS) to be evaluated using the REFPROP EOS library

Note that the REFPROP library is provided by ANSYS; it is REFPROP v10.0 from NIST.

WARNING: Since the *MODULE capability is not yet working in the Windows build of LS-DYNA, do not attempt to use a Windows DLL version of the REFPROP shared library that comes with REFPROP v10.0.

To use the REFPROP shared library with this keyword card, load this shared library into LS-DYNA using the *MODULE capability. The following *MODULE card needs to appear before a *DUALCESE_MODEL card (not inside the file hierarchy of any file specified with a *DUALCESE_MODEL card):

```
*MODULE_LOAD
UserA DUALCESE REFPROP
    < path to the installed REFPROP shared library >
```

In addition, for REFPROP to be able to find the appropriate EOS data, *DUALCESE_EOS_REFPROP_PATH must also be given somewhere inside a *DUALCESE_MODEL file hierarchy to point to the place in your filesystem where REFPROP has been installed.

## Card Summary:

Card 1. This card is required.

| EOSID | NCOMP | TYPE | PHASE | TABULAR |  |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

Card 2. Include as many cards as needed to specify mole fractions for the NCOMP components of the fluid.

| MOL_FR1 | MOL_FR2 | MOL_FR3 | MOL_FR4 | MOL_FR5 | MOL_FR6 | MOL_FR7 | MOL_FR8 |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

Card 3. This card is included when the TABULAR field is active on Card 1.

| N_T | N_DEN | DEN_LOW | DEN_HIGH | T_LOW | T_HIGH |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |

Card 4. This card is required.

## FLUIDNAME

## Data Card Definitions:

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | EOSID | NCOMP | TYPE | PHASE | TABULAR |  |  |  |
| Type | I | 1 | A | A | A |  |  |  |
| Default | none | none | none | GAS | optional |  |  |  |

VARIABLE
EOSID ID for this EOS
NCOMP Number of components in the fluid composition
TYPE Fluid type:
EQ.PURE: A single component fluid (default)
EQ.PSEUDOPURE: A predefined fluid mixture
EQ.MIXTURE: A fluid mixture with NCOMP components
PHASE Phase of the fluid:
EQ.GAS: Gas phase
EQ.LIQUID: Liquid phase
TABULAR Type of lookup tables to build for this EOS:
EQ. $<$ BLANK>: No table lookup (default)
EQ.P_EIN: Build tables of pressure and internal energy, both as a function of density and temperature.

REFPROP Parameters by Fluid Component. Repeat this card as many times as needed to input mole fractions for the NCOMP components of the fluid.

| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | MOL_FR1 | MOL_FR2 | MOL_FR3 | MOL_FR4 | MOL_FR5 | M0L_FR6 | MOL_FR7 | MOL_FR8 |
| Type | $F$ | F | F | F | F | $F$ | $F$ | $F$ |
| Default | none | none | none | none | none | none | none | none |

## VARIABLE

## DESCRIPTION

MOL_FR $i \quad$ Mole fraction of the $i^{\text {th }}$ component of the fluid
REFPROP EOS Table Density and Temperature Ranges. This card is included when the TABULAR field is active on Card 1.

| Card 3 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | N_T | N_DEN | DEN_LOW | DEN_HIGH | T_LOW | T_HIGH |  |  |
| Type | I | I | F | F | F | F |  |  |
| Default | none | none | none | none | none | none |  |  |
| Remarks | 1 | 1 | 2 | 2 | 2 | 2 |  |  |

## VARIABLE

N_T

DEN_LOW
DEN_HIGH
T_LOW
T_HIGH

N_DEN Number of density values (on a log scale) in the table

## DESCRIPTION

Number of temperature values in the tables

Minimum density available in the tables (in model units)
Maximum density available in the tables (in model units)
Minimum temperature available in the tables (in model units)
Maximum temperature available in the tables (in model units)

Name of REFPROP fluid. This card is required.

| Card 4 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |
| Variable | FLUIDNAME |  |  |  |  |  |  |  |
| Type |  |  |  |  |  |  |  |  |

## VARIABLE

FLUID-
NAME

## DESCRIPTION

Name of a fluid that has an EOS in REFPROP. For a list of the supported pure and pseudo-pure fluids, see the directory of supported fluids that comes with the REFPROP v10.0 library from ANSYS.

Note that the predefined fluid mixtures are not supported at this time.

## Remarks:

1. Number of Values in the Lookup Table. The number of values of density and temperature axes of the tables should not be too few to give good resolution of the EOS. Note that the cost of building the EOS from these tables rises with these numbers, as well as the computer memory required. Nevertheless, if these numbers are too small $(<20)$, then the accuracy may suffer, while larger numbers of density and temperature points improves the accuracy.
2. Valid Value Ranges for the Lookup Tables. For many equations of state in the REFPROP library, a range of densities and temperatures are valid. Thus, the low and high limits for the table densities and temperatures should not lie outside these ranges. Please refer to the REFPROP documentation for that information.

## *DUALCESE_EOS_REFPROP_PATH

Purpose: Provide the file path to the directory where the REFPROP EOS system is installed.

Note that in any problem where a *DUALCESE_EOS_REFPROP card is used, you must also provide a *DUALCESE_EOS_REFPROP_PATH card somewhere inside a *DUALCESE_MODEL file hierarchy to point to the place in your filesystem where REFPROP has been installed so that the appropriate EOS data can be loaded.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |
| Variable | PATH |  |  |  |  |  |  |  |
| Type | A |  |  |  |  |  |  |  |

## VARIABLE

PATH

## DESCRIPTION

Path giving the directory where the REFPROP data is installed.

## *DUALCESE_EOS_SET

Purpose: Define a set of equations of state that are used together to compute the thermodynamic state of a multiphase fluid for the dual CESE solver.

WARNING: As of version R14, this keyword is deprecated. Instead, use *DUALCESE_SOLVER_HYBRID_MULTIPHASE for the hybrid multiphase case and *DUALCESE_-SOLVER_TWO-PHASE_MULTIPHASE for the twophase multiphase case.

Include one card for each dual CESE multiphase mesh. This input ends at the next keyword ("*") card.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | EOSSID | EOSINID | EOSRCTID | EOSPRDID |  |  |  |  |
| Type | 1 | 1 | 1 | 1 |  |  |  |  |
| Default | none | none | none | none |  |  |  |  |

## VARIABLE

EOSSID

## DESCRIPTION

Set ID of the EOS mixture of a given multiphase fluid

EOSINID
EOSRCTID
EOSPRDID

EOS ID of the inert component of the multiphase mixture EOS ID of the reactant phase of the multiphase mixture EOS ID of the product phase of the multiphase mixture

## *DUALCESE_EOS_STIFFENED_GAS

Purpose: Define a stiffened gas type fluid EOS for use by the dual CESE solver.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | EOSID | GA | BT |  |  |  |  |  |
| Type | I | F | F |  |  |  |  |  |
| Default | none | none | none |  |  |  |  |  |

## VARIABLE

EOSID
GA Adiabatic exponent, $\gamma$. Must be $>$ 1.0.
BT $\quad$ Reference pressure, $\beta$. Must be $\geq 0.0$

## Remark:

The stiffened gas equation of state:

$$
p(\rho, e)=(\gamma-1) \rho e-\gamma \beta
$$

provides a fundamental characterization of material properties of fluids. Here $e$ is the internal energy per unit mass, $\rho$ is the density, and $\gamma$ and $\beta$ are two thermodynamic constants. $\gamma$ and $\beta$ can be determined by a fitting procedure from laboratory data. A typical set of parameter values for water are: $\gamma=7$ and $\beta=3000 \mathrm{~atm}$ while for human blood are: $\gamma=5.527$ and $\beta=614.6 \mathrm{MPa}$. In addition to the modelling of a liquid, it is often used to describe other type of materials, including many compressible solids of practical importance.

## *DUALCESE_EOS_VAN_DER_WAALS_GENERALIZED

Purpose: Define a Van Der Waals generalized type fluid EOS for use in the dual CESE solver.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | EOSID | A | B | GA | BT |  |  |  |
| Type | I | F | F | F | F |  |  |  |
| Default | none | none | none | none | none |  |  |  |

## VARIABLE

EOSID

A

B
GA Ratio of specific heats, $\gamma$. Must be $>$ 1.0.
BT Reference pressure, $\beta$. Must be $\geq 0.0$. See *DUALCESE_EOS_STIFFENED_GAS.

## Remark:

The generalized Van der Waals equation of state can be written as:

$$
p(\rho, e)=\frac{(\gamma-1)}{1-b \rho}\left(\rho e-\beta+a \rho^{2}\right)-\left(\beta+a \rho^{2}\right)
$$

Here $e$ denotes the specific internal energy, $\gamma$ is the ratio of specific heats $(\gamma>1), \beta$ is a reference pressure, and the quantities $a$ and $b$ are the van der Waals gas constants for molecular cohesive forces and the finite size of molecules, respectively ( $a \geq 0,0 \leq b<$ $1 / \rho)$. This EOS is often used to deal with possible real-gas effect (without phase transition) when both the temperature and pressure are high. When $\beta=0$, the generalized van der Waals equation of state becomes a standard Van der Waals equation of state. If $a=b=$ 0 , the stiffened gases EOS will be recovered.

## References:

[1] G. Allaire, S. Clerc, S. Kokh, A five-equation model for the simulation of interfaces between compressible fluids. J. Comp. Phys. 181 (2) (2002) 577-616.
[2] K.M. Shyue, A fluid-mixture type algorithm for compressible multicomponent flow with van der Waals equation of state, J. Comp. Phys. 156, 43 (1999)

## *DUALCESE_FSI_EXCLUDE

Purpose: Provide a list of mechanics solver parts that are not involved in the dual CESE FSI calculation. This keyword is intended for increasing computational efficiency by excluding parts that will not involve significant FSI interactions with the dual CESE compressible fluid solver.

Include as many cards as needed. This input ends at the next keyword ("*") card.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | PID1 | PID2 | PID3 | PID4 | PID5 | PID6 | PID7 | PID8 |
| Type | 1 | 1 | 1 | 1 | 1 | 1 | 1 | 1 |
| Default | none | none | none | none | none | none | none | none |

VARIABLE
PIDn

## DESCRIPTION

IDs of mechanics parts that will be excluded from the FSI interaction calculation with the dual CESE solver

## *DUALCESE_INCLUDE_MODEL

Purpose: Provide the filename of a file containing additional keywords belonging to a dual CESE model. Any number of these *DUALCESE_INCLUDE_MODEL keywords may be used in a single dual CESE model, where at the top level the overall model begins with a *DUALCESE_MODEL card.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |
| Variable |  |  |  |  |  |  |  |  |
| Type | FILENAME |  |  |  |  |  |  |  |

VARIABLE
FILENAME

## DESCRIPTION

Filename of the keyword file containing more of the dual CESE model. This card is only allowed inside a file that is given in one instance of a *DUALCESE_MODEL keyword card.

## *DUALCESE_INITIAL

Purpose: Specify constant initial conditions for flow variables at the centroid of each dual CESE fluid element.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | U | V | W | RH | P | T |  | IFUNC |
| Type | F | F | F | F | F | F |  | I |
| Default | 0 | 0.0 | 0.0 | 1.225 | 0.0 | 0.0 |  | none |

## VARIABLE

U, V, W
RHO Density, $\rho$
P Pressure, $P$
T Temperature, $T$
IFUNC Option to define initial conditions using *DEFINE_FUNCTION cards:

EQ.0: Not in use.
EQ.1: All values for initial velocity, pressure, density, and temperature now refer to *DEFINE_FUNCTION IDs. In these functions, the following parameters are allowed: $f(x, y, z)$, meaning that each variable's initial profile is a function of position.

## Remarks:

1. Required Input. Usually, only two of $\rho, P$, and $T$ are needed to be specified (besides the velocity). If all three are given, only $\rho$ and $P$ will be used.
2. Applicable Elements. These initial conditions will be applied in those elements that have not been assigned a value by *DUALCESE_INITIAL_OPTION cards for individual elements or sets of elements.

## *DUALCESE_INITIAL_SET

Purpose: Specify initial conditions for the flow variables at the centroid of each element in an element subset of the dual CESE mesh.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | ESID | IFUNC |  |  |  |  |  |  |
| Type | 1 | 1 |  |  |  |  |  |  |
| Default | none | none |  |  |  |  |  |  |


| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | U | V | W | RH 0 | P | T |  |  |
| Type | F | F | F | F | F | F |  |  |
| Default | none | none | none | none | none | none |  |  |

## VARIABLE

ESID Element set ID (see *DUALCESE_ELEMENTSET)
IFUNC Option to define initial conditions using *DEFINE_FUNCTION cards:

EQ.0: Not in use.
EQ.1: All values for initial velocity, pressure, density, and temperature now refer to *DEFINE_FUNCTION IDs. In these functions, the following parameters are allowed: $f(x, y, z)$, meaning that each variable's initial profile is a function of position.
$\mathrm{U}, \mathrm{V}, \mathrm{W} \quad x, y$, and $z$ velocity components, respectively
RHO Density, $\rho$
P Pressure, $P$

VARIABLE
T Temperature, $T$

## Remarks:

1. Required Input. Usually, only two of $\rho, P$ and $T$ are needed to be specified (besides the velocity). If all three are given, only $\rho$ and $P$ will be used.
2. Keyword Priority. The priority of this card is higher than *DUALCESE_INITIAL. Thus, if an element is assigned an initial value by this card, *DUALCESE_INITIAL will no longer apply to that element.

## *DUALCESE_INITIAL_HYBRID

Purpose: Specifies values to use for velocity, pressure, etc. for problem initialization of a hybrid multiphase model on the dual CESE mesh. Note that these values can be overridden in some mesh elements by use of the *DUALCESE_INITIAL_HYBRID_SET card.

The hybrid multiphase model involves two materials, separated by a material surface. These materials do not mix. The first material is inert while the second material is an explosive mixture, composed of reactants and products. This model is useful for simulations with high explosives. See Michael and Nikiforakis 2016 for details about this model.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | Z1 | RA | UIC | VIC | WIC | RH01 | RHO_A | RH0_B |
| Type | F | F | F | $F$ | $F$ | $F$ | $F$ | $F$ |
| Default | none | none | none | none | none | none | none | none |


| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | PIC | TIC |  | IFUNC |  |  |  |  |
| Type | F | F |  | I |  |  |  |  |
| Default | none | none |  | none |  |  |  |  |

## VARIABLE

RA Mass fraction of the reactant (material $\alpha$ ) with respect to material 2 (the explosive mixture)

UIC, VIC, Multiphase flow velocity components in the $x, y$, and $z$-directions, WIC

RHO1 Density of material 1

## VARIABLE

RHO_A Density of the reactant (material $\alpha$ )
RHO_B Density of the product (material $\beta$ )
PIC Equilibrium multifluid pressure
TIC

IFUNC Option to define initial conditions using *DEFINE_FUNCTION cards:

EQ.0: Not in use.
EQ.1: All values for initial velocity, pressure, and temperature now refer to *DEFINE_FUNCTION IDs. In these functions, the following parameters are allowed: $f(x, y, z)$, meaning that each variable's initial profile is a function of position.

## *DUALCESE_INITIAL_HYBRID_SET

Purpose: Specifies values to use for velocity, pressure, etc. for problem initialization of a hybrid multiphase model in an element subset of the dual CESE mesh. The values specified here override the values set on *DUALCESE_INITIAL_HYBRID for the element subset.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | ESID | IFUNC |  |  |  |  |  |  |
| Type | I | I |  |  |  |  |  |  |
| Default | none | none |  |  |  |  |  |  |


| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | Z1 | RA | UIC | VIC | WIC | RH01 | RHO_A | RHO_B |
| Type | F | F | F | F | $F$ | $F$ | $F$ | $F$ |
| Default | none | none | none | none | none | none | none | none |


| Card 3 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | PIC | TIC |  |  |  |  |  |  |
| Type | F | F |  |  |  |  |  |  |
| Default | none | none |  |  |  |  |  |  |

VARIABLE
ESID Element set ID (see *DUALCESE_ELEMENTSET)
IFUNC Option to define initial conditions using *DEFINE_FUNCTION cards:

## DESCRIPTION

EQ.0: Not in use.
EQ.1: All values for initial velocity, pressure, and temperature now refer to *DEFINE_FUNCTION IDs. In these functions, the following parameters are allowed: $f(x, y, z)$, meaning that each variable's initial profile is a function of position.

Z1 Volume fraction of material 1 (or color function). This is usually a value of 0 or 1 . For numerical stability, however, use a small value instead of 0 .

RA Mass fraction of the reactant (material $\alpha$ ) with respect to material 2 (the explosive mixture)

UIC, VIC, Multiphase flow velocity components in the $x, y$, and $z$-directions, WIC respectively

RHO1 Density of material 1
RHO_A Density of the reactant (material $\alpha$ )
RHO_B Density of the product (material $\beta$ )
PIC Equilibrium multifluid pressure
TIC Equilibrium multifluid temperature

## *DUALCESE_INITIAL_TWO-PHASE

Purpose: Specify values to use for velocity, pressure, etc. for problem initialization of a two-phase multifluid model on the dual CESE mesh. Note that these values can be overridden in some mesh elements by use of the *DUALCESE_INITIAL_TWO-PHASE_SET card.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | Z1 | UIC | VIC | WIC | RH0_1 | RH0_2 | PIC | TIC |
| Type | F | F | F | F | F | F | F | F |
| Default | none | none | 0. | 0. | 0. | none | none | none |


| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | IFUNC |  |  |  |  |  |  |  |
| Type | I |  |  |  |  |  |  |  |
| Default | none |  |  |  |  |  |  |  |

## VARIABLE

Z1
UIC, VIC, WIC

RHO_1 Density of fluid 1
RHO_2

PIC

TIC

IFUNC respectively.

Density of fluid 2

## DESCRIPTION

Volume fraction of material 1 (or color function)

Multiphase flow velocity components in the $x, y$, and $z$-directions

Equilibrium multifluid pressure
Equilibrium multifluid temperature
Option to define initial conditions using *DEFINE_FUNCTION cards:

## DESCRIPTION

EQ.0: Not in use.
EQ.1: All values for initial velocity, pressure, and temperature now refer to *DEFINE_FUNCTION IDs. In these functions, the following parameters are allowed: $f(x, y, z)$, meaning that each variable's initial profile is a function of position.

## *DUALCESE_INITIAL_TWO-PHASE_SET

Purpose: Specify values to use for velocity, pressure, etc. for problem initialization of a two-phase multifluid model in an element subset of the dual CESE mesh.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | ESID | IFUNC |  |  |  |  |  |  |
| Type | I | 1 |  |  |  |  |  |  |
| Default | none | none |  |  |  |  |  |  |


| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | Z1 | UIC | VIC | WIC | RHO_1 | RHO_2 | PIC | TIC |
| Type | F | F | F | F | F | F | F | F |
| Default | none | none | none | none | none | none | none | none |

## VARIABLE

ESID
IFUNC
Option to define initial conditions using *DEFINE_FUNCTION cards:

EQ.0: Not in use.
EQ.1: All values for initial velocity, pressure, and temperature now refer to *DEFINE_FUNCTION IDs. In these functions, the following parameters are allowed: $f(x, y, z)$,
meaning that each variable's initial profile is a function of tions, the following parameters are allowed: $f(x, y, z)$,
meaning that each variable's initial profile is a function of position.

Z1 Volume fraction of material 1 (or color function)
UIC, VIC, Multiphase flow velocity components in the $x, y$, and $z$-directions, WIC

## DESCRIPTION

## Element set ID (see *DUALCESE_ELEMENTSET)

## VARIABLE

## DESCRIPTION

RHO_1 Density of material 1
RHO_2 Density of material 2
PIC Equilibrium multiphase flow pressure
TIC Equilibrium multiphase flow temperature

## *DUALCESE_MAT_GAS

Purpose: Define the fluid (gas) properties in a viscous flow for the dual CESE solver. In this model, the dynamic viscosity is determined using Sutherland's formula for viscosity, and the thermal conductivity is determined using the Prandtl Number.

Include as many cards as needed. This input ends at the next keyword ("*") card.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | MID | C1 | C2 | PRND |  |  |  |  |
| Type | I | F | F | F |  |  |  |  |
| Default | none | $1.458 \mathrm{E}-6$ | 110.4 | 0.72 |  |  |  |  |

## VARIABLE

## DESCRIPTION

MID Material ID
C1, C2 Two coefficients in the Sutherland's formula for viscosity:

$$
\mu=\frac{C_{1} T^{3 / 2}}{T+C_{2}}
$$

Here $C_{1}$ and $C_{2}$ are constants for a given gas. For example, for air at moderate temperatures

$$
C_{1}=1.458 \times 10^{-6} \mathrm{~kg} / \mathrm{msK}^{1 / 2}, \quad C_{2}=110.4 \mathrm{~K}
$$

PRND
Prandtl Number (used to determine the coefficient of thermal conductivity). It is approximately constant for most gases. For air at standard conditions PRND $=0.72$.

## Remarks:

1. Inviscid Flows. C1 and C2 are only used to calculate the viscosity in viscous flows, so for inviscid flows, this material card is not needed.
2. Thermal Coupling. The Prandtl number is used to extract the thermal conductivity. It is only needed when thermal coupling with the structure is activated.
3. Unit Consistency. As with other solvers in LS-DYNA, you are responsible for unit consistency. For example, if dimensionless variables are used, $C_{1}$ and $C_{2}$ should be replaced by the corresponding dimensionless ones. If the dual CESE
model has a specified system of units either directly from the *DUALCESE_MODEL card or inherited from the overall problem input, then these values need to be given in that unit system. Also, note that the formulas here require the temperature be given in either Kelvin or Rankine units.

## *DUALCESE_MAT_GAS_0

Purpose: Define the fluid (gas) properties in a viscous flow for the dual CESE solver. In this model, the dynamic viscosity and thermal conductivity are specified constants.

Material Definition Cards. Include one card for each instance of this material type. This input ends at the next keyword ("*") card.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | MID | MU | K |  |  |  |  |  |
| Type | I | F | F |  |  |  |  |  |
| Default | none | none | none |  |  |  |  |  |

## VARIABLE

MID
MU
K

## DESCRIPTION

Material ID
Fluid dynamic viscosity. $\mathrm{MU}=1.81 \times 10^{-5} \mathrm{~kg} / \mathrm{ms}$ for air at $15^{\circ} \mathrm{C}$.
Thermal conductivity of the fluid

## Remarks:

1. Fields that Depend on Problem Physics. The viscosity is only used viscous flows, so for inviscid flows, it is not necessary to define it. The thermal conductivity is only used to calculate the heat transfer between the structure and the thermal solver when coupling is activated.
2. Unit Consistency. As with other solvers in LS-DYNA, you are responsible for unit consistency. For example, if dimensionless variables are used, MU should be replaced by the corresponding dimensionless one. If the dual CESE model has a specified system of units either directly from the *DUALCESE_MODEL card or inherited from the overall problem input, then these values need to be given in that unit system.

## *DUALCESE_MAT_GAS_2

Purpose: Define the fluid (gas) properties in a viscous flow for the dual CESE solver. This model determines dynamic viscosity and thermal conductivity by combining Sutherland's formula with the Power law for dilute gases.

Material Definition Cards. Include one card for each instance of this material type. This input ends at the next keyword ("*") card.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | MID | MU0 | SMU | K0 | SK | T0 |  |  |
| Type | I | F | F | F | F | F |  |  |
| Default | none | $1.716 \mathrm{E}-5$ | 111. | 0.0241 | 194.0 | 273.0 |  |  |

## VARIABLE

MID
MU0 / SMU

## DESCRIPTION

Material ID

Two coefficients appearing in the equation derived by combining Sutherland's formula with the Power law for dilute gases:

$$
\frac{\mu}{\mu_{0}}=\left(\frac{T}{T_{0}}\right)^{3 / 2} \frac{T_{0}+S_{\mu}}{T+S_{\mu}}
$$

Here $\mu$ is the dynamic viscosity, $\mu_{0}$ is a reference value, and $S_{\mu}$ is an effective temperature called the Sutherland constant which is characteristic of the gas. For air at moderate temperatures,

$$
\mu_{0}=1.716 \times 10^{-5} \mathrm{Ns} / \mathrm{m}^{2}, \quad S_{\mu}=111 \mathrm{~K}
$$

K0/SK Two coefficients appearing in the equation derived by combining Sutherland's formula with the Power law for dilute gases:

$$
\frac{k}{k_{0}}=\left(\frac{T}{T_{0}}\right)^{3 / 2} \frac{T_{0}+S_{k}}{T+S_{k}} .
$$

Here $k$ is the thermal conductivity, $k_{0}$ is a reference value, and $S_{k}$ is the Sutherland constant, which is characteristic of the gas. For air at moderate temperatures,

$$
k_{0}=0.0241 \mathrm{~W} / \mathrm{m}, \quad S_{k}=194 \mathrm{~K}
$$

Reference temperature, $T_{0}$. The default value (273.0) is for air in K .

## Remarks:

1. Fields that Depend on Problem Physics. The viscosity is only used for viscous flows, so for inviscid flows, it is not necessary to define it. The thermal conductivity is only used to calculate the heat transfer between the structure and the thermal solver when coupling is activated.
2. Unit Consistency. As with other solvers in LS-DYNA, you are responsible for unit consistency. For example, if dimensionless variables are used, MU should be replaced by the corresponding dimensionless one. If the dual CESE model has a specified system of units either directly from the *DUALCESE_MODEL card or inherited from the overall problem input, then these values need to be given in that unit system. Also, note that the formulas here require the temperature be given in either Kelvin or Rankine units.

## *DUALCESE_MESH_GEOMETRY

Purpose: Set general purpose control variables for the dual CESE compressible flow solver.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | GEOM | IDC |  |  |  |  |  |  |
| Type | A | F |  |  |  |  |  |  |
| Default | none | 0.25 |  |  |  |  |  |  |
| Remarks | 1 | 2 |  |  |  |  |  |  |

## VARIABLE

GEOM

IDC Contact interaction detection coefficient (for FSI and conjugate heat transfer problems)

## Remarks:

1. Mesh and Boundary Conditions for 2D Problems. If you want to use the 2D $(G E O M=2 \mathrm{D})$ or 2D axisymmetric (GEOM=AXI) solver, the mesh should only be distributed in the $x y$-plane with the boundary conditions given only at the $x y$ domain boundaries. Otherwise, a warning message will be given, and the 3D solver will be triggered instead.

The 2D axisymmetric case will work only if the 2D mesh and corresponding boundary conditions are properly defined with the $x$ and $y$ coordinates corresponding to the radial and axial directions, respectively.
2. Contact Interaction Detection Coefficient. IDC is the same type of variable that is input on the *ICFD_CONTROL_FSI card. For an explanation, see Remark 1 for the *ICFD_CONTROL_FSI card.
3. Dual CESE Solver and Restarts. The dual CESE solver is not currently supported for restarts.

## *DUALCESE_MESH_PART

Purpose: Define dual CESE solver mesh parts, that is, specify the dual CESE FSI algorithm to use for a mesh part. In other words, this keyword allows you to restrict the type of solver (immersed boundary FSI, moving mesh FSI, or non-FSI Eulerian) used on a region of a dual CESE mesh. The part ID specified with this keyword corresponds to the second field of Card 1 for either *DUALCESE_ELE2D or *DUALCESE_ELE3D.

Part Cards. Include one card for each dual CESE part. This input ends at the next keyword ("*") card.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | PID | FSITYPE | MMSHID |  |  |  |  |  |
| Type | I | A | 1 |  |  |  |  |  |
| Default | none | optional | 0 |  |  |  |  |  |

## VARIABLE

PID

FSITYPE
FSI type to use on this part:
EQ.<BLANK>: If left blank, no FSI performed.
EQ.IBM: Immersed boundary FSI solver
EQ.MOVMESH: Moving mesh FSI solver (FSITYPE = MMM may also be used for the same effect)

MMSHID ID for the mesh motion algorithm to use for the moving mesh FSI solver on this part (region of the current dual CESE mesh). This ID refers to the ID of an instantiation of *DUALCESE_CONTROL_MESH_MOV.

## *DUALCESE_MODEL

Purpose: Set the units used by a dual CESE compressible flow problem, along with the name of the file specifying the dual CESE model. There can be any number of such models (each with a separate mesh), and each such model must be in a different file.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | UNITSYS | FILENAME |  |  |  |  |  |  |
| Type | A |  | A |  |  |  |  |  |

## VARIABLE

UNITSYS

FILENAME Filename of the keyword file containing the dual CESE model. Note that only *DUALCESE_... keyword cards are allowed in this file.

## *DUALCESE_NODE2D

Purpose: Define a node and its coordinates in the global coordinate system. The nodal point ID must be unique relative to other nodes defined with *DUALCESE_NODE2D or *DUALCESE_NODE3D cards.

Node Cards. Include as many cards in the following format as desired. This input ends at the next keyword ("*") card.

| Card 1 | 1 | 2 |  | 3 | 4 |  | 5 | 6 | 7 | 8 | 9 | 10 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | NID | X |  |  | Y |  |  |  |  |  |  |  |
| Type | 1 | F |  |  | F |  |  |  |  |  |  |  |
| Default | none | 0. |  |  | 0. |  |  |  |  |  |  |  |

## VARIABLE

NID
$\mathrm{X} \quad x$ coordinate
$Y \quad y$ coordinate

## *DUALCESE_NODE3D

Purpose: Define a node and its coordinates in the global coordinate system. The nodal point ID must be unique relative to other nodes defined with *DUALCESE_NODE3D or *DUALCESE_NODE2D cards.

Node Cards. Include as many cards in the following format as desired. This input ends at the next keyword ("*") card.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 | 9 | 10 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | NID | X | Y | Z |  |  |  |  |  |  |
| Type | I | F | F | F |  |  |  |  |  |  |
| Default | none | 0. | 0. | 0. |  |  |  |  |  |  |

## VARIABLE

## DESCRIPTION

NID Node number
$\mathrm{X} \quad x$ coordinate
$Y \quad y$ coordinate
Z $\quad$ z coordinate

## *DUALCESE_NODESET

Purpose: Define a nodal set of dual CESE mesh nodes.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | NSID |  |  |  |  |  |  |  |
| Type | 1 |  |  |  |  |  |  |  |
| Default | none |  |  |  |  |  |  |  |

Node ID Cards. Set one value per node in the set. Include as many cards as needed. This input ends at the next keyword ("**") card.

| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | NID1 | NID2 | NID3 | NID4 | NID5 | NID6 | NID7 | NID8 |
| Type | 1 | 1 | 1 | 1 | 1 | 1 | 1 | 1 |

## VARIABLE

## DESCRIPTION

NSID
Set ID of new node set. All dual CESE node sets should have a unique set ID.

NID $i \quad$ Node ID $i$

## *DUALCESE_PART

Purpose: Define dual CESE solver parts, that is, specify the dual CESE material and EOS information for a part. It also provides a means to restrict the type of solver used on a region of a dual CESE mesh, meaning an immersed boundary FSI solver, a moving mesh FSI solver, or a non-FSI Eulerian solver can be specified for just this part.

WARNING: As of version R14, this keyword is deprecated. Instead, for the part ID and FSI solver details, use *DUALCESE_MESH_PART. For the material properties, use the *DUALCESE_SOLVER_... card corresponding to the choice made with *DUALCESE_SOLVER_SELECTION.

Part Cards. Include one card for each dual CESE part. This input ends at the next keyword ("*") card.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | PID | MID | EOSID | FSITYPE | MMSHID |  |  |  |
| Type | I | 1 | 1 | A | 1 |  |  |  |
| Default | none | none | none | optional | 0 |  |  |  |

## VARIABLE

MID Material ID referring to a *DUALCESE_MAT_... material (see Remark 1)

EOSID Equation of state ID referring to a *DUALCESE_EOS_... EOS
FSITYPE $\quad$ FSI type to use on this part:
EQ. <BLANK>: If left blank, no FSI performed.
EQ.IBM: Immersed boundary FSI solver
EQ.MOVMESH: Moving mesh FSI solver (FSITYPE = MMM may also be used for the same effect)

VARIABLE
MMSHID

## DESCRIPTION

ID for the mesh motion algorithm to use for the moving mesh FSI solver on this part (region of the current dual CESE mesh). This ID refers to a *DUALCESE_CONTROL_MESH_MOV card ID.

## Remarks:

1. MID for Inviscid Flows. Since material coefficients are only used in viscous flows, MID can be left blank for inviscid flows.

## *DUALCESE_PART_MULTIPHASE

Purpose: Define dual CESE multiphase solver parts, that is, specify the dual CESE material and EOS information for a part. It also provides a means to restrict the type of solver used on a region of a dual CESE mesh, meaning an immersed boundary FSI solver, a moving mesh FSI solver, or a non-FSI Eulerian solver can be specified for just this part.

WARNING: As of version R14, this keyword is deprecated. Instead, for the part ID and FSI solver details, use *DUALCESE_MESH_PART. For the material properties, use the *DUALCESE_SOLVER_... card corresponding to the choice made with *DUALCESE_SOLVER_SELECTION.

Part Cards. Include one card for each dual CESE multiphase solver part. This input ends at the next keyword ("*") card.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | PID | REACT_ID | EOSSID | MID | FSITYPE | MMSHID |  |  |
| Type | 1 | 1 | 1 | 1 | A | 1 |  |  |
| Default | none | none | none | none | optional | 0 |  |  |

VARIABLE
PID

REACT_ID ID of chemical reaction rate model (see *DUALCESE_REACTION_RATE_... cards)

EOSSID Set ID of multiphase EOS set specification (see *DUALCESE_EOS_SET)

MID Material ID defined by a *DUALCESE_MAT_... card

VARIABLE
FSITYPE

MMSHID

## DESCRIPTION

FSI type to use on this part:
EQ.<BLANK>: If left blank, no FSI is performed.
EQ.IBM: Immersed boundary FSI solver
EQ.MOVMESH: Moving mesh FSI solver (FSITYPE = MMM may also be used for the same effect)

ID of the mesh motion algorithm to use for the moving mesh FSI solver on this part (region of the current dual CESE mesh). This ID refers to a *DUALCESE_CONTROL_MESH_MOV card ID.

## *DUALCESE_POINTSET

Purpose: Define a list of points used to output at specified sample times variables from the chosen dual CESE solver to binary database binout.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | PNTSID |  |  |  |  |  |  |  |
| Type | I |  |  |  |  |  |  |  |
| Default | none |  |  |  |  |  |  |  |

Point Cards. Provide as many cards as necessary. This input ends at the next keyword ("*") card.

| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | X | Y | Z |  |  |  |  |  |
| Type | F | F | F |  |  |  |  |  |
| Default | none | none | none |  |  |  |  |  |

## VARIABLE

PNTSID
$\mathrm{X}, \mathrm{Y}, \mathrm{Z} \quad$ Coordinates of a point. As many points as desired can be specified

## *DUALCESE_REACTION_RATE_IG

Purpose: Define a reaction rate law for the Ignition and Growth model to describe the conversion of reactants to products in the modeling of a condensed phase explosive in a dual CESE multiphase model. See Michael and Nikiforakis 2016 and Tarver 2005 for details about this law.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | REACT_ID | IGN | AA | BB | XX | GROW1 | CC | DD |
| Type | I | F | F | F | F | F | $F$ | $F$ |
| Default | none | none | none | none | none | none | none | none |


| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | YY | GROW2 | EE | GG | ZZ | IGMAX | G1MAX | G2MAX |
| Type | F | F | F | F | F | F | F | F |
| Default | none | none | none | none | none | none | none | none |

## VARIABLE

## DESCRIPTION

REACT_ID ID of reaction rate law
IGN Reaction ignition term parameter, $I$
$\mathrm{AA} \quad$ Reaction ignition term parameter, $a$
BB Reaction ignition term parameter, $b$
$X X \quad$ Reaction ignition term parameter, $x$
GROW1 Reaction growth term parameter, $G_{1}$
CC Reaction growth term parameter, $c$
DD Reaction growth term parameter, $d$
YY Reaction growth term parameter, $y$

## VARIABLE

GROW2
EE
GG
ZZ
IGMAX

G1MAX Maximum mass fraction of the product for reaction growth term, $\Phi_{\text {G1max }}$

G2MAX Maximum $\Phi$ for reaction completion term, $\Phi_{G 2 \max }$

## Remarks:

Using the notation of Michael and Nikiforakis 2016, the reaction rate law can be given as:

$$
\begin{aligned}
\frac{d \Phi}{d t}=I(1-\Phi)^{b}(\rho & -1-a)^{x} H\left(\Phi_{\mathrm{IG} \max }-\Phi\right) \\
& +G_{1}(1-\Phi)^{c} \Phi^{d} p^{y} H\left(\Phi_{\mathrm{G} 1 \max }-\Phi\right)+G_{2}(1-\Phi)^{e} \Phi^{g} p^{z} H\left(\Phi-\Phi_{\mathrm{G} 2 \max }\right)
\end{aligned}
$$

where $H$ is the Heaviside function. Here $\Phi$ is the mass fraction of the products, $p$ is the pressure, and $\rho$ is the density of the explosive mixture. Note that the pressure is assumed to be in equilibrium between the phases. $I, G_{1}, G_{2}, a, b, c, d, e, g, x, y$, and $z$ are constants. They depend on the explosive as well as the part of the detonation process being modeled; see Michael and Nikiforakis 2016 for details.

The constants $\Phi_{\mathrm{IGmax}}, \Phi_{\mathrm{G} 1 \text { max }}$, and $\Phi_{\mathrm{G} 2 \max }$ in the Heavyside functions indicate when each of the three stages of the reaction are dominant. It was developed by Tarver 2005 to describe the reaction of pressed solid explosives during shock initiation and detonation. Tarver 2005 developed this reaction law for pressed solid explosives to describe the observed reaction stages during shock initiation and detonation See Tarver 2005 for a description of each stage of the reaction.

## *DUALCESE_REACTION_RATE_IG_REDUCED

Purpose: Define a reduced form reaction rate law compared to that defined with *DUALCESE_REACTION_RATE_IG for describing the conversion of reactants to products in the modeling of a condensed phase explosive in a dual CESE multiphase model. See Michael and Nikiforakis 2016 for details about this reaction rate law.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | REACT_ID | GROW1 | CC | DD | YY | PHIO |  |  |
| Type | I | F | F | F | F | F |  |  |
| Default | none | none | none | none | none | none |  |  |

## VARIABLE

REACT_ID
GROW1 Reaction growth term parameter, $G_{1}$
CC Reaction growth term parameter, $c$
DD Reaction growth term parameter, $d$
YY Reaction growth term parameter, $y$
PHIO Additional parameter to account for the non-zero amount of reaction when the mass fraction of the products, $\phi$, is zero

## Remarks:

Michael and Nikiforakis 2016 simplified the Ignition and Growth model reaction law to a pressure dependent law with a single stage to make the reduced model. They excluded the first and third terms from the Ignition and Growth model to makes this law. However, to account for a finite amount of reaction from ignition when $\phi$ is zero, they added a constant $\phi_{0}$. This reduced law has the form:

$$
\frac{d \phi}{d t}=G_{1}(1-\phi)^{c}\left(\phi+\phi_{0}\right)^{d} p^{y} .
$$

All the parameters are the same as the Ignition and Growth model except $\Phi_{0}$. See *DUALCESE_REACTION_RATE_IG and Michael and Nikiforakis 2016 for details.

## *DUALCESE_REACTION_RATE_P_DEPEND

Purpose: Define an explicitly pressure-dependent reaction rate law for describing the conversion of reactants to products in the modeling of a condensed phase explosive in a dual CESE multiphase model. This law is from Banks et al 2008.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | REACT_ID | SIGMA | NU | N |  |  |  |  |
| Type | I | F | F | F |  |  |  |  |
| Default | none | none | none | none |  |  |  |  |

## VARIABLE

## DESCRIPTION

REACT_ID ID of reaction rate law
SIGMA A positive constant, $\sigma$
NU A positive constant, $v$
$\mathrm{N} \quad$ A positive constant, $n$

## Remarks:

Using the notation of Michael and Nikiforakis 2016, this simple pressure dependent reaction rate law by Banks et al 2008 can be stated as:

$$
\frac{d \phi}{d t}=\sigma \phi^{v} p^{n}
$$

Here $\phi$ is the mass fraction of the products, $p$ is the pressure, and $\sigma, v$, and $n$ are positive constants found from experimental data. As discussed by Banks et al 2008, this type of reaction rate law that depends explicitly on pressure is often preferred for modeling the detonation of solid explosives because the constants can be found by fitting experimental data and other physical variables are difficult to measure.

## *DUALCESE_SEGMENTSET

Purpose: Define a set of segments. For three-dimensional geometries, a segment can be triangular or quadrilateral. For two-dimensional geometries, a segment is a line defined by two nodes.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | SID |  |  |  |  |  |  |  |
| Type | I |  |  |  |  |  |  |  |
| Default | none |  |  |  |  |  |  |  |

Segment Cards. For each segment in the set include one card of this format. Include as many cards as necessary. This input ends at the next keyword ("*") card.

| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | N1 | N2 | N3 | N4 |  |  |  |  |
| Type | I | I | I | I |  |  |  |  |

## VARIABLE

SID
N1
N2 Nodal point $n_{2}$
N3 Nodal point $n_{3}$. To define a line segment, set N3 $=$ N2.
N4 Nodal point $n_{4}$. To define a triangular segment, set $\mathrm{N} 4=\mathrm{N} 3$. To define a line segment, set $\mathrm{N} 4=\mathrm{N} 2$.

| FACE | Hexahedron | Pentahedron | Pyramid | Tetrahedron |
| :---: | :---: | :---: | :---: | :---: |
| 1 | N1, N5, N8, N4 | N1, N2, N5 | N1, N4, N3, N2 | N1,N2, N4 |
| 2 | N2,N3, N7,N6 | N4, N6, N3 | N1,N2,N5 | N2,N3,N4 |


| FACE | Hexahedron | Pentahedron | Pyramid | Tetrahedron |
| :---: | :---: | :---: | :---: | :---: |
| 3 | N1, N2, N6, N5 | N1, N4, N3, N2 | N2, N3, N5 | N1,N3, N2 |
| 4 | N4, N8, N7, N3 | N2, N3, N6, N5 | N3, N4, N5 | N1,N4, N3 |
| 5 | N1,N4,N3,N2 | N1,N5,N6,N4 | N4,N1,N5 |  |
| 6 | N5,N6, N7,N8 |  |  |  |

Table 5-1. Face definitions for volume dual CESE elements

## *DUALCESE_SOLVER_SELECTION

Purpose: Set general purpose control variables for the dual CESE compressible flow solver.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | EQNS |  |  |  |  |  |  |  |
| Type | A |  |  |  |  |  |  |  |
| Default | EULER |  |  |  |  |  |  |  |
| Remark | 1 |  |  |  |  |  |  |  |
| VARIABLE |  |  |  |  |  |  |  |  |

EQNS Select the equations being solved with the dual CESE solver:
EQ.NS:
Navier-Stokes equations
EQ.EULER: Euler equations
EQ.HYBRID: Hybrid multiphase model
EQ.TWO-PHASE: Two-phase multiphase model
EQ.CAVITATION: Cavitation solver

## Remarks:

1. Dual CESE Solver and Restarts. The dual CESE solver is not currently supported for restarts.

## *DUALCESE_SOLVER_CAV_EQNS

Purpose: Set the properties of the cavitating flow mixture being solved with the cavitation solver.

Property Card. Include one card.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | EOSID |  |  |  |  |  |  |  |
| Type | 1 |  |  |  |  |  |  |  |
| Default | none |  |  |  |  |  |  |  |

VARIABLE
EOSID

DESCRIPTION
Equation of state (EOS) ID that must refer to an EOS defined with *DUALCESE_EOS_CAV_HOMOG_EQUILIB

## *DUALCESE_SOLVER_EULER_EQNS

Purpose: Set the properties of the gas flow being solved with the Euler equations solver.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | EOSID |  |  |  |  |  |  |  |
| Type | 1 |  |  |  |  |  |  |  |
| Default | none |  |  |  |  |  |  |  |

## VARIABLE

EOSID

## DESCRIPTION

Equation of state (EOS) ID that refers to an EOS defined by one of the following keywords: *DUALCESE_EOS_IDEAL_GAS, *DUALCESE_EOS_INFLATOR1, or *DUALCESE_EOS_INFLATOR2.

## *DUALCESE_SOLVER_HYBRID_MULTIPHASE

Purpose: Define the properties of the fluid and condensed phase materials via equations-of-state that are used together to compute the thermodynamic state in the hybrid multiphase dual CESE solver. Also, specify the single-step reaction rate model for the chemical reactions.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | EOSINID | EOSRCTID | EOSPRDID | REACT_ID |  |  |  |  |
| Type | 1 | 1 | 1 | 1 |  |  |  |  |
| Default | none | none | none | none |  |  |  |  |
| Remark | 1 | 1 | 1 | 2 |  |  |  |  |

## VARIABLE

## DESCRIPTION

EOSINID
EOS ID of the inert component of the multiphase mixture
EOSRCTID EOS ID of the reactant phase of the multiphase mixture
EOSPRDID EOS ID of the product phase of the multiphase mixture
REACT_ID
ID of chemical reaction rate model

## Remarks:

1. Supported EOS Types. Each EOS can be one of the following types:
*DUALCESE_EOS_IDEAL_GAS
*DUALCESE_EOS_STIFFENED_GAS
*DUALCESE_EOS_VAN_DER_WAALS_GENERALIZED
*DUALCESE_EOS_COCHRAN_CHAN
*DUALCESE_EOS_JWL.
2. Supported Reaction Rate Types. The reaction rate can be one of the following types:

*DUALCESE_REACTION_REAT_IG<br>*DUALCESE_REACTION_RATE_IG_REDUCED<br>*DUALCESE_REACTION_RATE_P_DEPEND

## *DUALCESE_SOLVER_NAVIER_STOKES

Purpose: Set the properties of the fluid flow being solved with the Navier-Stokes equations solver. That is, specify the dual CESE material and EOS information to be used.

Property Card. Include one card.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | EOSID | MID |  |  |  |  |  |  |
| Type | 1 | 1 |  |  |  |  |  |  |
| Default | none | none |  |  |  |  |  |  |

VARIABLE
EOSID

MID Material ID referring to a *DUALCESE_MAT_... material card (see Remark 1)

## Remarks:

1. MID for Inviscid Flows. Since material coefficients are only used in viscous flows, MID can be left blank for inviscid flows.

## *DUALCESE_SOLVER_TWO-PHASE_MULTIPHASE

Purpose: Define the properties of the fluid phase materials through equations-of-state that are used together to compute the thermodynamic state in the two-phase multiphase dual CESE solver. This solver does not compute phase changes.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | EOSID1 | EOSID2 |  |  |  |  |  |  |
| Type | 1 | 1 |  |  |  |  |  |  |
| Default | none | none |  |  |  |  |  |  |
| Remarks | 1 | 1 |  |  |  |  |  |  |

## VARIABLE

DESCRIPTION
EOSID1
EOS ID of the first inert component of the multiphase mixture
EOSID2
EOS ID of the second inert component of the multiphase mixture

## Remarks:

1. Supported EOS Types. Each EOS can be one of the following types:
*DUALCESE_EOS_IDEAL_GAS
*DUALCESE_EOS_STIFFENED_GAS
*DUALCESE_EOS_VAN_DER_WAALS_GENERALIZED
*DUALCESE_EOS_COCHRAN_CHAN
*DUALCESE_EOS_JWL

## *EM

The *EM keyword cards provide input for the electromagnetism module. This module is for solving 3D eddy-current, inductive heating or resistive heating problems. It can be coupled with the mechanical and thermal solvers. Typical applications include magnetic metal forming and welding. This module also includes coupling the boundary element method to the finite element method coupling. We intend this coupling for simulations involving a conductor interacting with air so that the air does not need to be meshed. The conductor is modeled with finite elements while the air is modeled with boundary elements.

*EM_2DAXI<br>*EM_BOUNDARY<br>*EM_BOUNDARY_PRESCRIBED<br>*EM_CIRCUIT<br>*EM_CIRCUIT_CONNECT<br>*EM_CIRCUIT_ROGO<br>*EM_CONTACT<br>*EM_CONTACT_RESISTANCE<br>*EM_CONTACT_SUBDOM<br>*EM_CONTROL<br>*EM_CONTROL_CONTACT<br>*EM_CONTROL_COUPLING<br>*EM_CONTROL_EROSION<br>*EM_CONTROL_MAGNET<br>*EM_CONTROL_SOLUTION<br>*EM_CONTROL_SWITCH<br>*EM_CONTROL_SWITCH_CONTACT<br>*EM_CONTROL_TIMESTEP

*EM_DATABASE_CIRCUIT<br>*EM_DATABASE_CIRCUITOD<br>*EM DATABASE_ELOUT<br>*EM_DATABASE_FIELDLINE<br>*EM_DATABASE_GLOBALENERGY<br>*EM_DATABASE_NODOUT<br>*EM_DATABASE_PARTDATA<br>*EM_DATABASE_POINTOUT<br>*EM_DATABASE_ROGO<br>*EM_DATABASE_TIMESTEP<br>*EM_EP_CELLMODEL_DEFINEFUNCTION<br>*EM_EP_CELLMODEL_FENTONKARMA<br>*EM_EP_CELLMODEL_FIZHUGHNAGUMO<br>*EM_EP_CELLMODEL_TENTUSSCHER<br>*EM_EP_CELLMODEL_TOMEK<br>*EM_EP_CELLMODEL_USERMAT<br>*EM_EP_CREATEFIBERORIENTATION<br>*EM_EP_ECG<br>*EM_EP_FIBERINITIAL<br>*EM_EP_PURKINJE_NETWORK<br>*EM_EP_TENTUSSCHER_STIMULUS<br>*EM_EOS_BURGESS<br>*EM_EOS_MEADON<br>*EM_EOS_PERMEABILITY<br>*EM_EOS_TABULATED1<br>*EM_EOS_TABULATED2

*EM_EXTERNAL_FIELD
*EM_ISOPOTENTIAL
*EM_ISOPOTENTIAL_CONNECT
*EM_ISOPOTENTIAL_ROGO
*EM_MAT_001
*EM_MAT_002
*EM_MAT_003
*EM_MAT_004
*EM_MAT_005
*EM_MAT_006
*EM_OUTPUT
*EM_OUTPUT_VTK
*EM_PERMANENT_MAGNET
*EM_POINT_SET
*EM_RANDLES_BATMAC
*EM_RANDLES_EXOTHERMIC_REACTION
*EM_RANDLES_MESHLESS
*EM_RANDLES_TSHELL
*EM_RANDLES_SHORT
*EM_RANDLES_SOLID
*EM_ROTATION_AXIS
*EM_SOLVER_BEM
*EM_SOLVER_BEMMAT
*EM_SOLVER_FEM
*EM_SOLVER_FEMBEM
*EM_SOLVER_FEMBEM_MONOLITHIC

## *EM_2DAXI

Purpose: Sets up the electromagnetism solver as 2D axisymmetric instead of 3D, on a given part, in order to save computational time as well as memory.

The electromagnetism is solved in 2D on a given cross section of the part (defined by a segment set), with a symmetry axis defined by its direction (at this time, it can be the $x$, $y$, or $z$ axis). The EM forces and Joule heating are then computed over the full 3D part by rotations. The part needs to be compatible with the symmetry, i.e. each node in the part needs to be the child of a parent node on the segment set, by a rotation around the axis. Only the conductor parts (with a *EM_MAT_... of type 2 or 4 ) should be defined as 2D axisymmetric.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | PID | SSID |  |  | STARSSID | ENDSSID | NUMSEC |  |
| Type | 1 | 1 |  |  | 1 | 1 | 1 |  |
| Default | none | none |  |  | none | none | none |  |

VARIABLE
PID
SSID Segment Set ID : Segment that will define the 2D cross section of the part where the EM field is solved

STARSSID, Used by the 2D axisymmetric solver to make the connection beENDSSID

NUMSEC Number of Sectors. This field gives the ratio of the full circle to the angular extension of the mesh. This has to be a power of two. For example, NUMSEC $=4$ means that the mesh of the part represents one fourth of the total circle. If this value is set to 0 , then the value from *EM_ROTATION_AXIS is used instead.

## Remarks:

1. At this time, either all or none of the conductor parts should be 2 D axisymmetric. In the future, a mix between 2 D axisymmetric and 3D parts will be allowed.

## *EM_BOUNDARY

Purpose: Define some boundary conditions for the electromagnetism problems.
Include as many cards as needed. This input ends at the next keyword ("*") card.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | SSID | BTYPE |  |  |  |  |  |  |
| Type | 1 | 1 |  |  |  |  |  |  |
| Default | none | none |  |  |  |  |  |  |

VARIABLE
SSID
BTYPE

DESCRIPTION
Segment Set Id
EQ.9: The faces of this segment set are eliminated from the BEM calculations (used for example for the rear or side faces of a workpiece).

## *EM_BOUNDARY_PRESCRIBED

Purpose: Prescribe a local boundary condition applied on nodes.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | BPID | BPTYPE | SETTYPE | SETID | VAL | LCID |  |  |
| Type | I | 1 | 1 | 1 | F | 1 |  |  |
| Default | none | none | none | none | 0. | 0 |  |  |

Optional Card.

| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | BIRTHT | DEATHT |  |  |  |  |  |  |
| Type | F | F |  |  |  |  |  |  |
| Default | 0. | $1 . \mathrm{e} 28$ |  |  |  |  |  |  |

## VARIABLE

ISOID
BPTYPE

## DESCRIPTION

ID of the prescribed boundary
Prescribed boundary type:
EQ.1: Short (scalar potential set to 0.)
EQ.2: Prescribed resistance (Robin B.C.)
EQ.3: Prescribed scalar potential (Dirichlet B.C.)
EQ.4: Prescribed current density (Neumann B.C.)

SETTYPE

SETID

Set type:
EQ.1: Segment set
EQ.2: Node set
EQ.3: Fluid part (see *ICFD_PART)

VAL

LCID

BIRTHT / Birth and death times for that prescribed boundary

## DESCRIPTION

Value of the resistance, current density or potential depending on BPTYPE. Ignored if LCID is defined.

Load curve ID defining the value of the resistance, voltage, or current as a function of time. If a negative value is entered, a *DEFINE_FUNCTION will be expected. The following parameters are allowed: f(time, emdt, curr, pot, cond, temp, potglob, currglob, areaglob, area, x, y, z). Pot/curr/area and potglob/currglob/areaglob are the local value of the scalar potential/current/area and the global averaged value on the prescribed boundary, respectfully. Cond is the local electrical conductivity, and $x, y$, and $z$ are the local coordinates.

DEATHT

## Remarks:

1. Supported Solvers. This keyword is only currently available for the resistive heating solver $(\mathrm{EMSOL}=3)$.

## *EM_CIRCUIT_\{OPTION\}

Available options include

## SOURCE

Purpose: Define an electrical circuit.
For the SOURCE option, the current will be considered uniform in the circuit. In general, this feature is used to model stranded conductors carrying a source current (in which case Amperes become Ampere-turns). This feature can also be useful for saving computational time in cases with a low frequency current and where the diffusion of the EM fields is a very fast process. In contrast, for the general case, the current density in a circuit is modeled in accordance with the solver type defined in EMSOL of *EM_CONTROL. For example, if an eddy current solver is selected, the diffusion of the current in the circuit is taken into account. See Remark 4 for a discussion of the available circuit types for source circuits.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | CIRCID | CIRCTYP | LCID | R/F | L/A | C/to | V0 | T0 |
| Type | I | I | I | F | F | F | F | F |
| Default | none | none | none | none | none | none | none | 0. |


| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | SIDCURR | SIDVIN | SIDVOUT | PARTID | IFREQST |  |  |  |
| Type | 1 | 1 | 1 | 1 | 1 |  |  |  |
| Default | none | none | none | none | 1 |  |  |  |

## VARIABLE

CIRCID
CIRCTYP

## DESCRIPTION

Circuit ID
Circuit type (see Remark 4 for source circuits):
EQ.1: Imposed current vs time defined by a load curve.

## VARIABLE

EQ.2: Imposed voltage vs time defined by a load curve. If a negative value is entered for LCID, its absolute value will refer to a DEFINE FUNCTION for a user defined circuit equation. If a DEFINE_FUNCTION is used, the following parameters are accepted: f(time, emdt, curr, curr1, curr2, pot1, pot2). emdt is the current timestep; curr, curr1, and curr2 refer to the current value at $t, t-1$ and $t-2$, respectfully; and pot1 and pot2 refer to the scalar potential at $t-1$ and $t-2$, respectfully.
EQ.3: R, L, C, V0 circuit (not available for source circuits)
EQ.11: Imposed current defined by an amplitude $A$, frequency $F$ and initial time $t_{0}: I=A \sin \left[2 \pi F\left(t-t_{0}\right)\right]$
EQ.12: Imposed voltage defined by an amplitude $A$, frequency $F$ and initial time $t_{0}: V=A \sin \left[2 \pi F\left(t-t_{0}\right)\right]$

EQ.21: Imposed current defined by a load curve over one period and a frequency $F$. See Remark 3.

EQ.22: Imposed voltage defined by a load curve over one period and a frequency F. See Remark 3.

LCID Load curve ID for CIRCTYP $=1,2,21$ or 22
R/F Value of the circuit resistance for CIRCTYP $=3$
Value of the frequency for CIRCTYP $=11,12,21$ or 22 . For CIRCTYP $=11$ or 12 , to have the frequency specified as a function of time with a load curve, a negative value can be entered with the absolute value corresponding to the load curve ID.

L/A Value of the circuit inductance for CIRCTYP $=3$
Value of the amplitude for CIRCTYP $=11$ or 12 . To have the amplitude specified as a function of time with a load curve, a negative value can be entered with the absolute value corresponding to the load curve ID.
$\mathrm{C} / \mathrm{t} 0 \quad$ Value of the circuit capacity for CIRCTYP $=3$
Value of the initial time $t_{0}$ for CIRCTYP $=11$ or 12
Value of the circuit initial voltage for CIRCTYP $=3$.
Starting time for CIRCTYPE $=3$. Default is at the beginning of the run.

## VARIABLE

SIDCURR

## DESCRIPTION

Segment set ID for the current. It uses the orientation given by the normal of the segments. To use the opposite orientation, use a '-' (minus) sign in front of the segment set ID.

CIRCTYP.EQ.1/11/21: The current is imposed through this segment set.
CIRCTYP.EQ.3: The current needed by the circuit equations is measured through this segment set.
CIRCTYP.EQ.2/12/22: Optional segment set that the current is forced through. See Remark 2.

SIDVIN Segment set ID for input voltage when CIRCTYP $=2,3,12$, or 22 or for input current when CIRCTYP $=1,11$, or 21 . The input voltage or current is oriented to enter the structural mesh, irrespective of the orientation of the segments.

SIDVOUT $\quad$ Segment set ID for output voltage when CIRCTYP $=2,3,12$, or 22 or for output current when CIRCTYP $=1,11$, or 21 . The output voltage or current is oriented to leave the structural mesh, irrespective of the orientation of the segment.

PARTID Part ID associated to the circuit. It can be any part ID associated to the circuit.

IFREQST
Frequency for recomputing the source terms for the SOURCE keyword option only. The source terms are recalculated every IFREQST time steps. By default, the source terms are recomputed every EM time step.

LT.0: |IFREQST| is a load curve ID giving the frequency for recomputing as a function of time.

## Remarks:

1. Imposed Current with Closed Loop Geometry. When defining a circuit with an imposed current (CIRCTYP 1, 11 or 21) in cases of a closed loop geometry (torus), SIDVIN and SIDVOUT cannot be defined and thus, only SIDCURR is necessary.
2. SIDCURR with Imposed Tension. When defining a circuit with an imposed tension (CIRCTYP 2, 12, 22), it is possible to also define SIDCURR. This can be

| Variable | Circuit Type (CIRCTYP) |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: |
|  | Imposed <br> 1: Current | Imposed <br> 2: Voltage | 3: R, L, C | 11: F, A, to | 12: F, A, t0 |
| LCID | M | M | - | - | - |
| R/L/C/V0 | - | - | M | - | - |
| F | - | - | - | M | M |
| A/t0 | - | - | - | M | M |
| SIDCURR | M | 0 | M | M | $\bigcirc$ |
| SIDVIN | M* | M | M | M ${ }^{\text {* }}$ | M |
| SIDVOUT | M* | M | M | M* | M |
| PARTID | M | M | M | M | M |
| Variable | 21: LCID, F | 22 : LCID, F |  |  |  |
| LCID | M | M | - | - | - |
| R/L/C/Vo | - | - | - | - | - |
| F | M | M | - | - | - |
| A/t0 | - | - | - | - | - |
| SIDCURR | M | $\bigcirc$ | - | - | - |
| SIDVIN | M* | M | - | - | - |
| SIDVOUT | M* | M | - | - | - |
| PARTID | M | M | - | - | - |

Table 6-1. Correspondence between circuit type and card entries. " M " indicates mandatory, " $\mathrm{M}^{* \prime}$ mandatory with exceptions (see Remark 1), " O " indicates optional, and "-" indicates ignored.
useful in circuits where various flow paths are possible for the current in order to force the entire current to go through SIDCURR.
3. CIRCTYP = $\mathbf{2 1}$ and 22. Circuit types 21 and 22 are for cases where the periodic current/tension does not exactly follow a perfect sinusoidal. You must provide the shape of the current/tension over one period through a LCID as well as the frequency.
4. Circuit Types for Source Circuits. For source circuits, CIRCTYP = 1, 2, 11, 12, 21, and 22 are supported. For imposed currents (CIRCTYP = 1, 11, or 21), source
circuits must be associated with an insulator material (MTYPE $=1$ ) in *EM_MAT_001. The solver relies on the Biot-Savart integration method to retrieve the influence of the source circuit on other conductors. This method is fast, but it is not possible to calculate the force on the coil.

You can also define a source circuit with an imposed voltage with CIRCTYP $=2$, 12 , or 22 (associated to a resistance value and a number of windings). The source circuit must be used with the monolithic solver (*EM_SOLVER_FEMBEM_MONOLITHIC) and become part of the FEM/BEM system. Therefore, it must be associated with a conductor material (MTYPE = 2) on *EM_MAT_001. The imposed voltage is slower than the imposed current, but a force applied to the coil can be calculated. Defining source circuits by their voltage instead of their current is very useful in cases where the current is not known, such as for actuators or electric motors.

## *EM_CIRCUIT_CONNECT

Purpose: This keyword connects several circuits together by imposing a linear constraint on the global currents of circuit pairs

$$
c_{1} i_{1}+c_{2} i_{2}=0 .
$$

This is especially useful for 2D axisymmetric models involving spiral or helical coils.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | CONID | CONTYPE | CIRC1 | CIRC2 | C 1 | C 2 |  |  |
| Type | I | I | I | I | F | F |  |  |
| Default | none | none | none | none | none | none |  |  |

## VARIABLE

CONID
CONTYPE

C1/C2

## DESCRIPTION

Id of the Circuit Connect
Type of connection between circuits. For the moment, it is only possible to combine circuits by imposing a linear constraint on the global current (=1).

Values of the linear constraints if CONTYPE $=1$.

## *EM_CIRCUIT_ROGO

Purpose: Define Rogowsky coils to measure a global current vs time through a segment set or a node set.

Include as many cards as needed. This input ends at the next keyword ("*") card.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | ROGID | SETID | SETTYPE | CURTYP |  |  |  |  |
| Type | 1 | 1 | 1 | 1 |  |  |  |  |
| Default | 0 | 0 | 0 | 0 |  |  |  |  |

VARIABLE
ROGID Rogowsky coil ID
SETID Segment or node set ID
SETTYPE Type of set:
EQ.1: Segment set
EQ.2: Node set (not available yet)
CURTYP Type of current measured:
EQ.1: Volume current
EQ.2: Surface current (not available yet)
EQ.3: Magnetic field flow (B field times Area)

## Remarks:

1. An ASCII file "em_rogo_xxx", with $x x x$ representing the rogoId, is generated for each *EM_CIRCUIT_ROGO card giving the value of the current or the magnetic field vs time.

## *EM_CONTACT

Purpose: Optional define and specify options on electromagnetic contacts between two sets of parts. Generally, it is used with the *EM_CONTACT_RESISTANCE. Fields left empty on this card default to the value of the equivalent field for *EM_CONTROL_CONTACT.

Contact Definition Cards. Include one card for each contact definition. This input ends at the next keyword ("*") card.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | CONTID | DTYPE | PSIDR | PSIDT | EPS1 | EPS2 | EPS3 | D0 |
| Type | I | I | I | I | F | F | F | F |
| Default | none | 0 | none | none | 0.3 | 0.3 | 0.3 | none |

## VARIABLE

CONTID

DTYPE

PSIDR

PSIDT Tracked surface part set ID
EPSi Contact coefficients for contact detection conditions. See the discussion below.

D0 Parameter for contact condition 3 when DTYPE = 1 (see remarks)

## Remarks:

In these remarks, we discuss the conditions for contact detection. For reference, Figure 0-1 illustrates which geometric values help determine contact. In this figure and discussion, $f_{r}$ is a face of the reference surface, and $f_{t}$ is a face of the tracked surface. Contact is detected when all of the following three conditions are satisfied:


Figure 0-1. Contact detection conditions between two faces.

1. Contact condition 1 :

$$
\mathbf{n}_{r} \cdot \mathbf{n}_{t} \leq-1+\varepsilon_{1}
$$

Here $\mathbf{n}_{r}$ and $\mathbf{n}_{t}$ are the normal vectors of faces $f_{r}$ and $f_{t}$, respectively, and $\varepsilon_{1}$ is an input parameter. See Figure 0-1.
2. Contact condition 2:

$$
\begin{aligned}
& -\varepsilon_{2} \leq \alpha_{1} \leq 1+\varepsilon_{2} \\
& -\varepsilon_{2} \leq \alpha_{2} \leq 1+\varepsilon_{2} \\
& -\varepsilon_{2} \leq \alpha_{3} \leq 1+\varepsilon_{2}
\end{aligned}
$$

( $\alpha_{1}, \alpha_{2}, \alpha_{3}$ ) are the local coordinates of point $P$ (see Figure 0-1). $P$ is the projection of point $a_{2}$ on face $f_{r} . \varepsilon_{2}$ is an input parameter.
3. Contact condition 3 depends on the contact type. For either possible condition, let $d$ be the distance between $P$ and $a_{t}$ (see Figure 0-1).
a) For contact type 0:

$$
d \leq \varepsilon_{3} S_{r}
$$

where $\varepsilon_{3}$ is an input parameter and $S_{r}$ is the minimum side length for $f_{r}$ :

$$
S_{r}=\min \left[d\left(a_{r}, b_{r}\right), d\left(b_{r}, c_{r}\right), d\left(c_{r}, a_{r}\right)\right] .
$$

b) For contact type 1 :

$$
d \leq D_{0},
$$

where $D_{0}$ is an input parameter.

## *EM_CONTACT_RESISTANCE

Purpose: Calculate the electric contact resistance of a previously defined EM contact in *EM_CONTACT.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | CRID | CONTID | CTYPE |  | JHRTYPE |  |  |  |
| Type | 1 | 1 | 1 |  | 1 |  |  |  |
| Default | none | none | none |  | none |  |  |  |


| Cards 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | DFID |  |  |  |  |  |  |  |
| Type | I |  |  |  |  |  |  |  |
| Default | none |  |  |  |  |  |  |  |

VARIABLE
CRID Resistive contact ID
CONTID EM contact ID defined in *EM_CONTACT
CTYPE Contact resistance type:
EQ.1: Electric contact resistance defined by user defined define function.

DFID
JHRTYPE

## DESCRIPTION

Define function ID (see Remark 1)
Indicates how the Joule heating calculated by the contact resistance shall be taken into account:

EQ.0: No addition. The Joule heating calculated by the contact resistance is not taken into account.


Figure 6-2. Electrode coming into contact with workpiece (RSW application).

## VARIABLE

## DESCRIPTION

EQ.1: The Joule heating coming from the contact resistance is divided and distributed evenly among all elements neighboring the contact surface.

## Remarks:

1. Define Function Parameters. In the *DEFINE_FUNCTION, the following parameters are allowed: f(time, emdt, arealoc, areatot, ctdist, currloc, currglob, rho_mst, rho_slv, cond_mst, cond_slv, ctpress_mst, ctpress_slv, temp_mst, temp_slv, vmstress_mst, vmstress_slv, press_mst, press_slv). Here,
time current time step
emdt EM time step
arealoc Local area associated with each face in contact
areatot Total contact area
ctdist Contact distance between the two faces in contact
currloc Local current density at the contact area
currglob Total current flowing through the contact area

| rho_mst | Density of the elements associated with the reference sur- <br> face side of the contact |
| :--- | :--- |
| rho_slv | Density of the elements associated with the tracked sur- <br> face side of the contact |
| cond_mst | Electrical conductivity of the elements associated with the <br> reference surface side of the contact |
| cond_slv | Electrical conductivity of the elements associated with the <br> tracked surface side of the contact |
| ctpress_mst | Contact pressure of the elements associated with the ref- <br> erence surface side of the contact |
| ctpress_slv | Contact pressure of the elements associated with the <br> tracked surface side of the contact |
| temp_mst | Temperature of the elements associated with the reference <br> surface side of the contact |
| temp_slv | Temperature of the elements associated with the tracked <br> surface side of the contact |
| vmstress_mst | von Mises stress of the elements associated with the refer- <br> ence surface side of the contact |
| vmstress_slv | von Mises stress of the elements associated with the <br> tracked surface side of the contact |
| press_mst | Pressure of the elements associated with the reference sur- <br> face side of the contact |
| Pressure of the elements associated with the tracked sur- <br> face side of the contact |  |

## *EM_CONTACT_SUBDOM

Purpose: Optional card used for defining a specific region where EM contact will be active. This allows saving some calculation time by limiting the contact search area. Must be used in conjunction with *EM_CONTROL_CONTACT.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | SDTYPE | MVTYPE | LCIDX/NID | LCIDY | LCIDZ |  |  |  |
| Type | 1 | 1 | 1 | 1 | 1 |  |  |  |
| Default | none | 0 | none | none | none |  |  |  |


| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | R | PMINX | PMINY | PMINZ | PMAXX | PMAXY | PMAXZ |  |
| Type | F | F | F | F | F | F | F |  |
| Default | none | none | none | none | none | none | none |  |

## VARIABLE

## DESCRIPTION

SDTYPE
Subdomain definition type :
EQ.1: Defined by box.
EQ.2: Defined by cylinder.
EQ.3: Defined by sphere.
MVTYPE Movement type of subdomain:
EQ.0: Static subdomain (Default).
EQ.1: Domain translates in the three directions by the velocities given by LCIDX,LCIDY,LCIDZ.

EQ.2: Domain follows the displacements of the node ID given by NID.

VARIABLE
LCIDX/NID

LCIDY/Z

R

PMINX/Y/Z

PMAXX/Y/Z

## DESCRIPTION

Time dependent load curve ID for the translational velocity in the X direction for MVTYPE $=1$, Node ID for MVTYPE $=2$.

Time dependent load curve IDs for MVTYPE $=1$ in the Y and Z directions.

Radius of the sphere if SDTYPE $=3$ or the cylinder if SDTYPE $=2$.

Point of minimum coordinates if SDTYPE $=1$. Origin point if SDTYPE $=3$. Axis head point if SDTYPE $=2$.

Point of maximum coordinates if SDTYPE $=1$. Axis tail point if SDTYPE $=2$.

## *EM_CONTROL

Purpose: Enable the EM solver and set its options.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | EMSOL | NUMLS | MACRODT | DIMTYPE | NPERIO |  | NCYLFEM | NCYLBEM |
| Type | 1 | 1 | F | 1 | 1 |  | 1 | 1 |
| Default | 0 | 100 | none | 0 | 2 |  | 5000 | 5000 |

## VARIABLE

EMSOL

NUMLS

MACRODT Macro time step when EMSOL $=2$
DIMTYPE EM dimension type:
EQ.0: 3D solve
EQ.1: 2D planar with zero thickness shell elements
EQ.3: 2D axisymmetric ( $Y$-axis only) with zero thickness elements

NPERIO

Number of periods for which the last is used to calculate the average Joule heat rate when $\mathrm{EMSOL}=2 . \mathrm{NPERIO}=2$ means that two periods of NUMLS steps will be calculated. Only the last period of

VARIABLE

NCYLFEM

NCYLBEM

## DESCRIPTION

NPERIO is used for the average Joule heat calculation. See Remark 1.

Number of electromagnetism cycles between the recalculation of FEM matrices. If negative the absolute value refers to a load curve that gives the number of electromagnetism cycles as a function of time. See Remark 2.

Number of electromagnetism cycles between the recalculation of BEM matrices. If negative, the absolute value refers to a load curve that gives the number of electromagnetism cycles as a function of time. See Remark 2.

## Remarks:

1. Number of periods and the average Joule heating calculation. The purpose of using more than one period to calculate the average Joule heating is to allow the different fields to adopt the correct amplitude and time shift when starting from 0.0 at $t=0$. NPERIO $=2$ means that two periods will be calculated of which only last one, being the second one in this case, will be used for the average Joule heat calculation. In some cases, using higher values might be required to achieve good accuracy and conversely, in others, NPERIO $=1$ might yield sufficient accuracy.
2. Order of precedence for NCYLFEM and NCYLBEM. You can set NCYLFEM and NCYLBEM with *EM_CONTROL_SOLUTION, *EM_SOLVER_FEM/BEM, or *EM_CONTROL. *EM_CONTROL_SOLUTION has the highest priority for setting these fields, while *EM_CONTROL has the lowest priority. If a field is left as the default on *EM_CONTROL_SOLUTION, LS-DYNA looks at the setting of that field on *EM_SOLVER_FEM/BEM. If left as default on *EM_SOLVER_FEM/BEM, LS-DYNA looks at the setting on *EM_CONTROL.

## *EM_CONTROL_CONTACT

Purpose: Activate the electromagnetism contact algorithms, which detect contact between conductors. Electromagnetic fields flow from one conductor to another when found to be in contact.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | EMCT | CCONLY | CTYPE | DTYPE | EPS1 | EPS2 | EPS3 | D0 |
| Type | I | I | I | I | F | F | F | F |
| Default | 0 | 0 | 0 | 0 | 0.3 | 0.3 | 0.3 | none |

## VARIABLE

## DESCRIPTION

EMCT
EM contact activation flag:
EQ.O: No contact detection
EQ.1: Contact detection
CCONLY Determines parts for which EM contact is active:
EQ.0: Contact detection between all active parts associated with a conducting material. (Default)
EQ.1: Only look for EM contact between parts associated through the *EM_CONTACT card. In some cases, this option can reduce the calculation time.

CTYPE Contact type:
EQ.-1: Node-to-node contact based on constraints on the scalar potential. See Remark 1.

EQ.0: Node-to-node penalty-based contact on the scalar potential.

EQ.1: Discrete mortar penalty contact on the scalar potential.
EQ.2: Continuous mortar penalty contact on the scalar potential and the vector potential (when active).

DTYPE Detection type. If *EM_CONTACT is not defined, the solver will look for global contact options in *EM_CONTROL_CONTACT. See *EM_CONTACT for details.

## VARIABLE

EQ.O: Contact type 0 (default).
EQ.1: Contact type 1.
EPSi Global contact coefficients used if the equivalent fields in *EM_CONTACT are empty. See *EM_CONTACT for details.

D0 Global parameter for contact condition 3 when DTYPE $=1$. See *EM_CONTACT for details.

## Remarks:

1. Contact types. In versions prior to R12, CTYPE $=0$ was the default EM contact for the resistive heating solver, while CTYPE $=-1$ was the default EM contact for the Eddy current solver. In versions R12 and later, CTYPE $=0$ became the default for all solvers. We recommend CTYPE $=1$ and 2 for the best accuracy.
2. Contact between BEM surfaces with the Eddy current solver. When the Eddy current solver is active and contact occurs between BEM surfaces, the solver automatically removes the faces on the contact surface and internally stitches the two BEM surfaces together to achieve a continuous closed BEM mesh.
*EM_CONTROL_COUPLING

## *EM_CONTROL_COUPLING

Purpose: Control couplings between various solvers with the EM solver.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | THCPL | SMCPL | THLCID | SMLCID | THCPLFL | SMCPLFL |  |  |
| Type | 1 | 1 | 1 | 1 | 1 | 1 |  |  |
| Default | 0 | 0 | 0 | 0 | 0 | 0 |  |  |

Optional Card.

| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | SMMOD | DFX | DFY | DFZ |  |  |  |  |
| Type | 1 | 1 | 1 | 1 |  |  |  |  |
| Default | 0 | none | none | none |  |  |  |  |

VARIABLE
THCPL

## DESCRIPTION

Coupling to the thermal solver. When turned on, the EM solver will transfer the Joule heating terms to the solid mechanics thermal solver.

EQ.O: Coupling on.
EQ.1: Coupling off.
SMCPL Coupling to the solid mechanics solver. When turned on, the EM solver will transfer forces to the solid mechanics solver.

EQ.O: Coupling on. Lorentz force density is transferred.
EQ.1: Coupling off.
EQ.2: Coupling on. Magnetic force surface density is transferred. More accurate representation of EM forces in cases involving magnets or nonlinear ferromagnets. See *EM_SOLVER_FEMBEM_MONOLITHIC.

VARIABLE

THLCID

SMLCID

THCPLFL

SMCPLFL

SMMOD Coupling to the solid mechanics solver. When turned on, the EM solver will transfer forces to the solid mechanics solver.

EQ.0: Off.
EQ.1: Force calculation at element level is decided by *DEFINE_FUNCTION. See DFX, DFY and DFZ.

EQ.2: Force calculation at element level is decided by usermat routine. See dyn21em.f and user_getEMForceArray routine.

Define function IDs for the force three components if $\mathrm{SMMOD}=1$. Arguments for the define functions are the same as in *EM_EOS_TABULATED2.

## *EM_CONTROL_EROSION

Purpose: Allows the EM solver to take eroded elements into account

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | ECTRL |  |  |  |  |  |  |  |
| Type | 1 |  |  |  |  |  |  |  |
| Default | 0 |  |  |  |  |  |  |  |

## VARIABLE

## DESCRIPTION

ECTRL
Erosion search :
EQ.O: Off. This means that the EM solver will ignore eroded elements and still consider them part of the EM problem.

EQ.1: On. The EM solver will look for potential elements that are eroded and remove them from the EM solve by updating its matrix system.

## *EM_CONTROL_MAGNET

Purpose: Control how often the magnetization vector for magnets is recomputed. By default, the magnetization vector for each magnet is computed only once at the beginning of the calculation (see Remark 1).

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | MCOMP | NCYCM |  |  |  |  |  |  |
| Type | 1 | 1 |  |  |  |  |  |  |
| Default | 0 | 0 |  |  |  |  |  |  |

## VARIABLE

MCOMP

NCYCM

Magnetization vector recomputation frequency. A value of 1 means recomputation at every EM time step. If a negative value is entered, $\mid$ NCYCM $\mid$ is the ID of a load curve giving the value as a function of time.

## Remark:

1. MCOMP. In most applications involving magnets, the magnetization vector associated to each magnet needs only to be calculated once at the beginning of the analysis. The magnetization direction is then scaled by the Coercive force value (See EM_PERMANENT_MAGNET) to correctly estimate the contribution of each magnet. In certain specific applications, for examples in cases involving magnet deformations or changes in magnet boundary conditions, it may be needed to periodically update the initially computed magnetization vector. This can be achieved by setting MCOMP to 1 and defining NCYCM.

## *EM_CONTROL_SOLUTION

Purpose: Specify different conditions under which LS-DYNA reassembles the FEM and BEM matrices.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | NCYLFEM | NCYLBEM | AUTOFEM | AUTOBEM | TOL1FEM | TOL2FEM | TOL1BEM | TOL2BEM |
| Type | I | I | I | I | F | F | F | F |
| Default | 5000 | 5000 | 0 | 0 | 0.3 | 0.1 | 0.3 | 0.1 |

## VARIABLE

NCYLFEM

NCYLBEM Number of electromagnetism cycles between the recalculation of BEM matrices. See Remark 1.

LT.O: |NCYLBEM| refers to a load curve, giving the number of cycles as a function of time.

AUTOFEM

AUTOBEM In addition to NCYLBEM, AUTOBEM triggers an automatic recomputation of the BEM matrices based on an error calculation of the conductors' relative displacements. See TOL1BEM and TOL2BEM. See Remark 2.

EQ.0: Autorecomputation off
EQ.1: Autorecomputation on

VARIABLE

TOL1FEM

TOL2FEM

TOL1BEM

TOL2BEM

## DESCRIPTION

Only used if AUTOFEM $=1$. If a conducting element experiences a deformation or a conductivity change that reaches an error larger than TOL1FEM, LS-DYNA reassembles the FEM matrices. If negative, the absolute value refers to a load curve that gives the tolerance as a function of time.

Only used if AUTOFEM $=1$. If (number of conducting elements) $\times$ TOL2FEM elements experience a deformation or a conductivity change that reaches an error larger than TOL2FEM, LS-DYNA recomputes the FEM matrices. If negative, the absolute value refers to a load curve that gives TOL2FEM as a function of time.

Only used if AUTOBEM = 1. If a conducting element experiences a displacement that reaches an error larger than TOL1BEM, then LS-DYNA reassembles the BEM matrices. If negative, the absolute value refers to a load curve that gives the tolerance as a function of time.

Only used if AUTOBEM $=1$. If (number of conducting elements) $\times$ TOL2BEM elements experience a displacement that reaches an error larger than TOL2BEM, then the BEM matrices will be recomputed. If negative, the absolute value refers to a load curve that gives TOL2BEM as a function of time.

## Remarks:

1. Order of precedence for NCYLFEM and NCYLBEM. You can set NCYLFEM and NCYLBEM with *EM_CONTROL_SOLUTION, *EM_SOLVER_FEM/BEM, or *EM_CONTROL. *EM_CONTROL_SOLUTION has the highest priority for setting these fields, while *EM_CONTROL has the lowest priority. If you leave one of these fields as the default value on *EM_CONTROL_SOLUTION, LS-DYNA then looks at the setting of that field on *EM_SOLVER_FEM/BEM. If left as the default on *EM_SOLVER_FEM/BEM, LS-DYNA looks at the setting on *EM_CONTROL.
2. EM contact with recomputation. When EM contact occurs and you have enabled automatic FEM or BEM matrix recomputation, LS-DYNA simultaneously reassembles both the FEM and BEM matrices when any criterion is reached to maintain consistency. When you do not enable automatic recomputation, we recommend the FEM and BEM systems have the same value for the recomputation cycle.

## *EM_CONTROL_SWITCH

Purpose: Create a control "switch" that will shut down the solver based on information from a load curve. LS-DYNA supports complex types of curves (See *DEFINE_CURVE_FUNCTION) that allow the setting up of complex on/off switches, for instance, by using a nodal temperature value.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | LCID | FEMCOMP | BEMCOMP |  |  |  |  |  |
| Type | I | 1 | 1 |  |  |  |  |  |
| Default | 0 | 0 | 0 |  |  |  |  |  |

## VARIABLE

LCID

FEMCOMP

BEMCOMP

## DESCRIPTION

Load curve or define curve function ID. Negative values switch the solver off; positive values switch it back on.

Flag to determine if LS-DYNA recomputes the FEM matrices each time the EM solver is turned back on:

EQ.0: Recompute FEM matrices.
EQ.1: Do not recompute FEM matrices.
Flag to determine if LS-DYNA recomputes the BEM matrices each time the EM solver is turned back on:

EQ.O: Recompute BEM matrices.
EQ.1: Do not recompute the BEM matrices.

## *EM_CONTROL_SWITCH_CONTACT

Purpose: It is possible to active a control "switch" that will shut down the electromagnetic contact detection. This can be useful in order to save some calculation time in cases where the user knows when contact between conductors will occur or stop occurring.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | LCID | NCYLFEM | NCYLFEM |  |  |  |  |  |
| Type | 1 | 1 | 1 |  |  |  |  |  |
| Default | 0 | 0 | 0 |  |  |  |  |  |

VARIABLE
LCID

NCYLFEM

NCYLBEM Determines the number of cycles before BEM matrix recomputation. If defined this will overwrite the previous NCYCLBEM as long as the contact detection is turned on.

## *EM_CONTROL_TIMESTEP

Purpose: Controls the EM time step and its evolution.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | TSTYPE | DTCONST | LCID | FACTOR | TSMIN | TSMAX | RLCSF | MECATS |
| Type | I | F | I | F | F | F | I | 1 |
| Default | none | none | none | 1.0 | none | none | 25 | 0 |

## VARIABLE

TSTYPE

## DESCRIPTION

Time step type:
EQ.1: Constant time step given in DTCONST
EQ.2: Time step as a function of time given by a load curve specified in LCID

EQ.3: Automatic time step computation, depending on the solver type. This time step is then multiplied by FACTOR
EQ.4: EM time step forced to be the same as the thermal time step in problems with the EM solver coupled to the thermal solver. In other words, the thermal solver determines the EM time step.

DTCONST Constant value for the time step for TSTYPE $=1$
LCID Load curve ID giving the time step as a function of time for TSTYPE $=2$

FACTOR
TSMIN

TSMAX

RLCSF $\quad$ RLC Circuit time step scale factor. See Remark 2.

## VARIABLE

MECATS

## DESCRIPTION

Mechanical time step handling in cases where the EM solver time step becomes smaller (see Remark 3):

EQ.0: Default. The EM time step will go below the solid mechanics timestep, and several EM solves will occur between two solid mechanics time steps to ensure time consistency.

EQ.1: The solid mechanics time step will adapt and decrease to the EM time step value so that only one EM solve occurs between two solid mechanics solves.

## Remarks:

1. Eddy Current Solver Automatic Time Step. For an eddy current solver, the automatic time step is based on the diffusion equation for the magnetic field:

$$
\sigma \frac{\partial \vec{A}}{\partial t}+\vec{\nabla} \times \frac{1}{\mu} \vec{\nabla} \times \vec{A}+\sigma \vec{\nabla} \varphi=\vec{\jmath}_{S}
$$

It is computed as the minimal elemental diffusion time step over the elements. For a given element, the elemental diffusion time step is given as $d t_{e}=l_{e}^{2} /(2 D)$, where:

- $D$ is the diffusion coefficient $D=1 /\left(\mu_{0} \sigma_{e}\right)$,
- $\sigma_{e}$ is the element electrical conductivity,
- $\mu_{0}$ is the permeability of free space,
- $l_{e}$ is the minimal edge length of the element (minimal size of the element).

2. Automatic Time Step with RLC Circuit. When an automatic time step is defined and an RLC circuit is present, the EM solver will perform an additional check and calculate an approximation of the first current period based on a 0-D circuit solve. It will then limit the timestep by a factor $T_{\text {period }} /(4 \times$ RLCSF $)$. The default value of RLCSF ensures that 25 EM timesteps will be calculated for the first quarter period.
3. MECATS. In general, we recommend avoiding scenarios where the EM time step becomes smaller than the solid mechanics time step which are often the result of ill-defined input decks and parameters. This can, however, happen in cases where conducting elements have high deformations and an automatic EM time step is selected in which case you can choose between the two MECATS options.

## *EM_DATABASE_CIRCUIT

Purpose: Enable the output of EM data for every circuit defined.
Output options card

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | OUTLV | DTOUT |  |  |  |  |  |  |
| Type | 1 | F |  |  |  |  |  |  |
| Default | 1 | 0. |  |  |  |  |  |  |

## VARIABLE

OUTLV

## DESCRIPTION

Determines if LS-DYNA creates the output file:
EQ.O: Do not generate the output file.
EQ.1: Generate the output file.
DTOUT Time interval to print the output. If DTOUT is equal to 0.0 , LSDYNA uses the EM timestep.

## Remarks:

1. File name. The file name for this database is em_circuit_ $X X X$.dat with $X X X$ as the circuit ID.
2. ResistanceD. LS-DYNA calculates ResistanceD in the following way:
a) Imposes a scalar potential difference of 1 at the circuit's boundaries SIDVIN and SIDVOUT.
b) Solves $\nabla^{2} \varphi=0$ with $\varphi_{\text {SIDvIN }}=1$ and $\varphi_{\text {SIDvout }}=0$ at SIDCURR. We do not consider diffusive effects, meaning the current density can be written as $\mathbf{j}=$ $\nabla \varphi$ and the total current as $I=\mathbf{j} \cdot \mathbf{n} d A$.
c) Estimates the resistance using $R_{D}=U / I . U$ is the potential difference between $\varphi_{\text {SIDVIN }}$ and and $\varphi_{\text {SIDVOUT }}$, that is, equal to 1 . The circuit's geometry and conductivity solely determine this $R_{D}$ resistance. It is, therefore, equivalent to the resistance as commonly defined in the circuit equations:

$$
R_{D}=L / \sigma S,
$$

where $L$ is the length of the circuit and $S$ is its surface area.
3. ResistanceJ. LS-DYNA calculates ResistanceJ with the data provided during the EM solve: $R_{J}=J / I^{2}$ where $J$ and $I$ are, respectively, the joule heating and the current. Compared with ResistanceD, ResistanceJ is not so much a resistance calculation since it accounts for the resistive effects (when using the Eddy current solver). Rather, it corresponds to the resistance that the circuit would need in order to get the same Joule heating in the context of a circuit equation. If all EM fields are diffused or you are using the resistive heating solver, ResistanceJ should be close to ResistanceD.
4. Mutual inductances. Only the mutual inductances between the first three circuits defined are output.

## *EM_DATABASE_CIRCUITOD

Purpose: Enable the output of 0D EM data for every circuit defined.
Output options card

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | OUTLV | DTOUT |  |  |  |  |  |  |
| Type | I | F |  |  |  |  |  |  |
| Default | 0 | 0. |  |  |  |  |  |  |

## VARIABLE

OUTLV

## DESCRIPTION

Determines if LS-DYNA creates the output file.
EQ.O: Does not generate the output file.
EQ.1: Generates the output file.
DTOUT Time interval to print the output. If DTOUT equals 0.0 , then LSDYNA uses the EM time step.

## Remarks:

1. File name. The file name for this database is em_circuitOD_ $X X X$.dat with $X X X$ as the circuit ID.
2. $\mathbf{O D}$ solution. At the start of the run, based on the initial values of the meshes resistances and inductances, the solver calculates the results for a so-called " 0 D " solution, which does not consider the current's diffusion, the part's displacements, or the EM material property changes. It is, therefore, a crude approximation. LS-DYNA extrapolates the initial results through time which it outputs at DTOUT intervals. This result can be useful in some cases, especially in RLC circuits if you need a rough idea of how the source current will behave.
3. Cost. Since the calculation of this 0 D circuit can take time depending on the size of the problem, you should only use it in cases where the output results help with comprehension of the analysis.
4. Effect of this keyword. This keyword does not influence the results of the EM run itself.
*EM_DATABASE_ELOUT
Purpose: Enable the output of EM data on elements.
Output Options Card.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | OUTLV | DTOUT |  |  |  |  |  |  |
| Type | 1 | F |  |  |  |  |  |  |
| Default | 0 | 0. |  |  |  |  |  |  |

Include as many cards as needed. This input ends at the next keyword ("*") card.

| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | ELSID |  |  |  |  |  |  |  |
| Type | 1 |  |  |  |  |  |  |  |
| Default | none |  |  |  |  |  |  |  |

VARIABLE
OUTLV

## DESCRIPTION

Determines if LS-DYNA creates the output file:
EQ.O: Does not generate the output file.
EQ.1: Generates the output file.

DTOUT Time interval to print the output. If DTOUT equals 0.0, LS-DYNA uses the EM time step.

ELSID
Solid elements set ID

## Remarks:

1. File name. The file name for this database is em_elout.dat.

## *EM_DATABASE_FIELDLINE

Purpose: The EM solver uses a BEM method to calculate the EM fields between conductors. With this method, the magnetic field in the air or vacuum between conductors is therefore not explicitly computed. However, in some cases, it may be interesting to visualize some magnetic field lines for a better analysis. This keyword allows the output of field line data. It has no influence on the results of the EM solve.

Output Options Card.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | FLID | PSID | DTOUT | NPOINT |  |  |  |  |
| Type | I | I | F | I |  |  |  |  |
| Default | none | none | 0. | 100 |  |  |  |  |

## Remaining cards are optional. ${ }^{\dagger}$

| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | INTEG | H | HMIN | HMAX | TOLABS | TOLREL |  |  |
| Type | I | F | F | F | F | F |  |  |
| Default | 2 | 0. | 0. | 1 E 10 | $1 \mathrm{E}-3$ | $1 \mathrm{E}-5$ |  |  |


| Card 3 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | BTYPE |  |  |  |  |  |  |  |
| Type | 1 |  |  |  |  |  |  |  |
| Default | 2 |  |  |  |  |  |  |  |

## VARIABLE

FLID

PSID

DTOUT

NPOINT

INTEG

H

HMIN

HMAX

TOLABS

TOLREL

BTYPE

## DESCRIPTION

Field line set ID
Point Set ID associated to the field line set (See *EM_POINT_SET). The coordinates given by the different points will be the starting points of the field lines.

Time interval to print the output. If DTOUT is equal to 0.0 , then the EM time step will be used.

Number of points per field line. The points are regularly spaced.
Type of numerical integrator used to compute the field lines :
EQ.1: RK4, Runge Kutta 4. See Remark 2.
EQ.2: DOP853, Dormand Prince 8(5,3). See Remark 2.
Value of the step size. In case of an integrator with adaptive step size, it is the initial value of the step size.

Minimal step size value. Only used in the case of an integrator with adaptive step size.

Maximal step size value. Only used in the case of an integrator with adaptive step size.

Absolute tolerance of the integrator. Only used in the case of an integrator with adaptive step size.

Relative tolerance of the integrator. Only used in the case of an integrator with adaptive step size.

Method to compute the magnetic field :
EQ.1: Direct method (every contribution is computed by the Biot Savart Law and summed up : very slow).

EQ.2: Multipole method (approximation of the direct method using the multipole expansion).

EQ.3: Multicenter method (approximation of the direct method using a weighted subset of points only in order to compute the magnetic field).

## Remarks:

1. File Names. The file name for this database is em_fieldLine_XX_YYY.dat where $X X$ is the field line ID and $Y Y Y$ is the point set ID defined in *EM_POINT_SET.
2. Integrators. The Runge Kutta 4 integrator is an explicit iterative method for solving ODEs. It is a fourth order method with a constant step size. The Dormand Prince $8(5,3)$ integrator is an explicit iterative method for solving IDEs. Particularly, this integrator is an embedded Runge Kutta integrator of order 8 with an adaptive step size. This integrator allows a step size control which is done though an error estimate at each step. The Dormand Prince $8(5,3)$ is a Dormand Prince 8(6) for which the $6^{\text {th }}$ order error estimator has been replaced by a $5^{\text {th }}$ order estimator with $3^{\text {rd }}$ order correction in order to make the integrator more robust.
*EM_DATABASE_GLOBALENERGY
Purpose: Enable the output of global EM energies.
Output Options Card.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | OUTLV | DTOUT |  |  |  |  |  |  |
| Type | I | F |  |  |  |  |  |  |
| Default | 0 | 0. |  |  |  |  |  |  |

## VARIABLE

## DESCRIPTION

OUTLV
Determines if the output file should be dumped.
EQ.O: No output file is generated.
EQ.1: The output file is generated.

DTOUT Time interval to print the output. If DTOUT is equal to 0.0 , then the EM timestep will be used.

## Remarks:

1. Database Name. The file name for this database is em_globEnergy.dat.
2. Output Data. The output file includes the global EM energies of the mesh, the air, and the source circuit. It also includes the global kinetic energy and the global plastic work energy.

## *EM_DATABASE_NODOUT

Purpose: This keyword enables the output of EM data on nodes.
Output Options Card.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | OUTLV | DTOUT |  |  |  |  |  |  |
| Type | I | F |  |  |  |  |  |  |
| Default | 0 | 0. |  |  |  |  |  |  |

Include as many cards as needed. This input ends at the next keyword ("*") card.

| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | NSID |  |  |  |  |  |  |  |
| Type | 1 |  |  |  |  |  |  |  |
| Default | none |  |  |  |  |  |  |  |

## VARIABLE

OUTLV

## DESCRIPTION

Determines if the output file should be dumped.
EQ.O: No output file is generated.
EQ.1: The output file is generated.
DTOUT Time interval to print the output. If DTOUT is equal to 0.0 , then the EM timestep will be used.

NSID Node Set ID.

## Remarks:

1. The file name for this database is em_nodout.dat.
*EM_DATABASE_PARTDATA
Purpose: This keyword enables the output of EM data for every part defined. .

Output Options Card.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | OUTLV | DTOUT |  |  |  |  |  |  |
| Type | I | F |  |  |  |  |  |  |
| Default | 0 | 0. |  |  |  |  |  |  |

## VARIABLE

## DESCRIPTION

OUTLV
Determines if the output file should be dumped.
EQ.O: No output file is generated.
EQ.1: The output file is generated.

DTOUT Time interval to print the output. If DTOUT is equal to 0.0 , then the EM timestep will be used.

## Remarks:

1. The file name for this database is em_partData_ $X X X$.dat with $X X X$ the part ID.
2. Outputs the part EM energies of the part as well as the Lorentz force. Also outputs the part kinetic energy and the part plastic work energy.

## *EM_DATABASE_POINTOUT

Purpose: Enable the output of EM data on point sets.
Output Options Card.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | OUTLV | DTOUT |  |  |  |  |  |  |
| Type | 1 | F |  |  |  |  |  |  |
| Default | 0 | 0. |  |  |  |  |  |  |

Include as many cards as needed. This input ends at the next keyword ("*") card.

| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | PTSID |  |  |  |  |  |  |  |
| Type | 1 |  |  |  |  |  |  |  |
| Default | none |  |  |  |  |  |  |  |

VARIABLE
OUTLV

## DESCRIPTION

Determines if LS-DYNA creates the output file:
EQ.O: Do not generate the output file.
EQ.1: Generate the output file.
DTOUT Time interval to print the output. If DTOUT equals 0.0, LS-DYNA uses the ICFD time step.

PTSID Point set ID (See *EM_POINT_SET card).

## Remarks:

1. File name. The file name for this database is em_pointout.dat.
*EM_DATABASE_ROGO
Purpose: This keyword enables the output of EM data for every circuit defined. .
Output Options Card.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | OUTLV | DTOUT |  |  |  |  |  |  |
| Type | I | F |  |  |  |  |  |  |
| Default | 1 | 0. |  |  |  |  |  |  |

## VARIABLE

## DESCRIPTION

OUTLV
Determines if the output file should be dumped.
EQ.O: No output file is generated.
EQ.1: The output file is generated.
DTOUT Time interval to print the output. If DTOUT is equal to 0.0 , then the EM timestep will be used.

## Remarks:

1. The file name for this database is em_rogoCoil_ $X X X$.dat where $X X X$ is the rogo Coil ID.

## *EM_DATABASE_TIMESTEP

Purpose: This keyword enables the output of EM data regarding the EM timestep.
Output options card.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | OUTLV |  |  |  |  |  |  |  |
| Type | 1 |  |  |  |  |  |  |  |
| Default | 0 |  |  |  |  |  |  |  |

## VARIABLE

OUTLV

DESCRIPTION
Determines if the output file should be dumped.
EQ.O: No output file is generated.
EQ.1: The output file is generated.

## Remarks:

1. The file name for this database is em_timestep.dat.
2. Outputs the run's EM tim estep versus the time step calculated using the EM CFL condition as criteria (autotimestep). This can be useful in cases with big deformations and/or material property changes and a fixed time step is being used in case that time step becomes to big compared to the stability time step.

## *EM_EP_CELLMODEL_DEFINE_FUNCTION

Purpose: Define a user-defined ionic cell model for Electro-Physiology.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | MID | NSTATE | FSWITCH |  |  |  |  |  |
| Type | 1 | 1 | 1 |  |  |  |  |  |
| Default | none | none | none |  |  |  |  |  |


| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | DVDT | DU1DT | DU2DT | DU3DT | DU4DT | DU5DT | DU6DT | DU7DT |
| Type | 1 | 1 | 1 | 1 | 1 | 1 | 1 | 1 |
| Default | none | none | none | none | none | none | none | none |


| Card 3 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | V0 | U 1 | U 2 | U 3 | U 4 | U 5 | U 6 | U 7 |
| Type | I | I | I | I | I | I | I | I |
| Default | none | none | none | none | none | none | none | none |

VARIABLE
MID

FSWITCH

## DESCRIPTION

Material ID defined in the *MAT section
Flag for the ODE definition (see Remark 1):
EQ.0: Functions
EQ.1: Derivatives

## VARIABLE

NSTATE

DVDT

DUiDT

V0
$\mathrm{U} i$

## DESCRIPTION

Number of state variables $u_{1}, u_{2}, \ldots, u_{n}$. The maximum value is 7 (see Cards 2 and 3).

Function ID (see *DEFINE_FUNCTION) for evolution of $V$ (function $g$ in the equations in Remark 1).

Function ID (see *DEFINE_FUNCTION) for evolution of $u_{i}$ (function $f_{i}$ in the equations in Remark 1)

Function ID (see *DEFINE_FUNCTION) for initial value of $V(x, y, z)$

Function ID (see *DEFINE_FUNCTION) for initial value of $u_{i}(x, y, z)$

## Remarks:

1. Model. This keyword enables specifying a user-defined cell model through defined functions (see *DEFINE_FUNCTION). This model traces the evolution of the transmembrane potential, $V$, and $n$ state variables: $u_{1}, u_{2}, \ldots u_{n}$. Their temporal evolution depends upon FSWITCH.
a) If FSWITCH $=0$ :

$$
\begin{aligned}
& V(t)=g\left(t, d t, V(t-1), u_{1}(t-1), u_{2}(t-1), \ldots, u_{n}(t-1)\right) \\
& u_{1}(t)=f_{1}\left(t, d t, V(t-1), u_{1}(t-1), u_{2}(t-1), \ldots, u_{n}(t-1)\right) \\
& u_{2}(t)=f_{2}\left(t, d t, V(t-1), u_{1}(t-1), u_{2}(t-1), \ldots, u_{n}(t-1)\right) \\
& \vdots \\
& u_{n}(t)=f_{n}\left(t, d t, V(t-1), u_{1}(t-1), u_{2}(t-1), \ldots, u_{n}(t-1)\right)
\end{aligned}
$$

b) If FSWITCH $=1$ :

$$
\begin{gathered}
\frac{\partial V(t)}{\partial t}=g\left(t, d t, V(t-1), u_{1}(t-1), u_{2}(t-1), \ldots, u_{n}(t-1)\right) \\
\frac{\partial u_{1}(t)}{\partial t}=f_{1}\left(t, d t, V(t-1), u_{1}(t-1), u_{2}(t-1), \ldots, u_{n}(t-1)\right) \\
\frac{\partial u_{2}(t)}{\partial t}=f_{2}\left(t, d t, V(t-1), u_{1}(t-1), u_{2}(t-1), \ldots, u_{n}(t-1)\right) \\
\vdots \\
\frac{\partial u_{n}(t)}{\partial t}=f_{3}\left(t, d t, V(t-1), u_{1}(t-1), u_{2}(t-1), \ldots, u_{n}(t-1)\right)
\end{gathered}
$$

2. Benchmarks. You can use this model to perform the electrophysiology benchmarks presented in Pathmanathan and Gray [2014].

## References:

[1] Pathmanathan, P. and R.A. Gray, "Verification of computational models of cardiac electro-physiology," Int J Numer Method Biomed Eng, vol. 30, No. 5, pp. 524-544, (2014).

## *EM_EP_CELLMODEL_FENTONKARMA

Purpose: Define a Fenton-Karma ionic cell model for Electro-Physiology.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | MATID |  |  |  |  |  |  |  |
| Type | I |  |  |  |  |  |  |  |
| Default | none |  |  |  |  |  |  |  |


| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | TAUD | TAUR | TAUSI | TAU0 | TAUVP | TAUVM | TAUWP | TAUWM |
| Type | F | F | F | F | F | F | F | F |
| Default | none | none | none | none | none | none | none | none |


| Card 3 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | UC | UCSI | K |  |  |  |  |  |
| Type | F | F | F |  |  |  |  |  |
| Default | none | none | none |  |  |  |  |  |


| Card 4 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | U0 | V0 | W0 |  |  |  |  |  |
| Type | F | F | F |  |  |  |  |  |
| Default | none | none | none |  |  |  |  |  |

```
VARIABLE
```

MATID
TAUD

TAUR

TAUSI
TAU0

TAUVP

TAUVM Time constant $\tau_{v m}$ described in Equation 3
TAUWP Time constant $\tau_{w p}$ described in Equation 4
TAUWM $\quad$ Time constant $\tau_{w m}$ described in Equation 4
UC Threshold potential, $u_{c}$ for activation of $J_{\text {fi }}$ (the fast inward current) in Equations 3, 4, 5, and 6

UCSI Threshold potential $u_{c}^{\text {si }}$ for activation of $J_{\text {si }}$ (the slow inward current) in Equation 7

K Constant $k$ in Equation 7
U0/V0/W0 Initial values of $u, v$, and $w$, respectively

## Remarks:

```
The Fenton-Karma model is a simplified ionic model with three membrane currents that approximates well the restitution properties and spiral wave behavior of more complex ionic models of cardiac action potential (Beeler-Reuter and others). It was introduced in [1].
```

The total current flowing through the membrane is given by:

$$
\begin{equation*}
I_{\mathrm{ion}}=-C_{m} \frac{\partial V}{\partial t}=-J_{\mathrm{fi}} \tag{1}
\end{equation*}
$$

where $V$ is the transmembrane potential, $C_{m}$ is the specific capacitance of the cell membrane, and $J_{\mathrm{fi}}$ is the fast inward current.

The model depends on three state variables, $u, v$, and $w$, and three membrane currents, $J_{\mathrm{fi}} J_{\mathrm{so}}$ (slow outward current), and $J_{\mathrm{si}}$ (slow inward current), through the following equations:

$$
\begin{gather*}
\frac{d u}{d t}=-J_{\mathrm{fi}}-J_{\mathrm{so}}-J_{\mathrm{si}}  \tag{2}\\
\frac{d v}{d t}=\frac{\Theta\left(u_{c}-u\right)(1-v)}{\tau_{v m}}-\frac{\Theta\left(u-u_{c}\right) v}{\tau_{v p}}  \tag{3}\\
\frac{d w}{d t}=\frac{\Theta\left(u_{c}-u\right)(1-w)}{\tau_{w w m}}-\frac{\Theta\left(u-u_{c}\right) w}{\tau_{w p}}  \tag{4}\\
J_{\mathrm{fi}}=-\frac{\Theta\left(u_{c}-u\right)(1-u)\left(u-u_{c}\right)}{\tau_{d}}  \tag{5}\\
J_{\mathrm{so}}=\frac{u \Theta\left(u_{c}-u\right)}{\tau_{o}}+\frac{u \Theta\left(u-u_{c}\right)}{\tau_{r}}  \tag{6}\\
J_{\mathrm{si}}=-\frac{w\left(1+\tanh \left[k\left(u-u_{c}^{s i}\right]\right)\right.}{2 \tau_{s i}} \tag{7}
\end{gather*}
$$

In the above $\Theta$ is the Heaviside step function.

## References:

[1] Fenton, F. \& A. Karma, "Vortex dynamics in three-dimensional continuous myocardium with fiber rotation. Filament instability and fibrillations," Chaos, Solitons, and Fractals, Vol. 8, No. 1, pp. 661686, (1998).
[2] https://www.ibiblio.org/e-notes/html5/fk.html

## *EM_EP_CELLMODEL_FITZHUGHNAGUMO

Purpose: Define a Fitzhugh-Nagumo ionic cell model for Electro-Physiology.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | MATID |  |  |  |  |  |  |  |
| Type | I |  |  |  |  |  |  |  |
| Default | none |  |  |  |  |  |  |  |


| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | ALPHA | BETA | GAMMA | C | MU1 | MU2 |  |  |
| Type | F | F | F | F | F | F |  |  |
| Default | none | none | none | none | none | none |  |  |


| Card 3 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | V | R |  |  |  |  |  |  |
| Type | F | F |  |  |  |  |  |  |
| Default | none | none |  |  |  |  |  |  |

VARIABLE
MATID

ALPHA

BETA
GAMMA

C

DESCRIPTION
Material ID defined in *MAT

Excitation constant $\alpha$ described in Equation 1
Excitation constant $\beta$ described in Equation 2
Excitation constant $\gamma$ described in Equation 2
Excitation constant $c$ described in Equation 1

## VARIABLE

MU1
MU2 Excitation constant $\mu_{2}$ described in Equation 2
V Initial value of $V$
R Initial value of $r$

## Remarks:

In the Fitzhugh-Nagumo model, the excitation is defined by a cubic polynomial along with one recovery variable, $r$. The transmembrane current, $I_{\text {ion }}$, is given by:

$$
\begin{equation*}
I_{\text {ion }}=-C_{m} \frac{\partial V}{\partial t}=-c V(V-\alpha)(V-1)-r V \tag{1}
\end{equation*}
$$

Here $V$ is the transmembrane potential, $C_{m}$ is the specific capacitance of the cell membrane, and $c$ and $\alpha$ are excitation constants.

The recovery variable $r$ evolves according to:

$$
\begin{equation*}
\frac{d r}{d t}=\left(\gamma+\frac{r \mu_{1}}{\mu_{2}+V}\right)(-r-c V(V-\beta-1)) \tag{2}
\end{equation*}
$$

where $\beta, \gamma, \mu_{1}$ and $\mu_{2}$ are excitation constants.

## References:

[1] Aliev, R.R. and Panfilov, A.V., "A simple two-variable model of cardiac excitation," Chaos, Solitons, and Fractals, Vol 7, No 3, pp 293-301, (1996).
[2] Pullan, A.J., Cheng, L.K., and Buist, M.L., Mathematically Modelling the Electrical Activity of the Heart, World Scientific Publishing Co. Pte. Ltd., Singapore, pp 132-133, (2005).
[3] Baillargeon, B. et al., "The Living Heart Project: A robust and integrative simulator for human heart function," European Journal of Mechanics - A/Solids. Vol 48, pp 38-47, (2014).

## *EM_EP_CELLMODEL_TENTUSSCHER

Purpose: Define a ten Tusscher ionic cell model for Electro-Physiology.

## Card Summary:

Card 1. This card is required.

| MID |  |  |  |  |  |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

Card 2. This card is required.

| $R$ | $T$ | $F$ | CM | VC | VSR | VSS | PKNA |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |

Card 3. This card is required.

| KO | NAO | CAO |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |

Card 4. This card is required.

| GK1 | GKR | GKS | GNA | GBNA | GCAL | GBCA | GT0 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |

Card 5. This card is required.

| GPCA | GPK |  |  |  |  |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

Card 6. This card is required.

| PNAK | KMK | KMNA | KNACA | KSAT | ALPHA | GAMMA | KMCA |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |

Card 7. This card is required.

| KMNAI | KPCA |  |  |  |  |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

Card 8. This card is required.

| K1 | K2 | K3 | K4 | EC | MAXSR | MINSR |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |

Card 9. This card is required.

| VREL | VLEAK | VXFER | VMAXUP | KUP |  |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

Card 10. This card is required.

| BUFC | KBUFC | BUFSR | KBUFSF | BUFSS | KBUFSS |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

Card 11. This card is required.

| $V$ | KI | NAI | CAI | CASS | CASR | RPRI |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |

Card 12. This card is required.

| XR1 | XR2 | XS | M | H | J | D | F |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

Card 13. This card is required.

| F2 | FCASS | S | R |  |  |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

## Data Card Definitions:

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | MID |  |  |  |  |  |  |  |
| Type | 1 |  |  |  |  |  |  |  |
| Default | none |  |  |  |  |  |  |  |

VARIABLE
MID Material ID defined in *MAT section

| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | R | T | F | CM | VC | VSR | VSS | PKNA |
| Type | F | F | F | F | F | F | F | F |
| Default | none | none | none | none | none | none | none | none |

## VARIABLE

## DESCRIPTION

R
Gas constant $\left(\mathrm{J} \times \mathrm{K}^{-1} \times \mathrm{Mol}^{-1}\right)$
T
Temperature (K)

## VARIABLE

F

CM
VC
VSR
VSS
PKNA

Faraday constant $\left(\mathrm{C} \times \mathrm{mmol}^{-1}\right)$
Cell capacitance for unit surface area $\left(\mu \mathrm{F} \times \mathrm{Cm}^{-2}\right)$
Cytoplasmic volume ( $\mu \mathrm{m}^{3}$ )
Sarcoplasmic reticulum volume ( $\mu \mathrm{m}^{3}$ )
Subspace volume ( $\mu \mathrm{m}^{3}$ )
Relative $I_{\text {Ks }}$ permeability to $\mathrm{Na}^{+}$

| Card 3 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | KO | NAO | CAO |  |  |  |  |  |
| Type | F | F | F |  |  |  |  |  |
| Default | none | none | none |  |  |  |  |  |

VARIABLE
КО
$\mathrm{NAO} \quad$ Extracellular $\mathrm{Na}^{+}$concentration $(\mathrm{mM})$
CAO Extracellular $\mathrm{Ca}^{2+}$ concentration (mM)

| Card 4 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | GK1 | GKR | GKS | GNA | GBNA | GCAL | GBCA | GT0 |
| Type | F | F | F | F | F | F | $F$ | $F$ |
| Default | none | none | none | none | none | none | none | none |


| Card 5 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | GPCA | GPK |  |  |  |  |  |  |
| Type | F | F |  |  |  |  |  |  |
| Default | none | none |  |  |  |  |  |  |

VARIABLE
GK1, GKR, GKS, GNA, GBNA, GCAL, GBCA, GTO, GPCA, GPK

| Card 6 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | PNAK | KMK | KMNA | KNACA | KSAT | ALPHA | GAMMA | KMCA |
| Type | F | F | F | F | F | $F$ | $F$ | $F$ |
| Default | none | none | none | none | none | none | none | none |


| Card 7 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | KMNAI | KPCA |  |  |  |  |  |  |
| Type | F | F |  |  |  |  |  |  |
| Default | none | none |  |  |  |  |  |  |

## VARIABLE

PNAK

## DESCRIPTION

$P_{\mathrm{NaK}}$, parameter for calculating the $\mathrm{Na}^{+} / \mathrm{K}^{+}$pump current (units: $\mathrm{pA} \times \mathrm{pF}^{-1}$ ). See Reference [1].

VARIABLE
KMK,
KMNA
KNACA,
KSAT, AL-
PHA, GAM-
MA, KMNAI
KPCA

## DESCRIPTION

$K_{\mathrm{mK}}$ and $K_{\mathrm{mNa}}$, parameters for calculating the $\mathrm{Na}^{+} / \mathrm{K}^{+}$pump current (units: millimolar). See Reference [1].
$k_{\mathrm{NaCa}}, k_{\text {sat }}, \alpha, \gamma$, and $K_{\mathrm{mNai}}$, parameters for calculating the $\mathrm{Na}^{+} / \mathrm{Ca}^{2+}$ exchanger current (units: millimolar). See Reference [1].

| Card 8 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | K1 | K2 | K3 | K4 | EC | MAXSR | MINSR |  |
| Type | F | F | F | F | F | F | F |  |
| Default | none | none | none | none | none | none | none |  |
| Ref | 2 | 2 | 2 | 2 | 2 | 2 | 2 |  |

VARIABLE
K1
K2
K3

EC
MAXSR/MI
NSR

K4 I to O and RI to I $I_{\text {rel }}$ transition rate $\left(\mathrm{ms}^{-1}\right)$
$\mathrm{Ca}_{\text {SR }}$ half-saturation constant of $k_{\text {casr }}(\mathrm{mM})$

## DESCRIPTION

R to O and RI to I $I_{\text {rel }}$ transition rate $\left(\mathrm{mM}^{-2} \times \mathrm{ms}^{-1}\right)$
O to I and R to $\mathrm{RI} I_{\text {rel }}$ transition rate $\left(\mathrm{mM}^{-1} \times \mathrm{ms}^{-1}\right)$
O to R and I to RI $I_{\text {rel }}$ transition rate $\left(\mathrm{ms}^{-1}\right)$

Maximum and minimum values of $k_{\text {casr }}$

| Card 9 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | VREL | VLEAK | VXFER | VMAXUP | KUP |  |  |  |
| Type | F | F | F | F | F |  |  |  |
| Default | none | none | none | none | none |  |  |  |

VARIABLE
VREL,
VLEAK,
VXFER,
VMAXUP
KUP

## DESCRIPTION

Maximal $I_{\text {rel }}, I_{\text {leak }}, I_{\text {xfer }}$, and $I_{\text {up }}$ conductance $\left(\mathrm{mM} \times \mathrm{ms}^{-1}\right)$, respectively. See Reference [2].

Half-saturation constant of $I_{\text {up }}(\mathrm{mM})$. See Reference [2].

| Card 10 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | BUFC | KBUFC | BUFSR | KBUFSF | BUFSS | KBUFSS |  |  |
| Type | F | F | F | F | F | F |  |  |
| Default | none | none | none | none | none | none |  |  |
| Ref | 2 | 2 | 2 | 2 | 2 | 2 |  |  |

## VARIABLE

BUFC
KBUFC
BUFSR
KBUFSR
BUFSS
KBUFSS Cass half-saturation constant for subspace buffer (mM)

| Card 11 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | V | KI | NAI | CAI | CASS | CASR | RPRI |  |
| Type | F | F | F | F | F | F | F |  |
| Default | none | none | none | none | none | none | none |  |
| Ref | 2 | 1 | 1 | 2 | 2 | 2 | 2 |  |

## VARIABLE

KI Initial value of $\mathrm{K}_{\mathrm{i}}$, used in potassium dynamics (mM)
NAI Initial value of $\mathrm{Na}_{\mathrm{i}}$, used in sodium dynamics (mM)
CAI Initial value of $\mathrm{Ca}_{\mathrm{i}}$. used in calcium dynamics ( mM )
CASS Initial value of Cass, used in calcium dynamics (mM)
CASR Initial value of CasR, used in calcium dynamics (mM)
RPRI Initial value of $\mathrm{R}^{\prime}$, used in calcium dynamics

| Card 12 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | XR1 | XR2 | XS | M | H | J | D | F |
| Type | F | F | F | F | F | F | F | F |
| Default | none | none | none | none | none | none | none | none |
| Ref | 1 | 1 | 1 | 1 | 1 | 1 | 2 | 2 |

## VARIABLE

XR1

## DESCRIPTION

Initial value of $x_{r 1}$, used to compute the rapid time dependent $\mathrm{K}^{+}$ current

| VARIAB |  | DESCRIPTION |  |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| XR2 |  | Initial value of $x_{r 2}$, used to compute the rapid time dependent $\mathrm{K}^{+}$ current |  |  |  |  |  |  |
| XS |  | Initial value of $x_{s}$, used to compute slow time dependent $\mathrm{K}^{+}$current |  |  |  |  |  |  |
| M |  | Initial value of $m$, used to compute the fast $\mathrm{Na}^{+}$current |  |  |  |  |  |  |
| H |  | Initial value of $h$, used to compute the fast $\mathrm{Na}^{+}$current |  |  |  |  |  |  |
| J |  | Initial value of $j$, used to compute the fast $\mathrm{Na}^{+}$current |  |  |  |  |  |  |
| D |  | Initial value of $d$, used to compute the L-type $\mathrm{Ca}^{2+}$ current |  |  |  |  |  |  |
| F |  | Initial value of $f$, used to compute the L-type $\mathrm{Ca}^{2+}$ current |  |  |  |  |  |  |
| Card 13 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| Variable | F2 | FCASS | S | R |  |  |  |  |
| Type | F | F | F | F |  |  |  |  |
| Default | none | none | none | none |  |  |  |  |
| Ref | 2 | 2 | 1 | 1 |  |  |  |  |

## VARIABLE

F2
FCASS

S

R

## DESCRIPTION

Initial value of $f_{2}$, used to compute the L-type $\mathrm{Ca}^{2+}$ current Initial value of $f_{\text {cass }}$, used to compute the L-type $\mathrm{Ca}^{2+}$ current Initial value of $s$, used to compute the transient outward current Initial value of $r$, used to compute the transient outward current

## Remarks:

This is a model of the action potential of human ventricular cells that, while including a high level of electrophysiological detail, is computationally cost-effective enough to be applied in large-scale spatial simulations for the study of reentrant arrhythmias. Please see the references for details. This model is based on [2].

## References:

[1] ten Tusscher, K.H.W.J., D. Noble, P.J. Noble, and A.V. Panfilov, "A model for human ventricular tissue," Am J Physiol Heart Circ Physiol, vol 286, no 4, pp H1573-H1589, (2004).
[2] ten Tusscher, K.H.W.J. and A.V. Panfilov, "Alternans and spiral breakup in human ventricular tissue model," Am J Physiol Heart Circ Physiol, vol 291, no 3, pp H1088-H1100, (2006).

## *EM_EP_CELLMODEL_TOMEK

Purpose: Define a ToR-ORd model for cardiac electrophysiology [1].

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | MID | PHIEND | PHIMY |  |  |  |  |  |
| Type | I | F | F |  |  |  |  |  |
| Default | none | 0.0 | 0.0 |  |  |  |  |  |

## VARIABLE

MID
PHIEND

PHIMYO

## DESCRIPTION

Material ID: refers to MID in the *PART card.

Value between 0.0 and 1.0 giving the ratio of the cardiac tissue to be considered endocardial in the ToR-Ord cell model.

Value between 0.0 and 1.0 giving the ratio of the cardiac tissue to be considered myocardial in the ToR-Ord cell model.

## References:

[1] Tomek J., Bueno-Orovio A., Passini E., Zhou X., Minchole A., Britton O., Bartolucci C., Severi S., Shrier A., Virag L., Varro A., and Rodriguez B., "Development, calibration, and validation of a novel human ventricular myocyte model in health, disease, and drug block," Elife (2019).

## *EM_EP_CELLMODEL_USERMAT

Purpose: Specify a user material for an ionic cell model to be used in electrophysiology simulations.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | MID |  |  |  |  |  |  |  |
| Type | 1 |  |  |  |  |  |  |  |

VARIABLE
MID

DESCRIPTION
Material ID. A unique number must be specified (see *PART).

## *EM_EP_CREATEFIBERORIENTATION

Purpose: Define fiber orientation by solving a Laplace-Dirichlet system defined by *EM_EP_FIBERINITIAL. This feature is based on [1].

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | PSID | LDID1 | LDID2 | ALPHA | BETA | IEXPORT | IPRERUN |  |
| Type | I | I | I | F | F | 1 | I |  |
| Default | none | none | none | 0.0 | 0.0 | 0 | 0 |  |

## VARIABLE

PSID
LDID1 ID of the Laplace system that is solved in the transmural direction
LDID2 ID of the Laplace system that is solved in the apicobasal direction
ALPHA Helical angle with respect to the counterclockwise circumferential direction in the heart when looking from the base towards the apex.

LT.O: |ALPHA | is the ID for the *DEFINE_FUNCTION giving the helical angle. See Remark 1 for available arguments.

BETA Angle with respect to the outward transmural axis of the heart.
LT.O: |BETA | is the ID for the *DEFINE_FUNCTION giving the angle. See Remark 1 for available arguments.

IEXPORT Selects whether result files (ELEMENT_SOLID_ORTHO.k and vtk files) are exported:

EQ.O: Not exported
EQ.1: Exported
IPRERUN

## DESCRIPTION

Part set ID of the part set on which the system is solved

Select whether the run is stopped after creating fibers:

EQ.0: Do not stop after fiber creation
EQ.1: Stop after fiber creation

## Remarks:

1. *DEFINE_FUNCTION Arguments. The arguments for the function defined with *DEFINE_FUNCTION may include the following: f(x_ele, y_ele, z_ele, phi_len, phi_thi). Here phi_len and phi_thi are the potentials corresponding to the potentials solved in LDID1 and LDID2, respectively.

## References:

[1] Bayer, J.D., Blake, R. C., Plank, G., and Trayanova, N. A., "A novel rule-based algorithm for assigning myocardial fiber orientation to computational heart models," Annals of biomedical engineering, 40(10), 2243-2254 (2012).

## *EM_EP_ECG

Purpose: Compute pseudo-ECGs on a set of virtual points. LS-DYNA exports a file named em_ECG_\{ECGID\}.dat.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | ECGID | PSID |  |  |  |  |  |  |
| Type | 1 | 1 |  |  |  |  |  |  |
| Default | none | none |  |  |  |  |  |  |
| VARIABLE |  |  |  |  |  |  |  |  |

ECGID
PSID

ID of the ECG computation
Point set ID containing the list of virtual points on which the pseudo-ECGs are computed

## *EM_EP_FIBERINITIAL

Purpose: Define the boundary conditions of the Laplace-Dirichlet system to be solved to define fiber orientation, based on [1]. The system is solved by using *EM_EP_CREATEFIBERORIENTATION.

Include as many of this card as needed. The next keyword ("*") card terminates this input.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | LDID | PID | STYPE | SID1 | SID0 |  |  |  |
| Type | 1 | 1 | 1 | 1 | 1 |  |  |  |
| Default | none | none | none | none | none |  |  |  |

## VARIABLE

## DESCRIPTION

LDID

PID Part ID on which the system is solved

Set type for the boundary condition:
EQ.1: Segment set
EQ.2: Node set

SID1

SID0

Set on which a potential of value 1 is prescribed
Set on which a potential of value 0 is prescribed

## References:

[1] Bayer, J.D., Blake, R. C., Plank, G., and Trayanova, N. A., "A novel rule-based algorithm for assigning myocardial fiber orientation to computational heart models," Annals of biomedical engineering, 40(10), 2243-2254 (2012).

## *EM_EP_PURKINJE_NETWORK

Purpose: Define a Purkinje network that consists of conductive beams and lies on a given surface, based on Costabal et al [2016].

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | PURKID | BUILDNET | SSID | MID | POINTSTX | POINTSTY | POINTSTZ | EDGELEN |
| Type | I | I | I | I | F | F | F | F |


| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | NGEN | NBRINIT | NSPLIT | INODEID | IEDGEID |  |  |  |
| Type | 1 | 1 | 1 | 1 | 1 |  |  |  |

## VARIABLE

PURKID
BUILDNET

SSID

MID
POINTSTX

POINTSTY

POINTSTZ
EDGELEN
NGEN

NBRINIT

Segment set on which the Purkinje network is lying
Material ID defined in the *MAT section.
$X$ coordinate of the tree origin
$Y$ coordinate of the tree origin
Z coordinate of the tree origin
Edge length
Number of generations of branches
Number of branches attached to the tree origin


Figure 6-3. Example of part of a network. For this network, NSPLIT $=4$.

## VARIABLE

NSPLIT

INODEID
IEDGEID

## DESCRIPTION

Number of nodes between two consecutive branchings as shown in Figure 6-3

Initial node ID
Initial edge ID. These edges are internally generated by LS-DYNA.

## References:

Costabal, F.S., D. E. Hurtado, and E. Kuhl, "Generating Purkinje networks in the human heart," Journal of Biomechanics, vol. 49, issue 12, pp. 2455-2465, (2016).

## *EM_EP_TENTUSSCHER_STIMULUS

Purpose: Define a stimulation pattern of a Tentusscher cell model (requires the use of *EM_EP_CELLMODEL_TENTUSSCHER).

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | STIMID | SETTYPE | SETID |  |  |  |  |  |
| Type | 1 | 1 | 1 |  |  |  |  |  |


| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | STIMSTRT | STIMT | STIMDUR | STIMAMP |  |  |  |  |
| Type | F | F | F | F |  |  |  |  |


| VARIABLE | DESCRIPTION |  |
| :---: | :--- | :--- |
|  |  | STIMID |
| SETTYPE |  | ID of the stimulation |
|  |  | EQ.1: Segment set |
|  | EQ.2: Node set |  |
| SETID |  | Node set or segment set ID to be stimulated |
| STIMSTRT |  | Starting time of the stimulation |
| STIMT |  | Stimulation period |
| STIMDUR |  | Stimulation duration |
| STIMAMP | Stimulation amplitude (picoA/picoF) |  |

## *EM_EOS_BURGESS

Purpose: Define the parameters for a Burgess model giving the electrical conductivity as as a function of the temperature and the density, see:
T.J. Burgess, "Electrical resistivity model of metals", 4th International Conference on Megagauss Magnetic-Field Generation and Related Topics, Santa Fe, NM, USA, 1986

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | EOSID | V0 | GAMMA | THETA | LF | C 1 | C 2 | C 3 |
| Type | I | F | F | F | F | F | F | F |
| Default | none | none | none | none | none | none | none | none |


| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | C4 | K | EXPON | LGTUNIT | TIMUNIT | TEMUNI | ADJUST |  |
| Type | F | F | I | F | F | I | I |  |
| Default | none | none | none | none | none | none | none |  |

In the following, UUS stands for User Units System and BUS for Burgess Units

## VARIABLE

EOSID

V0
GAMMA0

THETA

LF Latent heat of fusion $\mathrm{L}_{\mathrm{F}}$ in kJoule/mol (BUS).
C1 $\quad$ C1 constant (BUS)
C2 C2 constant (no units)
*EM_EOS_BURGESS
VARIABLE

## DESCRIPTION

C3
C4
K
EXPON Exponent in equations (2) (see remarks)
LGTUNIT Length units for UUS (relative to meter, i.e. $=1 . \mathrm{e}-3$ if UUS in mm).
TIMUNIT Time units for UUS (relative to seconds).
TEMUNIT Temperature units
EQ.1: temperature in Celsius
EQ.2: temperature in Kelvins
ADJUST Conductivity modification
EQ.O: (default) The conductivity is given by the Burgess formula.

EQ.1: The conductivity is adjusted so that it is equal to the conductivity defined in ${ }^{*}$ EM_MAT card $\sigma_{\text {mat }}$ at room temperature:

$$
\sigma(\theta)=\sigma_{\text {Burgess }}(\theta) \frac{\sigma_{\text {mat }}}{\sigma_{\text {Burgess }}\left(\theta_{\text {room }}\right)}
$$

## Remarks:

1. The Burgess model gives the electrical resistivity vs temperature and density for the solid phase, liquid phase and vapor phase. At this time, only the solid and liquid phases are implemented. To check which elements are in the solid and in the liquid phase, a melting temperature is first computed by:

$$
\theta_{m}=\theta_{m, 0}\left(\frac{V}{V_{0}}\right)^{-\frac{1}{3}} e^{\left(2 \gamma_{0}-1\right)\left(1-\frac{V}{V_{0}}\right)}
$$

a) If $\mathrm{T}<\theta_{m}$ : solid phase model applies.

The solid phase electrical resistivity corresponds to the Meadon model:

$$
\begin{equation*}
\eta_{S}=\left(C_{1}+C_{2} \theta^{C_{3}}\right) f_{c}\left(\frac{V}{V_{0}}\right), \tag{1}
\end{equation*}
$$

where $\theta$ is the temperature, $V$ is the specific volume, and $V_{0}$ is the reference specific volume (zero pressure, solid phase). In (1), the volume dependence is given by:

$$
f_{c}\left(\frac{V}{V_{0}}\right)=\left\{\begin{array}{llr}
\left(\frac{V}{V_{0}}\right)^{2 \gamma-1} & \text { EXPON.EQ. }-1 & \text { (most materials) }  \tag{2}\\
\left(\frac{V}{V_{0}}\right)^{2 \gamma+1} & \text { EXPON.EQ. }+1 & \text { (tungsten) } \\
\left(\frac{V}{V_{0}}\right)^{2 \gamma} & \text { EXPON.EQ. } 0 & \text { (stainless steel) }
\end{array}\right.
$$

with

$$
\begin{equation*}
\gamma=\gamma_{0}-\left(\gamma_{0}-\frac{1}{2}\right)\left(1-\frac{V}{V_{0}}\right) \tag{3}
\end{equation*}
$$

b) If $\mathrm{T}>\theta_{\mathrm{m}}$ : liquid phase model:

$$
\begin{equation*}
\eta_{L}=\left(\eta_{L}\right)_{\theta_{m}}\left(\frac{\theta}{\theta_{m}}\right)^{C_{4}} \tag{4}
\end{equation*}
$$

with

$$
\left(\eta_{L}\right)_{\theta_{m}}=\Delta \eta\left(\eta_{S}\right)_{\theta_{m}}
$$

where

$$
\Delta \eta= \begin{cases}k e^{0.69 L_{F} / \theta_{m}} & k>0  \tag{5}\\ 1+0.0772\left(2-\theta_{m}\right) & k=-1 \\ 1+0.106\left(0.846-\theta_{m}\right) & k=-2 \quad \text { (stainless steel SS-304) }\end{cases}
$$

The following table reports some sets of parameters given by Burgess in his paper:

| Parameter | $\mathbf{C u}$ | Ag | Au | $\mathbf{W}$ | Al(2024) | SS(304) |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| $\left.\mathbf{V}_{\mathbf{0}} \mathbf{( c m}^{3} / \mathbf{g m}\right)$ | 0.112 | 0.0953 | 0.0518 | 0.0518 | 0.370 | 0.1265 |
| $\gamma_{0}$ | 2.00 | 2.55 | 3.29 | 1.55 | 2.13 | 2.00 |
| $\boldsymbol{\theta}_{\mathrm{m}, 0}$ (BUS) | 0.117 | 0.106 | 0.115 | 0.315 | 0.0804 | 0.156 |
| $\mathbf{L}_{\mathrm{F}}$ (BUS) | 0.130 | 0.113 | 0.127 | 0.337 | 0.107 | 0.153 |
| $\mathbf{C}_{\mathbf{1}}$ (BUS) | $-4.12 \mathrm{e}-5$ | $-3.37 \mathrm{e}-5$ | $-4.95 \mathrm{e}-5$ | $-9.73 \mathrm{e}-5$ | $-5.35 \mathrm{e}-5$ | 0 |


| Parameter | $\mathbf{C u}$ | $\mathbf{A g}$ | $\mathbf{A u}$ | $\mathbf{W}$ | Al(2024) | SS(304) |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: |
| $\mathrm{C}_{2}$ | 0.113 | 0.131 | 0.170 | 0.465 | 0.233 | 0.330 |
| $\mathrm{C}_{3}$ | 1.145 | 1.191 | 1.178 | 1.226 | 1.210 | 0.4133 |
| EXPON | -1 | -1 | -1 | +1 | -1 | 0 |
| $\mathbf{C}_{4}$ | 0.700 | 0.672 | 0.673 | 0.670 | 0.638 | 0.089 |
| $\mathbf{k}$ | 0.964 | 0.910 | 1.08 | -1. | 0.878 | -2. |

## *EM_EOS_MEADON

Purpose: Define the parameters for a Meadon model, giving the electrical conductivity as a function of the temperature and the density; see:
T.J. Burgess, "Electrical resistivity model of metals", 4th International Conference on Megagauss Magnetic-Field Generation and Related Topics, Santa Fe, NM, USA, 1986

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | EOSID | C 1 | C 2 | C 3 | TEMUNI | V0 | GAMMA | EXPON |
| Type | I | F | F | F | I | F | F | I |
| Default | none | none | none | none | none | none | none | none |


| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | LGTUNIT | TIMUNIT | ADJUST |  |  |  |  |  |
| Type | F | F | I |  |  |  |  |  |
| Default | none | none | none |  |  |  |  |  |

In the following, UUS stands for User Units System and BUS for Burgess Units.

| VARIABLE |  | EOSID |
| :---: | :--- | :--- |
|  |  | ID of the EM_EOS |
| C1 |  | C1 constant (BUS) |
| C2 |  | C2 constant (no units) |
| C3 |  | C3 constant (no units) |

TEMUNIT Temperature units
EQ.1: temperature in Celsius
EQ.2: temperature in Kelvins

## VARIABLE

V0
GAMMA0
EXPON
LGTUNIT
TIMUNIT
ADJUST: EQ.O: (default) the conductivity is given by the Burgess formula.
EQ.1: The conductivity is adjusted so that it is equal to the conductivity defined in the *EM_MAT card $\sigma_{\text {mat }}$ at room temperature:

$$
\sigma(\theta)=\sigma_{\text {Burgess }}(\theta) \frac{\sigma_{\text {mat }}}{\sigma_{\text {Burgess }}\left(\theta_{\text {room }}\right)}
$$

## Remarks:

1. The Meadon model is a simplified Burgess model with the solid phase equations only.

The electrical resistivity is given by:

$$
\begin{equation*}
\eta_{S}=\left(C_{1}+C_{2} \theta^{C_{3}}\right) f_{c}\left(\frac{V}{V_{0}}\right) \tag{6}
\end{equation*}
$$

where $\theta$ is the temperature, V is the specific volume, and $\mathrm{V}_{0}$ is the reference specific volume (zero pressure, solid phase).

In (6), the volume dependence is given by:
$f_{c}\left(\frac{V}{V_{0}}\right)=\left\{\begin{array}{llr}\left(\frac{V}{V_{0}}\right)^{2 \gamma-1} & \text { EXPON.EQ. }-1 & \text { (most materials) } \\ \left(\frac{V}{V_{0}}\right)^{2 \gamma+1} & \text { EXPON.EQ. }+1 & \text { (tungsten) } \\ \left(\frac{V}{V_{0}}\right)^{2 \gamma} & \text { EXPON.EQ. } 0 & \text { (stainless steel) } \\ 1 & \text { VO.EQ. } 0 & \text { (default value for } V_{0} \text { is zero) }\end{array}\right.$
(In this last case, only EOSID, C1, C2, C3, TEMUNIT, TIMUNIT and LGTUNIT need to be defined)
with,

$$
\begin{equation*}
\gamma=\gamma_{0}-\left(\gamma_{0}-\frac{1}{2}\right)\left(1-\frac{V}{V_{0}}\right) \tag{8}
\end{equation*}
$$

The following table reports some sets of parameters given by Burgess in his paper:

| Parameter | $\mathbf{C u}$ | $\mathbf{A g}$ | Au | $\mathbf{W}$ | Al(2024) | SS(304) |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: |
| $\mathbf{V}_{\mathbf{0}}\left(\mathbf{c m}^{3} / \mathbf{g m}\right)$ | 0.112 | 0.0953 | 0.0518 | 0.0518 | 0.370 | 0.1265 |
| $\gamma_{0}$ | 2.00 | 2.55 | 3.29 | 1.55 | 2.13 | 2.00 |
| $\mathrm{C}_{\mathbf{1}}$ (BUS) | $-4.12 \mathrm{e}-5$ | $-3.37 \mathrm{e}-5$ | $-4.95 \mathrm{e}-5$ | $-9.73 \mathrm{e}-5$ | $-5.35 \mathrm{e}-5$ | 0 |
| $\mathrm{C}_{2}$ | 0.113 | 0.131 | 0.170 | 0.465 | 0.233 | 0.330 |
| $\mathrm{C}_{3}$ | 1.145 | 1.191 | 1.178 | 1.226 | 1.210 | 0.4133 |
| EXPON | -1 | -1 | -1 | +1 | -1 | 0 |

## *EM_EOS_PERMEABILITY

Purpose: Define the parameters for the behavior of a material's permeability

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | EOSID | EOSTYPE | LCID |  |  |  |  |  |
| Type | 1 | 1 | 1 |  |  |  |  |  |
| Default | none | none | none |  |  |  |  |  |

VARIABLE
EOSID
EOSTYPE

LCID

DESCRIPTION
ID of the EM_EOS
Define the type of EOS:
EQ.1: Permeability defined by a $B$ function of $H$ curve $(B=\mu H)$
EQ.2: Permeability defined by a $H$ function of $B$ curve $(H=B / \mu)$

Load curve ID

## *EM_EOS_TABULATED1

Purpose: Define the electrical conductivity or permeability depending on the material referencing this EOS as a function of temperature by using a load curve.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | EOSID | LCID |  |  |  |  |  |  |
| Type | 1 | 1 |  |  |  |  |  |  |
| Default | none | none |  |  |  |  |  |  |

VARIABLE
EOSID
ID of the EM EOS
LCID Load curve ID. See Remark 1.

## Remarks:

1. Suggestions for the Load Curve. The load curve describes the electrical conductivity or permeability (ordinate) as a function of the temperature (abscissa). You need to make sure the temperature and the electrical conductivity / permeability given by the load curve are in the correct units. Also, we advise giving some bounds to the load curve (conductivities / permeabilities at very low and very high temperatures) to avoid bad extrapolations of the conductivity if the temperature gets out of the load curve bounds.

## *EM_EOS_TABULATED2

Purpose: Define the electrical conductivity or permeability (depending on the material model referencing this EOSID) as a function of time by using a load curve.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | EOSID | LCID | IFLAG |  |  |  |  |  |
| Type | 1 | 1 | 1 |  |  |  |  |  |
| Default | none | none | 0 |  |  |  |  |  |

## VARIABLE

EOSID
LCID Load curve ID (see Remark 1), function ID (see *DEFINE_FUNCTION), table ID or 2D table ID. For the arguments for the *DEFINE_FUNCTION, see Remark 2.

IFLAG If LCID is a table ID or 2D table ID, conductivity / permeability is a function of temperature and material density. This flag dictates how LS-DYNA interprets the table. In other words, it specifies which property (temperature or material density) is the value for the table and which is the ordinate for load curves in the table:

EQ.0: Temperature (value) indexes each conductivity/permeability (ordinate) versus material density (abscissa) load curve.

EQ.1: Material density indexes each conductivity/permeability (ordinate) versus temperature (abscissa) load curve.

## Remarks:

1. Suggestions for the Load Curve. The load curve describes the electrical conductivity / permeability (ordinate) as a function of time (abscissa). You need to make sure the time and the electrical conductivity / permeability given by the load curve are in the correct units. Also, we advise giving some bounds to the load curve (conductivities / permeabilities at $t=0$ and after a long time) to avoid bad extrapolations of the conductivity / permeability if the run time gets out of the load curve bounds.
2. *DEFINE_FUNCTION Arguments. LCID can also refer to a *DEFINE_FUNCTION ID. If a *DEFINE_FUNCTION is used, the following parameters are allowed: $f(v x$, vy, vz, temp, pres, vol, mass, Ex, Ey, Ex, Bx, By, Bz, Fx, Fy, Fz, JHrate, time, x, y, z). (Fx, Fy, Fz) refers to the Lorentz force vector.

## *EM_EXTERNAL_FIELD

Purpose: Define the components of a time dependent exterior field uniform in space applied on the conducting parts.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | FIELDID | FTYPE | FDEF | LCIDX | LCIDY | LCIDZ |  |  |
| Type | I | I | F | I | I | I |  |  |
| Default | 0 | 0 | 0 | 0 | 0 | 0 |  |  |

FDEF Field defined by:

## VARIABLE

FIELDID
FTYPE

LCID[X,Y,Z]

FDEF Field defined by:
EQ.1: load curves
EQ.2: define function (FTYPE $=3$ only). If a define function is
used, the following parameters are accepted: $x, y, z$, time,
EQ.2: define function (FTYPE $=3$ only). If a define function is
used, the following parameters are accepted: $x, y, z$, time, emdt, pot, curr, sigma.
EQ.1. magnetic field
EQ.2: electric field (not available yet)
EQ.3: charge density (resistive heating solver only)

Load curve ID defining the ( $X, Y, Z$ ) component of the field function of time for FTYPE $=1$. For FTYPE $=3$, only LCIDY is used and should be a simple a load curve or define function ID.

## Remarks:

1. Electrostatic Problems. FTYPE $=3$ is mostly used in electrostatic problem configurations. The material's conductivity then represents the permittivity.

## *EM_ISOPOTENTIAL

Purpose: Define an isopotential. In other words, constrain nodes so that they have the same scalar potential value. This keyword card can only be used with the type 3 EM solver.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | ISOID | SETTYPE | SETID | RDLTYPE |  |  |  |  |
| Type | 1 | 1 | 1 | 1 |  |  |  |  |
| Default | none | none | none | 0 |  |  |  |  |

## VARIABLE

ISOID ID of the isopotential

Set type:
EQ.1: Segment Set
EQ.2: Node Set
EQ.3: Fluid surface part. See *ICFD_PART.

SETID
RDLTYPE Used for the battery application (see *EM_RANDLES_BATMAC or *EM_RANDLES_TSHELL). Selects which layers of the underlying battery cell are associated with the isopotential:

EQ.O: Default. No specific treatment.
EQ.1: Current Collector Positive
EQ.2: Positive Electrode
EQ.3: Separator
EQ.4: Negative Electrode
EQ.5: Current Collector Negative
The function of a layer is defined in *EM_MAT_001.

## *EM_ISOPOTENTIAL_CONNECT

Purpose: Define a connection between two isopotentials or between an isopotential and the ground.

## Card Summary:

Card 1a. Include this card for CONTYPE = 1 (short circuit).

| CONID | CONTYPE | ISOID1 | ISOID2 |  |  |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

Card 1b. Include this card for CONTYPE $=2,3,4$, or 6 .

| CONID | CONTYPE | ISOID1 | ISOID2 | VAL | LCID |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

Card 1c. Include this card for CONTYPE $=5$ (meshless Randles circuit).

| CONID | CONTYPE | ISOID1 | ISOID2 |  | RDLID | PSID |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

Card 2. Include this card for CONTYPE $=6$ (RLC circuit).

| L | C | Vo |  |  |  |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

## Data Card Definitions:

Short Circuit Connection Card. Include this card if CONTYPE=1.

| Card 1a | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | CONID | CONTYPE | ISOID1 | ISOID2 |  |  |  |  |
| Type | 1 | 1 | 1 | 1 |  |  |  |  |
| Default | none | none | none | 0 |  |  |  |  |

## VARIABLE

## DESCRIPTION

CONID Connection ID
CONTYPE Connection type:
EQ.1: Short circuit
ISOID1 ID of the first isopotential to be connected

## VARIABLE

ISOID2

## DESCRIPTION

Optional ID of the second isopotential to be connected

Resistance, Voltage Source, Current Source, and RLC Circuit Connection Card. Include this card if CONTYPE $=2,3,4$, or 6 .

| Card 1b | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | CONID | CONTYPE | ISOID1 | ISOID2 | VAL | LCID |  |  |
| Type | 1 | 1 | 1 | 1 | $F$ | 1 |  |  |
| Default | none | none | none | 0 | none | 0 |  |  |

## VARIABLE

CONID
CONTYPE

ISOID1
ISOID2
VAL Value of the resistance, voltage, or current depending on CONTYPE. Ignored for CONTYPE $=2$ through 4 if LCID $\neq 0$.

LCID Load curve ID giving the value of the resistance, voltage, or current as a function of time. Only available for CONTYPE $=2$ through 4.

LT.O: |LCID| is a *DEFINE_FUNCTION ID. The following arguments are allowed: $f($ time, emdt, curr1, curr2, pot1, pot2, rmesh). pot1 and pot2 are the potential at the previous time step and at two previous time steps. curr1 and curr2 are the current at the previous time step and two previous time steps. rmesh is the mesh resistance calculated by the solver at this isopotential.

Meshless Randles Circuit Connection Card. Include this card if CONTYPE $=5$.

| Card 1c | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | CONID | CONTYPE | ISOID1 | ISOID2 |  | RDLID | PSID |  |
| Type | 1 | 1 | 1 | 1 |  | 1 | 1 |  |
| Default | none | none | none | 0 |  | none | 0 |  |

VARIABLE
CONID
CONTYPE

ISOID1
ISOID2
VAL

RDLID
PSID Used if the variable ROTOTH of *EM_RANDLES_MESHLESS equals 1 . Part set ID where LS-DYNA adds the joule heating corresponding to the resistance $r_{0}$ in *EM_RANDLES_MESHLESS, averaged over the part set.

RLC Circuit Parameters Card. Only defined if CONTYPE $=6$.

| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | L | C | Vo |  |  |  |  |  |
| Type | F | F | F |  |  |  |  |  |
| Default | none | none | none |  |  |  |  |  |

## VARIABLE

L/C/V0 Circuit inductance, capacity, and initial voltage. VAL gives the resistance.

## *EM_ISOPOTENTIAL_ROGO

Purpose: measures the total current flowing through a given section of the conductor and outputs it in an ASCII file called em_rogoCoil.dat

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | ISOID | SETTYPE | SETID |  |  |  |  |  |
| Type | 1 | 1 | 1 |  |  |  |  |  |
| Default | none | none | none |  |  |  |  |  |


| VARIABLE |  | D |
| :---: | :--- | :--- |
| ISOID |  | ID of the Rogo coil. |
| SETTYPE |  | Set type: |
|  |  | EQ.1: Segment Set. |
| SETID | Set ID |  |

## *EM_MAT_001

Purpose: Define the electromagnetic material type and properties for a material whose permeability equals the free space permeability.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | MID | MTYPE | SIGMA | EOSID |  |  | DEATHT | RDLTYPE |
| Type | I | I | F | I |  |  | $F$ | 1 |
| Default | none | none | none | none |  |  | $10^{28}$ | none |

## VARIABLE

MID
MTYPE Defines the electromagnetism type of the material:
EQ.0: Air or vacuum
EQ.1: Insulator material. These materials have the same electromagnetism behavior as MTYPE $=0$.

EQ.2: Conductor carrying a source. In these conductors, the eddy current problem is solved, which gives the actual current density. Typically, this material would correspond to the coil.

EQ.3: Fluid conductor. In this case, MID refers to the ID given in *ICFD_PART. See Remark 1.

EQ.4: Conductor not connected to any current or voltage source, where the Eddy current problem is solved. Typically, this material would correspond to the workpiece.

SIGMA Initial electrical conductivity of the material
EOSID Optional ID of the EOS to be used for the electrical conductivity (see *EM_EOS cards).

RDLTYPE Used for composite Tshell batteries modeled with *EM_RANDLES_TSHELL. RDLTYPE specifies the function of the layer associated to MID:

EQ.O: Default. Conductor which is not part of a battery cell.
EQ.1: Current Collector Positive
EQ.2: Positive Electrode
EQ.3: Separator
EQ.4: Negative Electrode
EQ.5: Current Collector Negative

## Remarks:

1. Coupling ICFD to EM. Only the resistive heating solver is currently available when coupling the ICFD solver with the EM solver (see *EM_CONTROL).

## *EM_MAT_002

Purpose: Define an electromagnetic material type and properties whose permeability is different than the free space permeability.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | MID | MTYPE | SIGMA | EOSID | MUREL | EOSMU | DEATHT |  |
| Type | A | 1 | F | I | F | 1 | F |  |
| Default | none | none | none | none | none | none | $10^{28}$ |  |

Optional card

| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable |  | EOSID2 |  |  |  |  |  |  |
| Type |  | 1 |  |  |  |  |  |  |
| Default |  | none |  |  |  |  |  |  |

## VARIABLE

MID

MTYPE Electromagnetism type of the material:
EQ.O: Air or vacuum
EQ.1: Insulator material. These materials have the same electromagnetism behavior as MTYPE $=0$.

EQ.2: Conductor carrying a source. In these conductors, the eddy current problem is solved, which gives the actual current density. Typically, this would correspond to the coil.

EQ.4: Conductor not connected to any current or voltage source, where the Eddy current problem is solved. Typically, this would correspond to the workpiece.

SIGMA
EOSID ID of the EOS to be used for the electrical conductivity (see *EM_EOS cards)

MUREL Relative permeability which is the ratio of the permeability of a specific medium to the permeability of free space ( $\mu_{r}=\mu / \mu_{0}$ )

EOSID2
ID of the EOS to be used to define the nonlinear behavior of $\mu$. Note: if EOSMU is defined, MUREL will be used for the initial value only. See EM_EOS_PERMEABILITY.

Death time for the material. After DEATHT, the material will no longer be considered a conductor and will be removed from the EM solve. If a negative value is entered, a *DEFINE_FUNCTION will be expected. The following parameters are allowed: (vx, vy, vz, temp, vol, mass, Ex, Ey, Ez, Bx, By, Bz, Fx, Fy, Fz, JHrate, time). Fx, Fy, and Fz refer to the components of the Lorentz force vector. A negative value returned by the *DEFINE_FUNCTION corresponds to a 'dead' or inactive element. Once an element has been removed from the EM solve, it cannot return.

Optional ID of the EOS for specifying the behavior of $\mu$ by an equa- tion of state. See *EM_EOS_TABULATED1 and *EM_EOS_TABULATED2.
*EM_MAT_003
Purpose: Define an electromagnetic material type whose electromagnetic conductivity is defined by a $(3 \times 3)$ tensor matrix. Applications include composite materials. This material only applies to solid elements.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | MID | MTYPE | SIGMA11 | SIGMA22 | SIGMA33 | BETA | CM |  |
| Type | I | I | F | F | F | F | F |  |


| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | SIGMA12 | SIGMA13 | SIGMA21 | SIGMA23 | SIGMA31 | SIGMA32 | AOPT | LAMBDA |
| Type | I | I | F | F | F | F | I | F |


| Card 3 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | XP | YP | ZP | A 1 | A2 | A3 | MACF |  |
| Type | F | F | F | F | F | F | I |  |


| Card 4 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | V1 | V2 | V3 | D1 | D2 | D3 |  |  |
| Type | F | F | F | F | F | F |  |  |

VARIABLE
MID

MTYPE

## DESCRIPTION

Material ID. A unique number must be specified (see *PART).
Defines the electromagnetism type of the material:
EQ.0: Air or vacuum

## VARIABLE

## DESCRIPTION

EQ.1: Insulator material. These materials have the same electromagnetism behavior as MTYPE $=0$.

EQ.2: Conductor carrying a source. For these conductors, the EM solver solves the eddy current problem, which gives the actual current density. Typically, this type would correspond to the coil. In electrophysiology (EP), it corresponds to the tissue where the monodomain equations are solved for $\mathrm{EMSOL}=11$ or $\mathrm{EMSOL}=13$. For this case, an *EM_EP_CELLMODEL must be associated to this *EM_MAT_003.

EQ.4: Conductor not connected to any current or voltage source, where the Eddy current problem is solved. Typically, this type would correspond to the workpiece. In electrophysiology (EP), for $\mathrm{EMSOL}=11,12$ or 13, it corresponds to the bath surrounding the tissue for which only the external potential is found. In the case of electrophysiology, no *EM_EP_CELLMODEL should be associated with this material.

SIGMA11 The 1,1 term in the $3 \times 3$ electromagnetic conductivity tensor matrix. Note that 1 corresponds to the a material direction.

LT.O.0: |SIGMA11| corresponds to the ID of a *DEFINE_FUNCTION. See Remark 2 for available parameters.

The 1,2 term in the $3 \times 3$ electromagnetic conductivity tensor matrix. Note that 2 corresponds to the $\mathbf{b}$ material direction.

LT.O.0: |SIGMA12| corresponds to the ID of a *DEFINE_FUNCTION. See Remark 2 for available parameters.

SIGMA33
The 3, 3 term in the $3 \times 3$ electromagnetic conductivity tensor matrix.

LT.O.O: |SIGMA33| corresponds to the ID of a *DEFINE_FUNCTION. See Remark 2 for available parameters.

BETA Surface to volume ratio of the cell membrane (to be used only when EMSOL = 11 or 12 in *EM_CONTROL).

CM Membrane capacitance per unit area (to be used only when EMSOL = 11 or 12 in EM_CONTROL).

## VARIABLE

AOPT

## DESCRIPTION

Material axes option (see *MAT_002 for a more detailed description):

EQ.0.0: Locally orthotropic with material axes determined by element nodes. The a-direction is from node 1 to node 2 of the element. The $\mathbf{b}$-direction is orthogonal to the $\mathbf{a}-$ direction and is in the plane formed by nodes 1,2 , and 4.

EQ.1.0: Locally orthotropic with material axes determined by a point in space, $P$, and the global location of the element center; this is the a-direction.

EQ.2.0: Globally orthotropic with material axes determined by vectors defined below, as with *DEFINE_COORDINATE_VECTOR.

EQ.3.0: Locally orthotropic material axes determined by a vector $\mathbf{v}$ and the normal vector to the plane of the element.The plane of a solid element is the midsurface between the inner surface and outer surface defined by the first four nodes and the last four nodes of the connectivity of the element, respectively. Thus, for solid elements, AOPT $=3$ is only available for hexahedrons. a is determined by taking the cross product of $\mathbf{v}$ with the normal vector, $\mathbf{b}$ is determined by taking the cross product of the normal vector with $\mathbf{a}$, and $\mathbf{c}$ is the normal vector. Then $\mathbf{a}$ and $\mathbf{b}$ are rotated about $\mathbf{c}$ by an angle BETA. BETA may be set in the keyword input for the element.

EQ.4.0: Locally orthotropic in cylindrical coordinate system with the material axes determined by a vector, $\mathbf{v}$, and an originating point, $P$, which define the centerline axis.

XP, YP, ZP Coordinates of point, $P$, for AOPT $=1$ and 4
A1, A2, A3 Components of vector, $\mathbf{a}$, for $\mathrm{AOPT}=2$
MACF

V1, V2, V3 Components of vector, $\mathbf{v}$, for $\mathrm{AOPT}=3$ and 4.
D1, D2, D3 Components of vector, $\mathbf{d}$, for $\mathrm{AOPT}=2$

LAMBDA
Intra- to extracellular conductivity ratio. When non-empty, the elliptic equation is solved to compute extracellular potentials (to be used only when EMSOL = 11 in *EM_CONTROL).

## Remarks:

1. Material Directions. See the manual page for *MAT_002 for a description of how the principal material directions, $\{\mathbf{a}, \mathbf{b}, \mathbf{c}\}$, are determined with AOPT. Note that *EM_MAT_003 only works for solid elements. The AOPT options illustrated in the AOPT figure of *MAT_002 can define the material directions for all elements of the parts that use the material.
2. Function Parameters. The available parameters for the *DEFINE_FUNCTIONs are: f(time, emdt, $x$ _ele, y_ele, z_ele, ieleuser). Here, time is the current EM time; emdt is the current EM time step; $x_{-}$ele, $y \_$ele, and $z \_$ele are the location of the element; and ieleuser is the element ID.
*EM_MAT_004
Purpose: Define the electromagnetic material type and properties for conducting shells in a 3D problem or in a 2D resistive heating problem.

Include as many cards as needed. This input ends at the next keyword ("*") card.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | MID | MTYPE | SIGMA | EOSID | NELE | MUREL | EOSMU | DEATHT |
| Type | I | I | F | 1 | 1 | $F$ | 1 | $F$ |
| Default | none | none | none | none | 1 | 1. | none | $10^{28}$ |

VARIABLE
MID

MTYPE Defines the electromagnetism type of the material:
EQ.O: Air or vacuum
EQ.1: Insulator material. These materials have the same electromagnetism behavior as MTYPE $=0$.

EQ.2: Conductor carrying a source. In these conductors, the eddy current problem is solved, which gives the actual current density. Typically, this would correspond to the coil.

EQ.4: Conductor not connected to any current or voltage source,
where the Eddy current problem is solved. Typically, this
EQ.4: Conductor not connected to any current or voltage source,
where the Eddy current problem is solved. Typically, this would correspond to the workpiece.

SIGMA Initial electrical conductivity of the material
EOSID ID of the EOS to be used for the electrical conductivity (see *EM_EOS cards).

NELE Number of elements in the thickness of the shell. Note that you must make sure your mesh fine enough to correctly capture the in-ductive-diffusive effects (see skin depth definition).

MUREL

## DESCRIPTION

Material ID: refers to MID in the *PART card.

Relative permeability which is the ratio of the permeability of a specific medium to the permeability of free space $\left(\mu_{r}=\mu / \mu_{0}\right)$.

## VARIABLE

EOSMU

DEATHT

## DESCRIPTION

ID of the EOS to be used to define the nonlinear behavior of $\mu$. Note that if EOSMU is defined, MUREL will be used for the initial value only. See *EM_EOS_PERMEABILITY.

Death time for the material. After DEATHT, the material will no longer be considered a conductor and will be removed from the EM solve. If a negative value is entered, a *DEFINE_FUNCTION will be expected. The following parameters are allowed: (vx, vy, vz, temp, vol, mass, Ex, Ey, Ez, Bx, By, Bz, Fx, Fy, Fz, JHrate, time). Fx, Fy, and Fz refer to the components of the Lorentz force vector. A negative value returned by the *DEFINE_FUNCTION corresponds to a 'dead' or inactive element. Once an element has been removed from the EM solve, it cannot return.
*EM_MAT_005
Purpose: Define an electromagnetic material for which two material conductivities are specified per EM node and electromagnetic conductivities are defined by a ( $3 \times 3$ ) tensor matrix. Applications of this material include the Randles Batmac model (see Remark 1) and the electrophysiology bidomain model. This material is only supported for solid elements.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | MID | MTYPE | SIGMAXXA | SIGMAYYA | SIGMAZZA |  |  |  |
| Type | I | I | F | F | F |  |  |  |


| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | SIGMAXYA | SIGMAXZA | SIGMAYXA | SIGMAYZA | SIGMAZXA | SIGMAZYA |  |  |
| Type | F | F | F | F | F | F |  |  |


| Card 3 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable |  |  | SIGMAXXB | SIGMAYYB | SIGMAZZB |  |  |  |
| Type |  |  | F | F | F |  |  |  |


| Card 4 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | SIGMAXYB | SIGMAXZB | SIGMAYXB | SIGMAYZB | SIGMAZXB | SIGMAZYB |  |  |
| Type | F | F | F | F | F | F |  |  |


| Card 5 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | AOPT | XP | YP | ZP | A1 | A2 | A3 | MACF |
| Type | F | F | F | F | F | F | I | I |


| Card 6 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | V1 | V2 | V3 | D1 | D2 | D3 |  |  |
| Type | F | F | F | F | F | F |  |  |

## VARIABLE

MID
MTYPE
Defines the electromagnetism type of the material:
EQ.O: Air or vacuum
EQ.1: Insulator material. These materials have the same electromagnetism behavior as MTYPE $=0$.
EQ.2: In electrophysiology, it corresponds to the tissue, where the bidomain equations will be solved for $E M S O L=12$ or EMSOL $=13$. An ${ }^{*}$ EM_EP_CELLMODEL must be associated with this material.

EQ.4: In electrophysiology, it corresponds to the bath where only the external potential is solved for. No *EM_EP_CELLMODEL should be associated with this material.

EQ.5: Material associated with *EM_RANDLES_BATMAC. See Remark 1.

SIG- The 1, 1 term in the $3 \times 3$ electromagnetic conductivity tensor maMAXXA/B

## DESCRIPTION

Material ID. A unique number must be specified (see *PART). trix for the two conductivities. For the BatMac model, A is for the potential on the positive current collector, and $B$ is for the potential on the negative current collector. For the bidomain model in Electrophysiology, A is for the intracellular potential, and B is for the extracellular potential. Note that 1 corresponds to the a material direction.

LT.0.0: |SIGMAXXA/B | corresponds to the ID of a *DEFINE_FUNCTION. See Remark 3 for available parameters.

SIG- The 1, 2 term in the $3 \times 3$ electromagnetic conductivity tensor ma-

SIG-

AOPT Material axes option (see *MAT_002 for a more detailed description):

EQ.0.0: Locally orthotropic with material axes determined by element nodes. The a-direction is from node 1 to node 2 of the element. The $\mathbf{b}$-direction is orthogonal to the $\mathbf{a}-$ direction and is in the plane formed by nodes 1,2 , and 4.

EQ.1.0: Locally orthotropic with material axes determined by a point in space, $P$, and the global location of the element center; this is the a-direction.

EQ.2.0: Globally orthotropic with material axes determined by vectors defined below, as with *DEFINE_COORDINATE_VECTOR.

EQ.3.0: Locally orthotropic material axes determined by a vector $\mathbf{v}$ and the normal vector to the plane of the element.The plane of a solid element is the midsurface between the inner surface and outer surface defined by the first four nodes and the last four nodes of the connectivity of the element, respectively. Thus, for solid elements, AOPT = 3 is only available for hexahedrons. $\mathbf{a}$ is determined by taking the cross product of $\mathbf{v}$ with the normal vector, $\mathbf{b}$ is determined by taking the cross product of the normal vector with $\mathbf{a}$, and $\mathbf{c}$ is the normal vector. Then $\mathbf{a}$ and $\mathbf{b}$ are rotated about $\mathbf{c}$ by an angle BETA. BETA may be set in the keyword input for the element.

EQ.4.0: Locally orthotropic in cylindrical coordinate system with the material axes determined by a vector, $\mathbf{v}$, and an originating point, $P$, which define the centerline axis.

XP, YP, ZP $\quad$ Define coordinates of point $\mathbf{p}$ for $\mathrm{AOPT}=1$ and 4.
A1, A2, A3 Define components of vector a for AOPT $=2$.
MACF Material axes change flag for solid elements:
EQ.1: No change, default
V1, V2, V3 Define components of vector $\mathbf{v}$ for $\mathrm{AOPT}=3$ and 4 .
D1, D2, D3 Define components of vector $\mathbf{d}$ for $\mathrm{AOPT}=2$.

## Remarks:

1. Using this Material with BatMac. When this material is used in conjunction with the battery BatMac model, then the conductivities must be adjusted by the current collector thickness ratio over the total thickness of the cell. For example, if $n_{p}$ is the number of positive current collectors, $t_{p}$ the thickness of each individual positive current collector and Th the total thickness of the cell, then the conductivity for the positive current collector must be scaled by:

$$
\frac{n_{p} \times t_{p}}{\mathrm{Th}}
$$

2. Material Directions. See the manual page for *MAT_002 for a description of how the principal material directions, $\{\mathbf{a}, \mathbf{b}, \mathbf{c}\}$, are determined with AOPT. Note that *EM_MAT_005 only works for solid elements. The AOPT options illustrated in the AOPT figure of *MAT_002 can define the material directions for all elements of the parts that use the material.
3. Function Parameters. The available parameters for the *DEFINE_FUNC-
 EM time; emdt is the current EM time step; $x_{-}$ele, $y \_$ele, and $z \_$ele are the location of the element; and ieleuser is the element ID.

## *EM_MAT_006

Purpose: Define two conductivities per EM node for special applications (Randles Batmac).

Include as many cards as needed. This input ends at the next keyword ("*") card.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | MID | MTYPE | SIGP | EOSP | SIGN | EOSN | DEATHT |  |
| Type | 1 | 1 | F | 1 | F | 1 | F |  |
| Default | none | none | none | none | none | none | $10^{28}$ |  |

VARIABLE
MID

MTYPE Defines the electromagnetism type of the material:
EQ.0: Air or vacuum
EQ.1: Insulator material. These materials have the same electromagnetism behavior as MTYPE $=0$.

EQ.5: Material associated to *EM_RANDLES_BATMAC

SIGP/SIGN
EOSP/EOSN
DEATHT

Conductivities of the positive / negative current collector materials
Optional ID of the EOS to be used for the two conductivities
Death time for the material. After DEATHT, the material will no longer be considered a conductor and removed from the EM solve. If a negative value is entered, a |DEATHT| is a *DEFINE_FUNCTION ID. The following parameters are allowed: (vx, vy, vz, temp, vol, mass, Ex, Ey, Ez, Bx, By, Bz, Fx, Fy, Fz, JHrate, time). The vector ( $F x, F y, F z$ ) refers to the Lorentz force vector. A negative value returned by the *DEFINE_FUNCTION corresponds to a "dead" or inactive element. Once an element has been removed from the EM solve, it cannot return.

## Remarks:

1. When this material is used in conjunction with the battery BatMac model, then the conductivities must be adjusted by the current collector thickness ratio over the total thickness of the cell. For example, if $n_{p}$ is the number of positive current collectors, $t_{p}$ is the thickness of each individual positive current collector and Th is the total thickness of the cell, then the conductivity for the positive current collector must be scaled by: $n_{p} \times t_{p} /$ Th.

## *EM_OUTPUT

Purpose: Define the level of EM related output on the screen and in the messag file.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | MATS | MATF | SOLS | SOLF | MESH | MEM | TIMING |  |
| Type | 1 | 1 | 1 | 1 | 1 | 1 | 1 |  |
| Default | 0 | 0 | 0 | 0 | 0 | 0 | 0 |  |

## VARIABLE

MATS

MATF

SOLS

SOLF

Level of solver output to the messag file:
EQ.0: No output

## VARIABLE

## DESCRIPTION

EQ.1: Global information at each FEM iteration
EQ.2: Detailed information at each FEM iteration
MESH Controls the output of the mesh data to the d3hsp file:
EQ.0: No mesh output written.
EQ.1: Mesh info written.

MEMORY Controls the output of information about the memory used by the EM solve to the messag file:

EQ.0: No memory information written.
EQ.1: Memory information written.
TIMING Controls the output of information about the time spent in the different parts of the EM solver to the messag file

EQ.0: No timing information written.
EQ.1: Timing information written.

## *EM_OUTPUT_VTK

Purpose: Cause the output of vtk files. It can be used in applications that require two material conductivities per EM node and whose electromagnetic conductivities are defined by a $(3 \times 3)$ tensor matrix. These applications include the Randles Batmac model and the Electrophysiology Bidomain model.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | VTKTYPE | VTKT |  |  |  |  |  |  |
| Type | I | F |  |  |  |  |  |  |

VARIABLE
VTKTYPE

DESCRIPTION
Type of the vtk files output.
EQ.1: A single .vtk file
EQ.2: Parallel unstructured points data (.pvtu files), recommended in mpp executions.

VTKT Time period at which vtk files are exported

## *EM_PERMANENT_MAGNET

## *EM_PERMANENT_MAGNET

Purpose: Defines a permanent magnet.

## Card Summary:

Card 1. This card is required.

| ID | PID | MTYPE | NORTH | SOUTH | HC |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

Card 2a. This card is included if MTYPE $=3$.

| $X$ | $Y$ | $Z$ |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |

Card 2b. This card is included if MTYPE $=4$.

| NID1 | NID2 |  |  |  |  |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

Card 2c. This card is included if MTYPE $=5$ or 6 .

| $X$ | $Y$ | $Z$ | NDIVIS | AXIS | DIR/X2 | Y2 | $Z 2$ |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |

## Data Card Definitions:

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | ID | PID | MTYPE | NORTH | SOUTH | HC |  |  |
| Type | 1 | 1 | 1 | 1 | 1 | 1 |  |  |
| Default | none | none | none | none | none | none |  |  |
| VARIABLE |  | DESCRIPTION |  |  |  |  |  |  |
| ID |  | ID of the magnet |  |  |  |  |  |  |
| PID |  | Part ID |  |  |  |  |  |  |
| MTYPE |  | Magnet definition type: |  |  |  |  |  |  |

## VARIABLE

NORTH $\quad$ Set ID of the magnet north face for MTYPE $=0$ and 1
SOUTH $\quad$ Set ID of the magnet south face for MTYPE $=0$ and 1
$\mathrm{HC} \quad$ Coercive force, $H_{c}$. See Remark 1.
LT.0.0: $|\mathrm{HC}|$ refers to a load curve ID giving the coercive force as a function of time.

MTYPE = 3 Card. This card is only included for MTYPE $=3$.

| Card 2a | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | X | Y | Z |  |  |  |  |  |
| Type | F | F | F |  |  |  |  |  |
| Default | 0. | 0. | 0. |  |  |  |  |  |

*EM_PERMANENT_MAGNET

## VARIABLE

## DESCRIPTION

$\mathrm{X}, \mathrm{Y}, \mathrm{Z} \quad$ Orientation of magnetization vector
MTYPE = 4 Card. This card is only included for MTYPE $=4$.

| Card 2b | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | NID1 | NID2 |  |  |  |  |  |  |
| Type | 1 | 1 |  |  |  |  |  |  |
| Default | none | none |  |  |  |  |  |  |

## VARIABLE

NID1 / NID2

## DESCRIPTION

Two node IDs defining the magnetization vector

Magnetic Gear Card. This card is only included if MTYPE $=5$ or 6. See Figure 6-4.

| Card 2c | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | X | Y | Z | NDIVIS | AXIS | DIR/X2 | Y2 | $\mathrm{Z2}$ |
| Type | F | F | F | I | I | $\mathrm{I} / \mathrm{F}$ | F | F |
| Default | 0. | 0. | 0. | none | none | none | none | none |

## VARIABLE

X, Y, Z
NDIVIS Number of subdivisions, that is, number of magnets around the circle

AXIS Normal vector to the magnets:
EQ.1: Global $X$ axis
EQ.2: Global $Y$ axis
EQ.3: Global $Z$ axis


Figure 6-4. Example of Magnetic Gear (MTYPE $=5$ or 6 )

VARIABLE
DIR

## DESCRIPTION

Directional vector giving the location of the starting magnet / starting magnetic orientation if $\mathrm{MTYPE}=6$ :

EQ.1: Global $X$ axis
EQ.2: Global $Y$ axis
EQ.3: Global $Z$ axis

X2, Y2, Z2 Directional vector coordinates giving the starting magnet / starting magnetic orientation if MTYPE $=5$

## Remark:

1. Coercive Force. The absolute value of coercive force $H_{c}$ applied to the magnet $(\mathrm{A} / \mathrm{m})$ relates to the Residual induction $B_{r}$ by the following relation:

$$
H_{c}=B_{r} / \mu
$$

Here, $\mu$ is the magnet's permeability defined in *EM_MAT_002 using a constant relative permeability $\mu=\mu_{r} \mu_{0}$.

The coercive force can also be expressed using the energy product $B H_{\max }$ :

$$
H_{c}=2 \sqrt{\frac{B H_{\max }}{\mu}}
$$

## *EM_POINT_SET

Purpose: Create a set of points that can be used by *EM_DATABASE_POINTOUT.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | PTSID | PTSTYPE | VX | VY | VZ |  |  |  |
| Type | I | I | F | F | F |  |  |  |
| Default | 0 | 0 | 0. | 0. | 0. |  |  |  |

Include as many cards as needed. This input ends at the next keyword ("*") card.

| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | PTID | X | Y | Z | IPOS |  |  |  |
| Type | I | F | F | F | I |  |  |  |
| Default | none | none | none | none | 0 |  |  |  |

VARIABLE
PSID

PTSTYPE

VX, VY, VZ Constant velocities used when PTSTYPE $=1$
PTID Point ID
$\mathrm{X}, \mathrm{Y}, \mathrm{Z} \quad$ Point initial coordinates
IPOS Position flag (for 2D, see Remark 1):
EQ.0: The solver determines if the point is inside or outside of the conductors.

## DESCRIPTION

EQ.1: The point is outside of the conductors during the entire simulation. The solver does not check; hence a gain in computation time.

## Remarks:

1. 2D axisymmetric. If using *EM_2DAXI, notice that the conductors represent the corresponding 3D conductors.

## *EM_RANDLES_BATMAC

Purpose: define the distributed Randles circuit parameters for a Randles cell when using the batmac model. The batmac model is a macro battery model where solid elements are retained for the solid mechanics and thermal solve and where each conducting node will have its own Randles circuit associated to it. Must be used with *EM_MAT_006 or *EM_MAT_005.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | RDLID | RDLTYPE | RDLAREA | PSID |  |  |  |  |
| Type | 1 | 1 | 1 | 1 |  |  |  |  |
| Default | none | none | none | none |  |  |  |  |


| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | Q | CQ | SOCINIT | SOCTOU |  |  |  |  |
| Type | F | F | F | F |  |  |  |  |
| Default | none | none | none | none |  |  |  |  |


| Card 3.a | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | ROCHA | RODIS | R10CHA | R10DIS | C10CHA | C10DIS |  |  |
| Type | F | F | F | F | F | F |  |  |
| Default | none | none | none | none | none | none |  |  |

Optional Card. This card only needs to be defined for RDLTYPE greater than 1.

| Card 3.b | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | R20CHA | R20DIS | C20CHA | C20DIS | R30CHA | R30DIS | C30CHA | C30DIS |
| Type | F | F | F | F | F | F | F | F |
| Default | 0. | 0. | 0. | 0. | 0. | 0. | 0. | 0. |

Optional thermal card.

| Card 4 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | TEMP | FRTHER | ROTOTH | DUDT | TEMPU |  |  |  |
| Type | F | I | 1 | F | 1 |  |  |  |
| Default | 0. | 0 | 0 | none | 0 |  |  |  |

Optional SOC shift card.

| Card 5 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | USESOCS | TAU | FLCID |  |  |  |  |  |
| Type | I | F | 1 |  |  |  |  |  |
| Default | none | none | none |  |  |  |  |  |

## VARIABLE

DESCRIPTION
RDLID Id of the Randles Cell

VARIABLE
RDLTYPE

PSID
RDLAREA

Q
CQ

SOCINIT Initial state of charge of the cell.
SOCTOU Equilibrium voltage (OCV):
GE.0.0: constant value
LT.O.O: |SOCTOU| is a load curve ID defining equilibrium voltage (OCV) as a function of the state of charge (SOC).

## VARIABLE <br> R0CHA/ <br> R10CHA/ C10CHA

R0DIS/
R10DIS/
C10DIS

R20CHA/
R30CHA/ C20CHA/ C30CHA

## DESCRIPTION

$r_{0} / r_{10} / c_{10}$ when the current flows in the charge direction:
GE.0.0: constant value
LT.O.O: absolute value is a define function or table ID. See Remark 4 for the accepted define function arguments. For a define table, the circuit parameters can be made function of the SOC and temperature.
$r_{0} / r_{10} / c_{10}$ when the current flows in the discharge direction:
GE.0.0: constant value.
LT.O.O: absolute value is a define function or table ID. See Remark 4 for the accepted define function arguments. For a define table, the circuit parameters can be made function of the SOC and temperature.
$r_{20} / r_{30} / c_{20} / c_{30}$ when the current flows in the charge direction:
GE.0.0: constant value.
LT.O.O: absolute value is a define function or table ID. See Remark 4 for the accepted define function arguments. For a define table, the circuit parameters can be made function of the SOC and temperature.

R20DIS/ $\quad r_{20} / r_{30} / c_{20} / c_{30}$ when the current flows in the discharge direction:
R30DIS/
C20DIS/
C30DIS

TEMP Constant temperature value used for the Randles circuit parameters in case there is no coupling with the thermal solver $($ FRTHER $=0)$

FRTHER From Thermal :
EQ.0: The temperature used in the Randles circuit parameters is TEMP

EQ.1: The temperature used in the Randles circuit parameter is the temperature from the thermal solver.

| VARIABLE | DESCRIPTION |
| :---: | :---: |
| R0TOTH | $r_{0}$ to Thermal: |
|  | EQ.O: The joule heating in the resistance r 0 is not added to the thermal solver |
|  | EQ.1: The joule heating in the resistance $r 0$ is added to the thermal solver |
| DUDT | If negative integer, load curve ID of the reversible heat as a function of SOC. |
| TEMPU | Temperature Unit : |
|  | EQ.0: The temperature is in Celsius |
|  | EQ.1: The Temperature is in Kelvin |
| USESOCS | Use SOC shift (See Remark 2) : |
|  | EQ.0: Don't use the added SOCshift |
|  | EQ.1: Use the added SOCshift |
| TAU | Damping time in the SOCshift equation. |

TAU
FLCID

Load curve giving $f(i)$ where I is the total current in the unit cell

## Remarks:

1. Model combinations. The batmac model cannot be mixed with the solid or thick shell Randles models. It can however be used in conjunction with the meshless model.
2. Accounting for Diffusion Limitations. Sometimes, an extra term called SOCshift (or SocS) can be added at high rate discharges to account for diffusion limitations. The SOCshift is added to SOC for the calculation of the OCV $u(\mathrm{SOC}+\mathrm{SOCshift})$ and $r_{0}(\mathrm{SOC}+\mathrm{SOCshift})$. SOCshift satisfies the following equation:

$$
\frac{d(\text { SOCshift })}{d t}+\frac{\text { SOCshift }}{\tau}=\frac{f(i(t))}{\tau}
$$

with $\operatorname{SOCshift}(t=0)=0$.
3. User defined ECMs. Randles circuits are based on the finite element Robin boundary condition, acting similarly to a convection boundary condition for the heat equation. As such, the circuit equation is decomposed into a term that will enter the left-hand-side of the stiffness matrix and a term that will be added to the right-hand-side. Changing the definition of those terms allows the user to replace the Randles circuits by any type of Equivalent Circuit model (ECM). This is the purpose of RDLTYPE $=-1$ where the term entering the l.h.s is defined by a negative integer referring to a *DEFINE_FUNCTION ID in R0CHA (unit consistency: Resistance) while a negative integer in R0DIS, associated to a *DEFINE_FUNCTION will give the term entering in the r.h.s (unit consistency : current).
4. DEFINE FUNCTION variables available in EM_RANDLES keywords :

| *DEFINE <br> FUNCTIONs: <br> Variable <br> names | Randles Circuit parameters ( $\left.r_{0}, r_{10}, c_{10} e t c\right)$ | RDLTYPE $=\mathbf{- 1}$ | Internal Short | Exothermic reaction |
| :---: | :---: | :---: | :---: | :---: |
| 'time' : Current EM time | All models | All models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models |
| $\begin{aligned} & \text { 'emdt' : Current EM } \\ & \text { timestep } \end{aligned}$ | All models | All models | Solid/Tshel1/Batma c models | Solid/Tshel1/Batma c models |
| 'x_sep,y_sep,z_sep' : Positive Electrode coordinates | Solid/Tshell models | Solid/Tshell models | Solid/Tshell models | Solid/Tshell models |
| ' $x_{-}$sen,y_sen,z_sen' : Negative Electrode coordinates | Solid/Tshell models | Solid/Tshell models | Solid/Tshell models | Solid/Tshell models |
| ' $x_{-} c c p, y_{-} c c p, z_{-} c c p$ ' : Positive Current collector coordinates | Solid/Tshell/Batma c models | Solid/Tshel1/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models |
| ' $x_{-} c c n, y_{-} c c n, z_{-} c c n '$ : Negative Current collector coordinates | Solid/Tshell models | Solid/Tshell models | Solid/Tshell models | Solid/Tshell models |
| 'pres': Local pressure | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models |
| 'rho' : Local density | Solid/Tshel1/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models |
| 'vmstress': Local von Mises stress | Solid/Tshell/Batma c models | Solid/Tshel1/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models |
| 'cond' : Local electrical conductivity | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models |
| 'temp': Local Temperature | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models |
| 'tempRand' : Temperature associated to Randles Circuit | Solid/Tshel1/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models |
| 'efstrain' : Local Effective strain | Solid/Tshell/Batma c models | Solid/Tshel1/Batma c models | Solid /Batmac models | Solid /Batmac models |
| 'strainLocX/Y/Z' : Local strain in the $\mathrm{X} / \mathrm{Y} / \mathrm{Z}$ directions | Solid/Tshell/Batma c models | Solid/Tshel1/Batma c models | Solid/Batmac models | Solid /Batmac models |


| 'soc,soceff' : Local state of charge, effective state of charge | All models | All models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models |
| :---: | :---: | :---: | :---: | :---: |
| 'current' : Transverse Randles current | All models | All models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models |
| 'ocv,vc,volt,r0' : open charge voltage, damping voltage, total voltage, r0 resistance. | All models | All models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models |
| 'H_ex' : Exothermal heating power integrated over time (=exothermal heating energy) when exothermic reaction keyword is present. | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models |
| 'short': Short gives the state of the Randles circuit. If short $=0$, then the circuit is not shorted, if short $=1$, then it is shorted. | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models |
| 'ero: Ero gives the state of the erosion. If ero $=1$, then the circuit is adjacent to an element which has been eroded, ero $=0$ otherwise. | Solid/Batmac models | Solid/Batmac models | Solid/Batmac models | Solid/Batmac models |
| 'areaCircuit,areaCell, <br> areashortGlob' : local <br> Randles circuit area, Total Randles Cell area, Total Cell Shorted area. | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models |
| 'save1,save2,save3...,save10 ' : ten local variables that the user can define and that will be saved during the run and associated to each local Randles circuits. | No | Yes | No | No |

## *EM_RANDLES_EXOTHERMIC_REACTION

Purpose: This keyword allows the user to add an extra heat source term to the Randles circuit nodes in order to account for thermal runaway situations.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | AREATYPE | FUNCTID |  |  |  |  |  |  |
| Type | 1 | 1 |  |  |  |  |  |  |
| Default | none | none |  |  |  |  |  |  |

## VARIABLE

AREATYPE

## DESCRIPTION

Works the same way as RDLAREA in *EM_RANDLES_SOLID or in *EM_RANDLES_TSHELL :

EQ.1: The heat source in FUNCTID is per unit area so that, for each local Randles circuit, the result returned by FUNCTID is multiplied by a factor areaLoc (areaLoc is the local area associated to each Randles circuit while areaGlob is the area of the whole cell) (W.m²).

EQ.2: Default. The heat source in FUNCTID is for the whole cell (the whole cell is shorted), so that, for each Randles circuit, the result returned by FUNCTID is multiplied by a factor areaLoc/areaGlob (W).

EQ.3: The heat source returned by FUNCTID is taken as is in each Randles circuit (W).

FUNCTID DEFINE_FUNCTION ID giving the local heat source function of local parameters for the local Randles circuit. See Remark 1.

## Remarks:

1. DEFINE FUNCTION variables available in EM_RANDLES keywords :
$\left.\begin{array}{|c|c|c|c|}\hline \text { *DEFINE_ } & \begin{array}{c}\text { Randles Circuit } \\ \text { parameters } \\ \left(r_{0}, r_{10}, c_{10} \text { etc }\right)\end{array} & \text { RDLTYPE }=\mathbf{- 1} & \text { Internal Short }\end{array} \begin{array}{c}\text { Exothermic reac- } \\ \text { tion }\end{array}\right]$

| 'time' : Current EM time | All models | All models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models |
| :---: | :---: | :---: | :---: | :---: |
| $\begin{aligned} & \text { 'emdt : Current EM } \\ & \text { timestep } \end{aligned}$ | All models | All models | Solid/Tshel1/Batma c models | Solid/Tshel1/Batma c models |
| 'x_sep,y_sep,z_sep' : Positive Electrode coordinates | Solid/Tshell models | Solid/Tshell models | Solid/Tshell models | Solid/Tshell models |
| ' $x$ _sen, $y_{-}$sen,z_sen' : Negative Electrode coordinates | Solid/Tshell models | Solid/Tshell models | Solid/Tshell models | Solid/Tshell models |
| ' $x_{-} c c p, y_{-} c c p, z_{-} c c p^{\prime}:$ Positive Current collector coordinates | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models |
| 'x_ccn, y_ccn,z_ccn' : Negative Current collector coordinates | Solid/Tshell models | Solid/Tshell models | Solid/Tshell models | Solid/Tshell models |
| 'pres' : Local pressure | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models |
| 'rho' : Local density | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models |
| 'vmstress': Local von Mises stress | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models |
| 'cond': Local electrical conductivity | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models |
| 'temp': Local Temperature | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models |
| 'tempRand' : Temperature associated to Randles Circuit | Solid/Tshell/Batma c models | Solid/Tshel1/Batma c models | Solid/Tshel1/Batma c models | Solid/Tshel1/Batma c models |
| 'efstrain' : Local Effective strain | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid /Batmac models | Solid /Batmac models |
| 'strainLocX/Y/Z' : Local strain in the $\mathrm{X} / \mathrm{Y} / \mathrm{Z}$ directions | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid /Batmac models | Solid /Batmac models |
| 'soc,soceff' : Local state of charge, effective state of charge | All models | All models | Solid/Tshel1/Batma c models | Solid/Tshell/Batma c models |
| 'current' : Transverse <br> Randles current | All models | All models | Solid/Tshel1/Batma c models | Solid/Tshell/Batma c models |
| 'ocv,vc,volt, $r 0^{\prime} \quad$ : open charge voltage, damping voltage, total voltage, r0 resistance. | All models | All models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models |


| 'H_ex' : Exothermal heating power integrated over time (=exothermal heating energy) when exothermic reaction keyword is present. | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models |
| :---: | :---: | :---: | :---: | :---: |
| 'short': Short gives the state of the Randles circuit. If short $=0$, then the circuit is not shorted, if short $=1$, then it is shorted. | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models |
| 'ero: Ero gives the state of the erosion. If ero $=1$, then the circuit is adjacent to an element which has been eroded, ero $=0$ otherwise. | Solid/Batmac models | Solid/Batmac models | Solid/Batmac models | Solid/Batmac models |
| 'areaCircuit,areaCell, <br> areashortGlob' : local <br> Randles circuit area, Total <br> Randles Cell area, Total Cell Shorted area. | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models |
| 'save1,save2,save3...,save10 ' : ten local variables that the user can define and that will be saved during the run and associated to each local Randles circuits. | No | Yes | No | No |

## *EM_RANDLES_MESHLESS

Purpose: define the distributed Randles circuit parameters for a Randles cell which is not associated with a mesh (lumped Randles circuit).

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | RDLID | RDLTYPE |  |  |  |  |  |  |
| Type | I | 1 |  |  |  |  |  |  |
| Default | none | none |  |  |  |  |  |  |


| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | Q | CQ | SOCINIT | SOCTOU |  |  |  |  |
| Type | F | F | F | F |  |  |  |  |
| Default | none | none | none | none |  |  |  |  |


| Card 3.a | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | R0CHA | R0DIS | R10CHA | R10DIS | C10CHA | C10DIS |  |  |
| Type | F | F | F | F | F | F |  |  |
| Default | none | none | none | none | none | none |  |  |

Optional Card. This card only needs to be defined for RDLTYPE greater than 1.

| Card 3.b | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | R20CHA | R20DIS | C20CHA | C20DIS | R30CHA | R30DIS | C30CHA | C30DIS |
| Type | F | F | F | F | F | F | F | F |
| Default | 0. | 0. | 0. | 0. | 0. | 0. | 0. | 0. |

Thermal Optional card.

| Card 4 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | TEMP |  |  | DUDT | TEMPU |  |  |  |
| Type | F |  |  | F | 1 |  |  |  |
| Default | 0. |  |  | none | 0 |  |  |  |

SOC shift Optional card.

| Card 5 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | USESOCS | TAU | FLCID |  |  |  |  |  |
| Type | I | F | 1 |  |  |  |  |  |
| Default | none | none | none |  |  |  |  |  |

RDLID Id of the Randles Cell

## VARIABLE

RDLTYPE

Q
CQ

SOCINIT
SOCTOU

R0CHA/
R10CHA/ C10CHA

R0DIS/
R10DIS/
C10DIS

R20CHA/
R30CHA/
C20CHA/
C30CHA

## DESCRIPTION

Type of Randles Cell
EQ.0: 0-order Randles Cell.
EQ.1: 1-order Randles Cell.
EQ.2: 2-order Randles Cell.
EQ.3: 3-order Randles Cell.

Cell capacity.
SOC conversion factor (\%/s), known to be equal to $1 / 36$ in S.I units.

Initial state of charge of the cell.
Equilibrium voltage (OCV):
GE.0.0: constant value
LT.O.O: |SOCTOU| is a load curve ID defining equilibrium voltage (OCV) as a function of the state of charge (SOC).
$r_{0} / r_{10} / c_{10}$ when the current flows in the charge direction:
GE.0.0: constant value
LT.0.0: absolute value is a define function or table ID. See Remark 3 for the accepted define function arguments. For a define table, the circuit parameters can be made function of the SOC and temperature.
$r_{0} / r_{10} / c_{10}$ when the current flows in the discharge direction:
GE.0.0: constant value
LT.O.O: absolute value is a define function or table ID. See Remark 3 for the accepted define function arguments. For a define table, the circuit parameters can be made function of the SOC and temperature.
$r_{20} / r_{30} / c_{20} / c_{30}$ when the current flows in the charge direction:
GE.0.0: constant value
LT.O.O: absolute value is a define function or table ID. See Remark 4 for the accepted define function arguments. For a define table, the circuit parameters can be made function of the SOC and temperature.

## VARIABLE

R20DIS/
R30DIS/
C20DIS/
C30DIS

TEMP

DUDT

TEMPU Temperature Unit :
EQ.O: The temperature is in Celsius
EQ.1: The Temperature is in Kelvin
USESOCS Use SOC shift (See Remark 1) :
EQ.O: Don't use the added SOCshift
EQ.1: Use the added SOCshift
TAU
FLCID Load curve giving $f(i)$ where $I$ is the total current in the unit cell

## Remarks:

1. Accounting for Diffusion Limitations. Sometimes, an extra term called SOCshift (or SocS) can be added at high rate discharges to account for diffusion limitations. The SOCshift is added to SOC for the calculation of the OCV $u(\mathrm{SOC}+\mathrm{SOCshift})$ and $r_{0}(\mathrm{SOC}+\mathrm{SOCshift})$. SOCshift satisfies the following equation:

$$
\frac{d(\text { SOCshift })}{d t}+\frac{\text { SOCshift }}{\tau}=\frac{f(i(t))}{\tau}
$$

with $\operatorname{SOCshift}(t=0)=0$.
2. User defined ECMs. Randles circuits are based on the finite element Robin boundary condition, acting similarly to a convection boundary condition for the heat equation. As such, the circuit equation is decomposed into a term that will
enter the left-hand-side of the stiffness matrix and a term that will be added to the right-hand-side. Changing the definition of those terms allows the user to replace the Randles circuits by any type of Equivalent Circuit model (ECM). This is the purpose of RDLTYPE $=-1$ where the term entering the l.h.s is defined by a negative integer referring to a *DEFINE_FUNCTION ID in R0CHA (unit consistency: Resistance) while a negative integer in R0DIS, associated to a *DEFINE_FUNCTION will give the term entering in the r.h.s (unit consistency : current).
3. DEFINE FUNCTION variables available in EM_RANDLES keywords :

| *DEFINE <br> FUNCTIONs: <br> Variable <br> names | Randles Circuit parameters ( $r_{0}, r_{10}, c_{10}$ etc) | RDLTYPE $=\mathbf{- 1}$ | Internal Short | Exothermic reaction |
| :---: | :---: | :---: | :---: | :---: |
| 'time' : Current EM time | All models | All models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models |
| 'emdt' : Current EM timestep | All models | All models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models |
| 'x_sep,y_sep,z_sep' : Positive Electrode coordinates | Solid/Tshell models | Solid/Tshell models | Solid/Tshell models | Solid/Tshell models |
| ' $x$ _sen, $y_{-}$sen,z_sen' : Negative Electrode coordinates | Solid/Tshell models | Solid/Tshell models | Solid/Tshell models | Solid/Tshell models |
| ' $x_{-} c c p, y_{-} c c p, z_{-} c c p^{\prime}$ : Positive Current collector coordinates | Solid/Tshell/Batma c models | Solid/Tshel1/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models |
| ' $x_{-} c c n, y_{-} c c n, z_{-} c c n ': ~ N e g a-$ tive Current collector coordinates | Solid/Tshell models | Solid/Tshell models | Solid/Tshell models | Solid/Tshell models |
| 'pres': Local pressure | Solid/Tshell/Batma c models | Solid/Tshel1/Batma c models | Solid/Tshell/Batma c models | Solid/Tshel1/Batma c models |
| 'rho' : Local density | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models |
| 'vmstress': Local von Mises stress | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models |
| 'cond': Local electrical conductivity | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models |
| 'temp' : Local Temperature | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models |


| 'tempRand' : Temperature associated to Randles Circuit | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models |
| :---: | :---: | :---: | :---: | :---: |
| 'efstrain' : Local Effective strain | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid /Batmac models | Solid /Batmac models |
| 'strainLoc $X / Y / Z$ ' : Local strain in the $\mathrm{X} / \mathrm{Y} / \mathrm{Z}$ directions | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid /Batmac models | Solid /Batmac models |
| 'soc,soceff' : Local state of charge, effective state of charge | All models | All models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models |
| 'current' : Transverse <br> Randles current | All models | All models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models |
| 'ocv,vc,volt,r0' : open charge voltage, damping voltage, total voltage, r0 resistance. | All models | All models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models |
| 'H_ex': Exothermal heating power integrated over time (=exothermal heating energy) when exothermic reaction keyword is present. | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models |
| 'short': Short gives the state of the Randles circuit. If short $=0$, then the circuit is not shorted, if short $=1$, then it is shorted. | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models |
| 'ero: Ero gives the state of the erosion. If ero $=1$, then the circuit is adjacent to an element which has been eroded, ero $=0$ otherwise. | Solid/Batmac models | Solid/Batmac models | Solid/Batmac models | Solid/Batmac models |
| 'areaCircuit,areaCell, <br> areashortGlob' : local Randles circuit area, Total Randles Cell area, Total Cell Shorted area. | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models |
| 'save1,save2,save3...,save10 $'$ : ten local variables that the user can define and that will be saved during the run and associated to each local Randles circuits. | No | Yes | No | No |

## *EM_RANDLES_TSHELL

Purpose: Define the distributed Randles circuit parameters for a Randles cell when using a composite tshell mechanical model.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | RDLID | RDLTYPE | RDLAREA | PSID |  |  |  |  |
| Type | 1 | 1 | 1 | 1 |  |  |  |  |
| Default | none | none | 2 | none |  |  |  |  |


| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | Q | CQ | SOCINIT | SOCTOU |  |  |  |  |
| Type | F | F | F | F |  |  |  |  |
| Default | none | none | none | none |  |  |  |  |


| Card 3.a | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | ROCHA | RODIS | R10CHA | R10DIS | C10CHA | C10DIS |  |  |
| Type | F | F | F | F | F | F |  |  |
| Default | none | none | none | none | none | none |  |  |

Optional Card. This card only needs to be defined for RDLTYPE greater than 1.

| Card 3.b | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | R20CHA | R20DIS | C20CHA | C20DIS | R30CHA | R30DIS | C30CHA | C30DIS |
| Type | F | F | F | F | F | F | F | F |
| Default | 0. | 0. | 0. | 0. | 0. | 0. | 0. | 0. |

Optional Thermal Card.

| Card 4 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | TEMP | FRTHER | ROTOTH | DUDT | TEMPU |  |  |  |
| Type | F | I | I | F | I |  |  |  |
| Default | 0. | 0 | 0 | 0.0 | 0 |  |  |  |

Optional SOCShift Card.

| Card 5 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | USESOCS | TAU | FLCID |  |  |  |  |  |
| Type | I | F | 1 |  |  |  |  |  |
| Default | 0 | 0.0 | 0 |  |  |  |  |  |

VARIABLE
RDLID ID of the Randles Cell

CQ SOC conversion factor (\%/s), known to be equal to $1 / 36$ in SI units

RDLTYPE

PSID
RDLAREA

Q

SOCINIT
SOCTOU

R0CHA/ R10CHA/ C10CHA

## DESCRIPTION

Type of Randles Cell:
EQ.-1: User defined equivalent circuit model. See Remark 3.
EQ.0: 0-order Randles Cell
EQ.1: 1-order Randles Cell
EQ.2: 2-order Randles Cell
EQ.3: 3-order Randles Cell
Part Set ID of all the parts composing the cell
Randles Area:
EQ.1: the parameters are per unit area and will be scaled in each Randle circuit by a factor depending on the local area of the circuit. Unit consistency in S.I : Ohms times square meters.

EQ.2: the parameters are defined for the whole cell and will be scaled in each Randle circuit by a factor depending on the local area of the circuit and the global area of the cell (default). Unit consistency in S.I : Ohms.
EQ.3: the parameters are not scaled by area factors. Unit consistency in S.I: Ohms.

Q Cell capacity Initial state of charge of the cell

Equilibrium voltage (OCV):
GE.0.0: constant value
LT.O.O: |SOCTOU| is a load curve ID defining equilibrium voltage (OCV) as a function of the state of charge (SOC).
$r_{0} / r_{10} / c_{10}$ when the current flows in the charge direction:
GE.0.0: constant value
LT.O.O: absolute value is a define function or table ID. See Remark 4 for the accepted define function arguments. For a define table, the circuit parameters can be made function of the SOC and temperature.

## VARIABLE

R0DIS/
R10DIS/ C10DIS

R20CHA/
R30CHA/ C20CHA/ C30CHA

R20DIS/
R30DIS/
C20DIS/
C30DIS

TEMP

FRTHER

R0TOTH $\quad r_{0}$ to thermal:
EQ.O: the joule heating in the resistance $r_{0}$ is not added to the thermal solver.

EQ.1: the joule heating in the resistance $r_{0}$ is added to the thermal solver.

DUDT If negative integer, load curve ID of the reversible heat as a function of SOC.

## VARIABLE

TEMPU

USESOCS

TAU Damping time in the SOCshift equation (see Remark 2)
FLCID Load curve giving $f(i)$ where $i$ is the total current in the unit cell

## Remarks:

1. Sectioning of Circuit. Each part of PSID is defined by *PART_COMPOSITE_TSHELL. With this keyword for defining the part, each layer of a part can serve a different function, namely, as a current collector positive, current collector negative, separator, negative electrode, or positive electrode. A given layer's function is defined in the RDLTYPE field of *EM_MAT_001.
2. Accounting for Diffusion Limitations. Sometimes, an extra term called SOCshift (or SocS) can be added at high rate discharges to account for diffusion limitations. The SOCshift is added to SOC for the calculation of the OCV $u(\mathrm{SOC}+\mathrm{SOCshift})$ and $r_{0}(\mathrm{SOC}+\mathrm{SOCshift})$. SOCshift satisfies the following equation:

$$
\frac{d(\text { SOCshift })}{d t}+\frac{\text { SOCshift }}{\tau}=\frac{f(i(t))}{\tau}
$$

with SOCshift $(t=0)=0$.
3. User defined ECMs. Randles circuits are based on the finite element Robin boundary condition, acting similarly to a convection boundary condition for the heat equation. As such, the circuit equation is decomposed into a term that will enter the left-hand-side of the stiffness matrix and a term that will be added to the right-hand-side. Changing the definition of those terms allows the user to replace the Randles circuits by any type of Equivalent Circuit model (ECM). This is the purpose of RDLTYPE $=-1$ where the term entering the l.h.s is defined by a negative integer referring to a *DEFINE_FUNCTION ID in R0CHA (unit consistency: Resistance) while a negative integer in R0DIS, associated to a *DE-

FINE_FUNCTION will give the term entering in the r.h.s (unit consistency : current).
4. DEFINE FUNCTION variables available in EM_RANDLES keywords :

| *DEFINE <br> FUNCTIONs: <br> Variable <br> names | Randles Circuit parameters ( $\left.r_{0}, r_{10}, c_{10} e t c\right)$ | RDLTYPE $=\mathbf{- 1}$ | Internal Short | Exothermic reaction |
| :---: | :---: | :---: | :---: | :---: |
| 'time' : Current EM time | All models | All models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models |
| $\begin{aligned} & \text { 'emdt' : Current EM } \\ & \text { timestep } \end{aligned}$ | All models | All models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models |
| ' $x_{-}$sep,y_sep,z_sep' : Positive Electrode coordinates | Solid/Tshell models | Solid/Tshell models | Solid/Tshell models | Solid/Tshell models |
| ' $x_{-}$sen, $y_{-}$sen,z_sen' : Negative Electrode coordinates | Solid/Tshell models | Solid/Tshell models | Solid/Tshell models | Solid/Tshell models |
| ' $x_{-} c c p, y_{-} c c p, z_{-} c c p$ ' : Positive Current collector coordinates | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models |
| ' $x_{-} c c n, y_{-} c c n, z_{-} c c n '$ : Negative Current collector coordinates | Solid/Tshell models | Solid/Tshell models | Solid/Tshell models | Solid/Tshell models |
| 'pres': Local pressure | Solid/Tshell/Batma c models | Solid/Tshel1/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models |
| 'rho' : Local density | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models |
| 'vmstress': Local von Mises stress | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models |
| 'cond': Local electrical conductivity | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models |
| 'temp': Local Temperature | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models |
| 'tempRand' : Temperature associated to Randles Circuit | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models |
| 'efstrain' : Local Effective strain | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid /Batmac models | Solid /Batmac mod- els |


| 'strainLocX/Y/Z' : Local strain in the $X / Y / Z$ directions | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid /Batmac models | Solid /Batmac models |
| :---: | :---: | :---: | :---: | :---: |
| 'soc,soceff' : Local state of charge, effective state of charge | All models | All models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models |
| 'current' : Transverse Randles current | All models | All models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models |
| 'ocv,vc,volt,r0' : open charge voltage, damping voltage, total voltage, r0 resistance. | All models | All models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models |
| 'H_ex' : Exothermal heating power integrated over time (=exothermal heating energy) when exothermic reaction keyword is present. | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models |
| 'short': Short gives the state of the Randles circuit. If short $=0$, then the circuit is not shorted, if short $=1$, then it is shorted. | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models |
| 'ero : Ero gives the state of the erosion. If ero = 1, then the circuit is adjacent to an element which has been eroded, ero $=0$ otherwise. | Solid/Batmac models | Solid/Batmac models | Solid/Batmac models | Solid/Batmac models |
| 'areaCircuit,areaCell, <br> areashortGlob' : local <br> Randles circuit area, Total Randles Cell area, Total Cell Shorted area. | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models |
| 'save1,save2,save3...,save10 ' : ten local variables that the user can define and that will be saved during the run and associated to each local Randles circuits. | No | Yes | No | No |

## *EM_RANDLES_SHORT

Purpose: For battery cell internal short, define conditions to turn on a Randles short (replace one or several Randles circuits by resistances), and to define the value of the short resistance.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | AREATYPE | FUNCTID |  |  |  |  |  |  |
| Type | 1 | 1 |  |  |  |  |  |  |
| Default | none | None |  |  |  |  |  |  |

## VARIABLE

AREATYPE

## DESCRIPTION

Works the same way as RDLAREA in *EM_RANDLES_SOLID or in *EM_RANDLES_TSHELL :

EQ.1: The resistance inverse in FUNCTID is per unit area so that, for each local Randles circuit, the result returned by FUNCTID is multiplied by a factor 1./areaLoc (areaLoc is the local area associated to each Randles circuit while areaGlob is the area of the whole cell). Unit consistency in S.I: Ohms times square meters.

EQ.2: Default. The resistance in FUNCTID is for the whole cell (the whole cell is shorted), so that, for each Randles circuit, the result returned by FUNCTID is multiplied by a factor areaGlob/areaLoc. Unit consistency in S.I: Ohms.

EQ.3: The resistance returned by FUNCTID is taken as is for each Randles circuit. Unit consistency in S.I : Ohms.

FUNCTID

DEFINE_FUNCTION ID giving the local resistance function of local parameters for the local Randles circuit. See Remark 2.

## Remarks:

1. If the return value of the function is zero, there is no short, the Randles circuit is maintained. A positive returned value will replace the Randles circuit by the returned short resistance. In order to ensure that the short is maintained even after the original criteria is no longer met, the default positive value may be
replaced by a negative value. The solver will then take the absolute value returned and adopt it as the new short resistance in case the original short criteria is no longer met rather than reverting to a Randles circuit.
2. The parameter description is :

| *DEFINE <br> FUNCTIONs: <br> Variable <br> names: | Randles Circuit parameters $\left(r_{0}, r_{10}, c_{10}\right.$ etc) | RDLTYPE = -1 | Internal Short | Exothermic reaction |
| :---: | :---: | :---: | :---: | :---: |
| 'time' : Current EM time | All models | All models | Solid/Tshel1/Batma c models | Solid/Tshell/Batma c models |
| 'emdt' : Current EM timestep | All models | All models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models |
| 'x_sep,y_sep,z_sep' : Positive Electrode coordinates | Solid/Tshell models | Solid/Tshell models | Solid/Tshell models | Solid/Tshell models |
| ' $x_{-}$sen, $y_{-}$sen,z_sen' : Negative Electrode coordinates | Solid/Tshell models | Solid/Tshell models | Solid/Tshell models | Solid/Tshell models |
| ' $x_{-} c c p, y_{-} c c p, z_{-} c c p$ ' : Positive Current collector coordinates | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models |
| ' $x_{-} c c n, y_{-} c c n, z_{-} c c n '$ : Negative Current collector coordinates | Solid/Tshell models | Solid/Tshell models | Solid/Tshell models | Solid/Tshell models |
| 'pres' : Local pressure | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models |
| 'rho' : Local density | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models |
| 'vmstress': Local von Mises stress | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models |
| 'cond' : Local electrical conductivity | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models |
| 'temp': Local Temperature | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshel1/Batma c models | Solid/Tshell/Batma c models |
| 'tempRand' : Temperature associated to Randles Circuit | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models |
| 'efstrain' : Local Effective strain | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid /Batmac models | Solid/Batmac models |
| 'strainLoc $X / Y / Z^{\prime}$ : Local strain in the $X / Y / Z$ directions | Solid/Tshel1/Batma c models | Solid/Tshel1/Batma c models | Solid /Batmac models | Solid/Batmac models |


| 'soc,soceff' : Local state of charge, effective state of charge | All models | All models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models |
| :---: | :---: | :---: | :---: | :---: |
| 'current' : Transverse <br> Randles current | All models | All models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models |
| 'ocv,vc,volt,r0' : open charge voltage, damping voltage, total voltage, r0 resistance. | All models | All models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models |
| 'H_ex' : Exothermal heating power integrated over time (=exothermal heating energy) when exothermic reaction keyword is present. | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models |
| 'short': Short gives the state of the Randles circuit. If short $=0$, then the circuit is not shorted, if short $=1$, then it is shorted. | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models |
| 'ero: Ero gives the state of the erosion. If ero $=1$, then the circuit is adjacent to an element which has been eroded, ero $=0$ otherwise. | Solid/Batmac models | Solid/Batmac models | Solid/Batmac models | Solid/Batmac models |
| 'areaCircuit,areaCell, <br> areashortGlob' : local <br> Randles circuit area, Total <br> Randles Cell area, Total <br> Cell Shorted area. | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models |
| 'save1,save2,save3...,save10 ' : ten local variables that the user can define and that will be saved during the run and associated to each local Randles circuits. | No | Yes | No | No |

## 3. An example of a function :

*DEFINE_FUNCTION
FID (Function Id)
Float resistance_short_randle(
float time,

```
float x_ccp,float y_ccp,float z_ccp,
float x_sep,float y_sep,float z_sep,
float x_sem,float y_sem,float z_sem,
float x_ccm,float y_ccm,float z_ccm)
{ float seThick0;
seThick0 = 1.e-5;
    seThick=(sqrt(x_sep-x_sem)^^2+(y_sep-y_sem)^2+(z_sep-
z_sem)^2);
if (seThick >= seThick0) then
return -1.e-3;
else
return 1.e-2;
endif
```

In this example, as long as seThick is smaller than seThick0, no short occurs. Once seThick becomes larger than seThick0, a short occurs and the short resistance is 1.e-2. If during the run, seThick once again becomes smaller than seThick0, the short is maintained and the short resistance becomes 1.e-3. Replacing 1.e-3 by 0 . would cause the short to revert to the original Randles circuit.

## *EM_RANDLES_SOLID

Purpose: define the distributed Randles circuit parameters for a Randles cell when using a solid mechanical model.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | RDLID | RDLTYPE | RDLAREA | CCPPART | CCNPART | SEPPART | PELPART | NELPART |
| Type | 1 | 1 | 1 | 1 | 1 | 1 | 1 | 1 |
| Default | none | none | none | none | none | none | none | none |


| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | Q | CQ | SOCINIT | SOCTOU |  |  |  |  |
| Type | F | F | F | F |  |  |  |  |
| Default | none | none | none | none |  |  |  |  |


| Card 3.a | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | ROCHA | RODIS | R10CHA | R10DIS | C10CHA | C10DIS |  |  |
| Type | $F$ | $F$ | F | F | F | $F$ |  |  |
| Default | none | none | none | none | none | none |  |  |

Optional Card. This card only needs to be defined for RDLTYPE greater than 1.

| Card 3.b | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | R20CHA | R20DIS | C20CHA | C20DIS | R30CHA | R30DIS | C30CHA | C30DIS |
| Type | F | F | F | F | F | F | F | F |
| Default | 0. | 0. | 0. | 0. | 0. | 0. | 0. | 0. |

Optional Thermal card.

| Card 4 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | TEMP | FRTHER | ROTOTH | DUDT | TEMPU |  |  |  |
| Type | F | I | 1 | F | 1 |  |  |  |
| Default | 0. | 0 | 0 | None | 0 |  |  |  |

Optional SOC shift card

| Card 5 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | USESOCS | TAU | FLCID |  |  |  |  |  |
| Type | I | F | 1 |  |  |  |  |  |
| Default | none | none | none |  |  |  |  |  |

## VARIABLE

DESCRIPTION
RDLID Id of the Randles Cell

RDLTYPE

RDLAREA

CCPPART Current Collector Positive Part ID
CCNPART Current Collector Negative Part ID
SEPPART

PELPART

NELPART Negative Electrode Part ID

SOCINIT Initial state of charge of the cell.
SOCTOU

## DESCRIPTION

Type of Randles Cell:

EQ.0: 0-order Randles Cell
EQ.1: 1-order Randles Cell
EQ.2: 2-order Randles Cell
EQ.3: 3-order Randles Cell

Randles Area: meters. fault). Unit consistency in S.I : Ohms. sistency in S.I : Ohms.

Separator Part ID
Positive Electrode Part ID

Cell capacity. units.

Equilibrium voltage (OCV):

EQ.-1: User defined equivalent circuit model. See Remark 3.

EQ.1: the parameters are per unit area and will be scaled in each Randle circuit by a factor depending on the local area of the circuit. Unit consistency in S.I : Ohms times square

EQ.2: the parameters are defined for the whole cell and will be scaled in each Randle circuit by a factor depending on the local area of the circuit and the global area of the cell (de-

EQ.3: the parameters are not scaled by area factors. Unit con-

SOC conversion factor (\%/s), known to be equal to $1 / 36$ in S.I

GE.0.0: constant value
LT.0.0: |SOCTOU| is a load curve ID defining equilibrium voltage (OCV) as a function of the state of charge (SOC).

## VARIABLE <br> R0CHA/ <br> R10CHA/ C10CHA

R0DIS/
R10DIS/
C10DIS

R20CHA/
R30CHA/ C20CHA/ C30CHA

## DESCRIPTION

$r_{0} / r_{10} / c_{10}$ when the current flows in the charge direction:
GE.0.0: constant value
LT.O.O: absolute value is a define function or table ID. See Remark 4 for the accepted define function arguments. For a define table, the circuit parameters can be made function of the SOC and temperature.
$r_{0} / r_{10} / c_{10}$ when the current flows in the discharge direction:
GE.0.0: constant value
LT.O.O: absolute value is a define function or table ID. See Remark 4 for the accepted define function arguments. For a define table, the circuit parameters can be made function of the SOC and temperature.
$r_{20} / r_{30} / c_{20} / c_{30}$ when the current flows in the charge direction:
GE.0.0: constant value
LT.O.O: absolute value is a define function or table ID. See Remark 4 for the accepted define function arguments. For a define table, the circuit parameters can be made function of the SOC and temperature.

R20DIS/ $\quad r_{20} / r_{30} / c_{20} / c_{30}$ when the current flows in the discharge direction:
R30DIS/
C20DIS/
C30DIS

TEMP Constant temperature value used for the Randles circuit parameters in case there is no coupling with the thermal solver $($ FRTHER $=0)$

FRTHER From Thermal :
EQ.0: The temperature used in the Randles circuit parameters is TEMP.

EQ.1: The temperature used in the Randle circuit parameter is the temperature from the thermal solver.

## VARIABLE

R0TOTH

TEMPU

TAU
FLCID

DUDT If negative integer, load curve ID of the reversible heat as a function of SOC.

USESOCS Use SOC shift (See Remark 2) :
EQ.0: Don't use the added SOCshift
EQ.1: Use the added SOCshift

## DESCRIPTION

$r_{0}$ to Thermal:
EQ.O: The joule heating in the resistance rO is not added to the thermal solver.

EQ.1: The joule heating in the resistance r 0 is added to the thermal solver.

Temperature Unit:
EQ.0: The temperature is in Celsius
EQ.1: The Temperature is in Kelvin

Damping time in the SOCshift equation (See Remark 1)
Load curve giving $f(i)$ where I is the total current in the unit cell

## Remarks:

1. Element Normal orientation. the solid element normals must all be oriented in the positive current collector to negative current collector direction in order to detect which current collector nodes are connected to one another. Furthermore, any number of layers can be modelled but the meshes of the CCP, anode, separator, cathode, CCN must be continuous and have merged nodes at the boundaries.
2. Accounting for Diffusion Limitations. Sometimes, an extra term called SOCshift (or SocS) can be added at high rate discharges to account for diffusion limitations. The SOCshift is added to SOC for the calculation of the OCV $u(\mathrm{SOC}+\mathrm{SOCshift})$ and $r_{0}(\mathrm{SOC}+\mathrm{SOCshift})$. SOCshift satisfies the following equation:

$$
\frac{d(\text { SOCshift })}{d t}+\frac{\text { SOCshift }}{\tau}=\frac{f(i(t))}{\tau}
$$

with SOCshift $(t=0)=0$.
3. User defined ECMs. Randles circuits are based on the finite element Robin boundary condition, acting similarly to a convection boundary condition for the heat equation. As such, the circuit equation is decomposed into a term that will enter the left-hand-side of the stiffness matrix and a term that will be added to the right-hand-side. Changing the definition of those terms allows the user to replace the Randles circuits by any type of Equivalent Circuit model (ECM). This is the purpose of RDLTYPE $=-1$ where the term entering the l.h.s is defined by a negative integer referring to a *DEFINE_FUNCTION ID in R0CHA (unit consistency: Resistance) while a negative integer in R0DIS, associated to a *DEFINE_FUNCTION will give the term entering in the r.h.s (unit consistency : current).
4. DEFINE FUNCTION variables available in EM_RANDLES keywords :

| *DEFINE <br> FUNCTIONs: <br> Variable <br> names | Randles Circuit parameters ( $\left.r_{0}, r_{10}, c_{10} e t c\right)$ | RDLTYPE $=\mathbf{- 1}$ | Internal Short | Exothermic reaction |
| :---: | :---: | :---: | :---: | :---: |
| 'time' : Current EM time | All models | All models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models |
| 'emdt' : Current EM timestep | All models | All models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models |
| 'x_sep,y_sep,z_sep' : Positive Electrode coordinates | Solid/Tshell models | Solid/Tshell models | Solid/Tshell models | Solid/Tshell models |
| ' $x$ _sen,y_sen,z_sen' : Negative Electrode coordinates | Solid/Tshell models | Solid/Tshell models | Solid/Tshell models | Solid/Tshell models |
| ' $x_{-} c c p, y_{-} c c p, z_{-} c c p^{\prime}:$ Positive Current collector coordinates | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models |
| ' $x_{-} c c n, y_{-} c c n, z_{-} c c n '$ : Negative Current collector coordinates | Solid/Tshell models | Solid/Tshell models | Solid/Tshell models | Solid/Tshell models |
| 'pres': Local pressure | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models |
| 'rho' : Local density | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models |
| 'vmstress': Local von Mises stress | Solid/Tshell/Batma c models | Solid/Tshel1/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models |
| 'cond' : Local electrical conductivity | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models |


| 'temp' : Local Temperature | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models |
| :---: | :---: | :---: | :---: | :---: |
| 'tempRand' : Temperature associated to Randles Circuit | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models |
| 'efstrain' : Local Effective strain | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid /Batmac models | Solid /Batmac models |
| 'strainLocX/Y/Z' : Local strain in the $X / Y / Z$ directions | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid /Batmac models | Solid /Batmac models |
| 'soc,soceff' : Local state of charge, effective state of charge | All models | All models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models |
| 'current' : Transverse Randles current | All models | All models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models |
| 'ocv,vc,volt,r0' : open charge voltage, damping voltage, total voltage, r0 resistance. | All models | All models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models |
| 'H_ex' : Exothermal heating power integrated over time (=exothermal heating energy) when exothermic reaction keyword is present. | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models |
| 'short': Short gives the state of the Randles circuit. If short $=0$, then the circuit is not shorted, if short $=1$, then it is shorted. | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models |
| 'ero: Ero gives the state of the erosion. If ero $=1$, then the circuit is adjacent to an element which has been eroded, ero $=0$ otherwise. | Solid/Batmac models | Solid/Batmac models | Solid/Batmac models | Solid/Batmac models |
| 'areaCircuit,areaCell, <br> areashortGlob' : local <br> Randles circuit area, Total <br> Randles Cell area, Total <br> Cell Shorted area. | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models | Solid/Tshell/Batma c models |
| 'save1,save2,save3...,save10 ' : ten local variables that the user can define and that will be saved during the run and associated to each local Randles circuits. | No | Yes | No | No |

## *EM_ROTATION_AXIS

Purpose: Define a rotation axis for the EM solver. This is used with the 2D axisymmetric solver. The axis is defined by a point and a direction.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | XP | YP | ZP | XD | YD | ZD | NUMSEC |  |
| Type | F | F | F | F | F | F | I |  |
| Default | none | none | none | none | none | none | none |  |

VARIABLE
XP, YP, ZP
XD, YD, ZD
NUMSEC

## DESCRIPTION

$x, y$, and $z$ coordinates of the point
$x, y$, and $z$ components of direction of the axis
Number of Sectors. This field gives the ratio of the full circle to the angular extension of the mesh. This has to be a power of two. For example, NUMSEC $=4$ means that the mesh of the part represents one fourth of the total circle. If NUMSEC $=0$ for *EM_2DAXI, the solver will replace it with this value.

## *EM_SOLVER_BEM

Purpose: Define the type of linear solver and pre-conditioner as well as tolerance for the EM_BEM solve.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | RELTOL | MAXITE | STYPE | PRECON | USELAST | NCYLBEM |  |  |
| Type | 1 | 1 | 1 | 1 | 1 | 1 |  |  |
| Default | $10^{-6}$ | 1000 | 2 | 2 | 1 | 5000 |  |  |

VARIABLE
RELTOL

MAXITER Maximum number of iterations for the iterative solvers
STYPE Solver type:
EQ.1: Direct solve. The matrices are considered dense.
EQ.2: Pre-conditioned gradient method (PCG). This method allows for block matrices with low-rank blocks and thus reduces the memory used.
EQ.3: GMRES method. This method allows for block matrices with low-rank blocks and thus reduces the memory used. The GMRES option only works in serial for now.

PRECON
Preconditioner type for PCG or GMRES iterative solves:
EQ.O: No preconditioner
EQ.1: Diagonal line
EQ.2: Diagonal block
EQ.3: Broad diagonal including all neighbor faces
EQ.4: LLT factorization

## VARIABLE

USELAST

NCYLBEM Number of electromagnetism cycles between the recalculation of BEM matrices. If negative, the absolute value refers to a load curve giving the number of cycles as a function of time. See Remarks 2 and 3 .

## Remarks:

1. USELAST. Using USELAST = 1 can save many iterations in the subsequent solves if the vector solution of the present solve can be assumed to be nearly parallel to the vector solution of the previous solve, as usually happens in timedomain eddy-current problems.
2. Moving conductors. Since the BEM matrices depend on (and only on) the surface node coordinates of the conductors, recalculating them when the conductors are moving is important. NCYLBEM controls the frequency with which they are updated. Note that very small values, for example, NCYLBEM $=1$, should, generally, be avoided since this calculation involves a high computational cost. However, when two conductors are moving and in contact with each other, we recommend recalculating the matrices at every time step.
3. Order of precedence for NCYLBEM. You can set NCYLBEM with *EM_CONTROL_SOLUTION, *EM_SOLVER_FEM/BEM, and *EM_CONTROL. *EM_CONTROL_SOLUTION has the highest priority for setting this field. If NCYLBEM is the default value on *EM_CONTROL_SOLUTION, LS-DYNA looks at the setting of NCYLBEM on *EM_SOLVER_BEM. If left as default on this keyword, LS-DYNA looks at the setting of NCYLBEM on *EM_CONTROL.

## *EM_SOLVER_BEMMAT

Purpose: Define the type of BEM matrices as well as the way they are assembled.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | MATID |  |  |  |  |  |  | RELTOL |
| Type | I |  |  |  |  |  |  |  |
| Default | none |  |  |  |  |  |  |  |

VARIABLE

## DESCRIPTION

MATID
Defines which BEM matrix the card refers to:
EQ.1: P matrix
EQ.2: Q matrix
EQ.3: W matrix

RELTOL Relative tolerance on the sub-blocks of the matrix when doing low rank approximations. The user should try to decrease these tolerances if the results are not accurate enough. More memory will then be needed.

## Remarks:

1. The $\mathbf{W}$ matrix only exists when the monolithic solver is activated (see *EM_SOLVER_FEMBEM_MONOLITHIC).

## *EM_SOLVER_FEM

Purpose: Define some parameters for the EM FEM solver.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | RELTOL | MAXITE | STYPE | PRECON | USELAST | NCYLFEM |  |  |
| Type | 1 | 1 | 1 | 1 | 1 | 1 |  |  |
| Default | $10^{-3}$ | 1000 | 1 | 1 | 1 | 5000 |  |  |

## VARIABLE

RELTOL

MAXITER
STYPE

PRECON

USELAST

NCYLFEM

## DESCRIPTION

Relative tolerance for the iterative solvers (PCG). If the results are not accurate enough, try decreasing this tolerance. More iterations will then be needed.

Maximum number of iterations for iterative solvers
Solver type:
EQ.1: Direct solve
EQ.2: Pre-conditioned Gradient Method (PCG)

Preconditioner type for PCG.
EQ.0: No preconditioner
EQ.1: Diagonal line
This is used only for iterative solvers (PCG).
EQ.-1: Starts from 0 as the initial solution of the linear system.
EQ.1: Starts from the previous solution normalized by the right-hand-side change. See Remark 1.

Number of electromagnetism cycles between the recalculation of FEM matrices. If negative, the absolute value refers to a load curve giving NCYCLFEM as a function of time. See Remarks 2 and 3.

## Remarks:

1. Starting from the previous solution. Using USELAST = 1 can save many iterations in the subsequent solves if the vector solution of the present solve can be assumed to be nearly parallel to the vector solution of the previous solve, as usually happens in time-domain eddy-current problems.
2. NCYLFEM. When the conductor parts are deforming or undergoing changes in their EM material properties (conductivity for example), the FEM matrices should be recalculated more often, so NCYLFEM may need to be changed.
3. Order of precedence for NCYLFEM. You can set NCYLFEM with *EM_CONTROL_SOLUTION, *EM_SOLVER_FEM, and *EM_CONTROL. *EM_CONTROL_SOLUTION has the highest priority for setting this field. If NCYLFEM is the default value on *EM_CONTROL_SOLUTION, LS-DYNA looks at the setting of NCYLFEM on *EM_SOLVER_FEM. If left as default on this keyword, LSDYNA looks at the setting on *EM_CONTROL.

## *EM_SOLVER_FEMBEM

Purpose: Define some parameters for the standard coupling between the EM_FEM and EM_BEM solvers.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | RELTOL | MAXITE | FORCON |  |  |  |  |  |
| Type | F | 1 | 1 |  |  |  |  |  |
| Default | $10^{-2}$ | 50 | 0 |  |  |  |  |  |

VARIABLE
RELTOL

MAXITER Maximal number of iterations
FORCON tolerance will, however, require more iterations.

Force convergence:

## DESCRIPTION

Relative tolerance for the FEM/BEM system solve. If the results are not accurate enough, try decreasing this tolerance. A smaller

EQ.0: The code stops with an error if no convergence.
EQ.1: The code continues to the next time step even if the RELTOL convergence criteria has not been reached.

## Remarks:

This keyword couples the FEM and BEM systems with the Richardson method. At each time step, the solver will iterate between the FEM and the BEM system until reaching convergence (based on the choice of RELTOL and MAXITER). The cost for this solve is low. However, to ensure stability, we recommend imposing a limit on the timestep based on the characteristic diffusion time (See *EM_CONTROL_TIMESTEP). Furthermore, it can be unstable whenever magnetic materials are involved (conductor's permeability different than vacuum permeability). The monolithic solver invoked with *EM_SOLVER_FEMBEM_MONOLITHIC aims to remove those two limitations by solving both the FEM and BEM systems in one single monolithic bloc. For such cases, it is, therefore, the recommended choice (See *EM_SOLVER_FEMBEM_MONOLITHIC).

## *EM_SOLVER_FEMBEM_MONOLITHIC

Purpose: Replaces *EM_SOLVER_FEMBEM and turns on the monolithic FEM-BEM solver.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | MTYPE | STYPE | ABSTOL | RELTOL | MAXIT |  |  |  |
| Type | I | I | F | F | I |  |  |  |
| Default | 0 | 0 | $10^{-6}$ | $10^{-4}$ | 500 |  |  |  |

## VARIABLE

MTYPE

STYPE Solver type:
EQ.0: MINRES iterative solver
EQ.1: GMRES iterative solver

## Remarks:

The monolithic solver aims to overcome the limitations of the classic Richardson iterative coupling between the FEM and BEM systems. The monolithic solver offers better stability for large timesteps and for simulations involving ferromagnetic materials. We recommend this method whenever the *EM_MAT_002 keyword is present.

## *ICFD

The keyword *ICFD covers all the different options available in the incompressible fluid solver. The keyword cards in this section are defined in alphabetical order:

```
*ICFD_BOUNDARY_CONJ_HEAT
*ICFD_BOUNDARY_CONVECTION_TEMP
*ICFD_BOUNDARY_FLUX_TEMP
*ICFD_BOUNDARY_FREESLIP
*ICFD_BOUNDARY_FSI
*ICFD_BOUNDARY_FSI_EXCLUDE
*ICFD_BOUNDARY_FSI_FIXED
*ICFD_BOUNDARY_FSWAVE
*ICFD_BOUNDARY_GROUND
*ICFD_BOUNDARY_NAVIERSLIP
*ICFD_BOUNDARY_NONSLIP
*ICFD_BOUNDARY_PERIODIC
*ICFD_BOUNDARY_PRESCRIBED_LEVELSET
*ICFD_BOUNDARY_PRESCRIBED_MOVEMESH
*ICFD_BOUNDARY_PRESCRIBED_PRE
*ICFD_BOUNDARY_PRESCRIBED_SPTRANSP_CONC
*ICFD_BOUNDARY_PRESCRIBED_TEMP
*ICFD_BOUNDARY_PRESCRIBED_TURBULENCE
*ICFD_BOUNDARY_PRESCRIBED_VEL
*ICFD_BOUNDARY_WEAKVEL
*ICFD_BOUNDARY_WINDKESSEL
*ICFD_CONTROL_ADAPT
```


## *ICFD

*ICFD_CONTROL_ADAPT_SIZE
*ICFD_CONTROL_BACKFLOW
*ICFD_CONTROL_CONJ
*ICFD_CONTROL_DEM_COUPLING
*ICFD_CONTROL_EMBEDSHELL
*ICFD_CONTROL_FSI
*ICFD_CONTROL_GAP
*ICFD_CONTROL_GENERAL
*ICFD_CONTROL_IMPOSED_MOVE
*ICFD_CONTROL_LEVELSET
*ICFD_CONTROL_LOAD
*ICFD_CONTROL_MESH
*ICFD_CONTROL_MESH_MOV
*ICFD_CONTROL_MONOLITHIC
*ICFD_CONTROL_OUTPUT
*ICFD_CONTROL_OUTPUT_SUBDOM
*ICFD_CONTROL_OUTPUT_VAR
*ICFD_CONTROL_PARTITION
*ICFD_CONTROL_POROUS
*ICFD_CONTROL_STEADY
*ICFD_CONTROL_SURFMESH
*ICFD_CONTROL_TAVERAGE
*ICFD_CONTROL_TIME
*ICFD_CONTROL_TRANSIENT
*ICFD_CONTROL_TURB_SYNTHESIS
*ICFD_CONTROL_TURBULENCE
*ICFD_DATABASE_AVERAGE
*ICFD_DATABASE_DRAG
*ICFD_DATABASE_FLUX
*ICFD_DATABASE_FLUXSURF
*ICFD_DATABASE_FORCE_DEM
*ICFD_DATABASE_HTC
*ICFD_DATABASE_NODEAVG
*ICFD_DATABASE_NODOUT
*ICFD_DATABASE_NTEMPOUT
*ICFD_DATABASE_POINTAVG
*ICFD_DATABASE_POINTOUT
*ICFD_DATABASE_RESIDUALS
*ICFD_DATABASE_SSOUT
*ICFD_DATABASE_SSOUT_EXCLUDE
*ICFD_DATABASE_TEMP
*ICFD_DATABASE_TIMESTEP
*ICFD_DATABASE_UINDEX
*ICFD_DATABASE_WETNESS
*ICFD_DEFINE_HEATSOURCE
*ICFD_DEFINE_NONINERTIAL
*ICFD_DEFINE_POINT
*ICFD_DEFINE_RESIDENCETIMESOURCE
*ICFD_DEFINE_SOURCE
*ICFD_DEFINE_SPTRANSPSOURCE
*ICFD_DEFINE_TURBSOURCE
*ICFD_DEFINE_WAVE_DAMPING

## *ICFD

*ICFD_INITIAL
*ICFD_INITIAL_LEVELSET
*ICFD_INITIAL_SPTRANSP
*ICFD_INITIAL_TEMPNODE
*ICFD_INITIAL_TURBULENCE
*ICFD_MAT
*ICFD_MODEL_NONNEWT
*ICFD_MODEL_POROUS
*ICFD_MODEL_SPECIES_TRANSPORT
*ICFD_PART
*ICFD_PART_VOL
*ICFD_SECTION
*ICFD_SET_NODE
*ICFD_SOLVER_SPLIT
*ICFD_SOLVER_TOL_FSI
*ICFD_SOLVER_TOL_LSET
*ICFD_SOLVER_TOL_MMOV
*ICFD_SOLVER_TOL_MOM
*ICFD_SOLVER_TOL_MONOLITHIC
*ICFD_SOLVER_TOL_PRE
*ICFD_SOLVER_TOL_TEMP
*ICFD_BOUNDARY_CONJ_HEAT
Purpose: Specify which boundary of the fluid domain will exchange heat with the solid.
Include as many cards as needed. This input ends at the next keyword ("*") card.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | PID | CTYPE | VAL | SFLCID |  |  |  |  |
| Type | 1 | 1 | F | 1 |  |  |  |  |
| Default | none | 0 | 0. | 0 |  |  |  |  |

## VARIABLE

PID
CTYPE

VAL Optional temperature drop if CTYPE $=0$ or interface heat transfer coefficient if CTYPE $=1$ (high value by default to ensure perfect contact).

SFLCID Load curve ID used to describe scale factor on VAL value as a function of time; see *DEFINE_CURVE, *DEFINE_CURVE_FUNCTION, or *DEFINE_FUNCTION. If a *DEFINE_FUNCTION is used, the following parameters are allowed: $f(x, y, z, v x, v y, v z$, temp, pres, time).

## *ICFD_BOUNDARY_CONVECTION_TEMP

Purpose: Impose a heat transfer coefficient on the boundary expressed as $h=\frac{q}{T_{s}-T_{b}}$
Include as many cards as needed. This input ends at the next keyword ("*") card.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | PID | HLCID | HSF | TBLCID | TBSF |  |  |  |
| Type | I | I | F | I | F |  |  |  |
| Default | none | none | 1. | none | 1.0 |  |  |  |

## VARIABLE

PID
HLCID

HSF Load curve scale factor applied on the heat transfer coefficient value. $($ default $=1.0)$

TBLCID

## DESCRIPTION

PID for a fluid surface.
Load curve ID to describe the heat transfer coefficient value versus time, see *DEFINE_CURVE,*DEFINE_CURVE_FUNCTION or *DEFINE_FUNCTION. If a DEFINE_FUNCTION is used, the following parameters are allowed: $f(x, y, z, v x, v y, v z$, temp, pres, time $)$.

Load curve ID to describe the environment (i.e bulk) temperature value versus time, see *DEFINE_CURVE,*DEFINE_CURVE_FUNCTION or *DEFINE_FUNCTION. If a DEFINE_FUNCTION is used, the following parameters are allowed: $f(x, y, z, v x, v y, v z$, temp, pres, time $)$.

TBSF Load curve scale factor applied on the environment value. (default $=1.0$ )
*ICFD_BOUNDARY_FLUX_TEMP
Purpose: Impose a heat flux on the boundary expressed as $q=-k \nabla T$
Include as many cards as needed. This input ends at the next keyword ("*") card.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | PID | LCID | SF | DEATH | BIRTH |  |  |  |
| Type | I | I | F | F | F |  |  |  |
| Default | none | none | 1. | $1 . \mathrm{E}+28$ | 0.0 |  |  |  |

## VARIABLE

PID
LCID

SF Load curve scale factor. $($ default $=1.0)$

Time at which the imposed motion/constraint is removed:
EQ.0.0: default set to 10 e 28

BIRTH Time at which the imposed pressure is activated starting from the initial abscissa value of the curve
*ICFD_BOUNDARY_FREESLIP
Purpose: Specify the fluid boundary with free-slip boundary condition.
Include as many cards as needed. This input ends at the next keyword ("*") card.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | PID |  |  |  |  |  |  |  |
| Type | I |  |  |  |  |  |  |  |
| Default | none |  |  |  |  |  |  |  |

## VARIABLE

PID

## DESCRIPTION

PID of the fluid surface where a free-slip boundary condition is applied.

## *ICFD_BOUNDARY_FSI

Purpose: This keyword defines which fluid surfaces will be considered in contact with the solid surfaces for fluid-structure interaction (FSI) analysis. This keyword should not be defined if *ICFD_CONTROL_FSI is not defined.

Include as many cards as needed. This input ends at the next keyword ("*") card.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | PID |  |  |  |  |  |  |  |
| Type | 1 |  |  |  |  |  |  |  |
| Default | none |  |  |  |  |  |  |  |
| VARIABLE |  |  |  |  |  |  |  |  |

PID
PID of the fluid surface in contact with the solid domain.

## *ICFD_BOUNDARY_FSI_EXCLUDE

Purpose: Specify which solid part IDs are excluded from the FSI search. No forces coming from the fluid will be transmitted to those parts.

Include as many cards as needed. This input ends at the next keyword ("*") card.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | PID |  |  |  |  |  |  |  |
| Type | 1 |  |  |  |  |  |  |  |
| Default | none |  |  |  |  |  |  |  |

VARIABLE
PID

## DESCRIPTION

Part ID of a part from the solid mechanics problem that is to be excluded from the FSI analysis

## *ICFD_BOUNDARY_FSI_FIXED

Purpose: Define fixed fluid surfaces that will be considered for contact with the solid surfaces for FSI. This keyword is similar to *ICFD_BOUNDARY_FSI, except the fluid surface cannot move. This restriction allows a solid surface to "slide" over the fluid and for the exchange of data, such as temperature in CHT applications. This keyword should not be defined if *ICFD_CONTROL_FSI is not defined.

Include as many cards as needed. This input ends at the next keyword ("*") card.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | PID |  |  |  |  |  |  |  |
| Type | 1 |  |  |  |  |  |  |  |
| Default | none |  |  |  |  |  |  |  |

VARIABLE
PID

## DESCRIPTION

Part ID of the fluid surface that can be considered for contact with the solid domain

## *ICFD_BOUNDARY_FSWAVE

Purpose: Impose a wave inflow boundary condition.

## Card Summary:

Card Sets. Include as many sets of the following cards as needed. This input ends with the next keyword ("*") card.

Card 1. This card is required.

| PID | WTYPE | HO | WAMP | WLENG | WMAX | SFLCID | WANG |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |

Card 2. This card is included if $\mathrm{WTYPE}=7$.

| WPEAK |  |  |  |  |  |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

## Data Card Definitions:

Include as many of this card and/or sets of this card with the next as needed. This input ends at the next keyword ("*") card.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | PID | WTYPE | H0 | WAMP | WLENG | WMAX | SFLCID | WANG |
| Type | I | I | F | F | F | F | । | F |
| Default | none | none | none | none | none | none | $\downarrow$ | none |

## VARIABLE

## DESCRIPTION

PID
WTYPE

Wave type:
EQ.1: Stokes wave of first order
EQ.2: Stokes wave of second order
EQ.3: Stokes wave of fifth order
EQ.4: Solitary wave
EQ.5: Irregular waves using JONSWAP spectrum

## VARIABLE

H0

WAMP

WLENG

WMAX

SFLCID

WANG

## DESCRIPTION

EQ.6: Irregular waves using One Parameter Pierson-Moskowitz spectrum

EQ.7: Irregular waves using Two Parameter Pierson-Moskowitz spectrum

Water level (from the bottom of the channel) for the unperturbed condition

Wave amplitude or height for WTYPE = 1 and 4 . Significant wave height for WTYPE $=5,6$, and 7 .

WTYPE.LE.2: Wave length
WTYPE.EQ.3: Wave period
WTYPE.EQ.4: Not used
WTYPE.GE.5: Minimum wave frequency in spectrum (rad/sec)

Maximum wave frequency in spectrum ( $\mathrm{rad} / \mathrm{sec}$ ) for $\mathrm{WTYPE}=5$, 6, and 7. Angle between the boundary and the incident waves (in degrees) for WTYPE $=3$.

Scale factor LCID on the wave amplitude for WTYPE $=1,2$ and 3. Number of wave modes $($ default $=1024)$ for $\mathrm{WTYPE}=5,6$, and 7 .

Angle between incoming wave direction and $x$-axis for $z$ - and $y$ aligned gravity vector, or angle between incoming wave direction and $y$-axis for $x$-aligned gravity vector.

Card included for WTYPE $=7$ only

| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | WPEAK |  |  |  |  |  |  |  |
| Type | F |  |  |  |  |  |  |  |
| Default | none |  |  |  |  |  |  |  |

VARIABLE

## Remarks:

1. Peak Wave Frequency for WTYPE = 6. For the irregular waves using the One Parameter Pierson-Moskowitz spectrum, the peak wave frequency in the spectrum LS-DYNA calculates the peak wave frequency with:

$$
0.4 \sqrt{\frac{g}{H_{s}}}
$$

Here $g$ the gravity and $H_{s}$ is the significant wave height input with WAMP.

## *ICFD_BOUNDARY_GROUND

Purpose: Specify the fluid boundary with a ground boundary condition. The ground boundary condition is similar to the nonslip boundary condition except that it will keep $\mathrm{V}=0$ in all circumstances, even if the surface nodes are moving. This is useful in cases where the nodes are allowed to move or translate (using ICFD_BOUNDARY_PRESCRIBED_MOVEMESH for example) but those displacements are only to accommodate for mesh movement and do not correspond to a physical motion.

Include as many cards as needed. This input ends at the next keyword ("*") card.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | PID |  |  |  |  |  |  |  |
| Type | I |  |  |  |  |  |  |  |
| Default | none |  |  |  |  |  |  |  |
| PARIABLE |  |  |  |  |  |  |  |  |

PID

| PID of the fluid surface where a ground boundary condition is ap- |
| :--- |
| plied. |

## *ICFD_BOUNDARY_NAVIERSLIP

Purpose: Specify the fluid boundary with the Navier slip boundary condition.
Include as many cards as needed. This input ends at the next keyword ("*") card.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | PID | FRIC |  |  |  |  |  |  |
| Type | I | F |  |  |  |  |  |  |
| Default | none | 0. |  |  |  |  |  |  |

VARIABLE
PID

FRIC Friction coefficient. If a negative value is entered, it will refer to a load curve ID used to describe the friction coefficient value as a function of time; see *DEFINE_CURVE, *DEFINE_CURVE_FUNCTION, or *DEFINE_FUNCTION. If a *DEFINE_FUNCTION is used, the following parameters are allowed: $f(x, y, z, v x$, vy, vz, temp, pres, time).

## Remarks:

1. About the Navier condition. It is similar to the regular free slip condition, except a local source term is added based on the choice of the friction coefficient. Since this extra shear is a function of the fluid's velocity, a smaller time step may be needed to reduce the effects of the introduced nonlinearity.

Purpose: Specify the fluid boundary with a non-slip boundary condition.
Include as many cards as needed. This input ends at the next keyword ("*") card.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | PID |  |  |  |  |  |  |  |
| Type | 1 |  |  |  |  |  |  |  |
| Default | none |  |  |  |  |  |  |  |
| VARIABLE |  |  |  |  |  |  |  |  |

PID
PID of the fluid surface where a non-slip boundary condition is applied.

## *ICFD_BOUNDARY_PERIODIC

Purpose: Impose various kinds of constraints between two fluid surfaces.
Include as many cards as needed. This input ends at the next keyword ("*") card.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | PID | PTYPE | PID2 | PDLCID | AXE | PTID | ANGLE |  |
| Type | 1 | 1 | 1 | 1 | 1 | 1 | $F$ |  |
| Default | none | none | none | none | none | none | none |  |

## VARIABLE

## DESCRIPTION

PID
Part ID for a fluid surface

## PTYPE Boundary type:

EQ.1: Periodic rotation boundary condition
EQ.2: Periodic reflective boundary condition
EQ.3: Sliding mesh boundary condition

PID2

PDLCID

AXE

Part ID for the second surface mesh. The boundary condition selected with PTYPE will be applied between PID and PID2. See Remark 1.

Optional load curve ID to describe the pressure drop value between PID and PID2 as a function of time. This curve can be specified with *DEFINE_CURVE,*DEFINE_CURVE_FUNCTION, or *DEFINE_FUNCTION. For *DEFINE_FUNCTION, the following parameters are allowed: $f(x, y, z, v x, v y, t e m p, p r e s, t i m e)$.

The meaning of AXE depends on PTYPE. It only applies for PTYPE $=1$ and 3 . For the periodic rotation boundary condition (PTYPE = 1):

EQ.1: Rotation around $X$-axis
EQ.2: Rotation around $Y$-axis
EQ.3: Rotation around Z-axis
For the sliding mesh boundary condition $($ PTYPE $=3)$ :

VARIABLE

PTID

ANGLE $\quad$ Rotation angle for PTYPE $=1$. Characterizes contact distance for PTYPE $=3$ and $A X E \neq 0$.

## Remarks:

1. Selection of PID and PID2. When the two meshes are of different densities, we recommend selecting the finer mesh to be PID and the coarser mesh to be PID2.

## *ICFD_BOUNDARY_PRESCRIBED_LEVELSET

Purpose: Prescribe the fluid height on a boundary.
Include as many cards as needed. This input ends at the next keyword ("*") card.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | PID | PTID | AXE |  |  |  |  |  |
| Type | 1 | 1 | 1 |  |  |  |  |  |
| Default | none | none | none |  |  |  |  |  |

## VARIABLE

PID
PTID Point ID specifying the origin of the fluid surface. See *ICFD_DEFINE_POINT.

AXE

## DESCRIPTION

PID of the fluid surface where a fluid height will be imposed.

Global axis specifying the direction of the fluid:

EQ.1: X-axis
EQ.2: $Y$-axis
EQ.3: Z-axis
*ICFD_BOUNDARY_PRESCRIBED_MOVEMESH
Purpose: Allows the node of a fluid surface to translate in certain directions using an ALE approach. This is useful in piston type applications or can also be used in certain cases to avoid big mesh deformation.

Include as many cards as needed. This input ends at the next keyword ("*") card.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | PID | dofx | dofy | dofz |  |  |  |  |
| Type | 1 | 1 | 1 | 1 |  |  |  |  |
| Default | none | 1 | 1 | 1 |  |  |  |  |

VARIABLE
PID
dofx, dofy, dofz

## DESCRIPTION

PID for a fluid surface.
Degrees of freedom in the $\mathrm{X}, \mathrm{Y}$ and Z directions :
EQ.O: degree of freedom left free (Surface nodes can translate in the chosen direction)

EQ.1: prescribed degree of freedom (Surface nodes are blocked)

## *ICFD_BOUNDARY_PRESCRIBED_PRE

Purpose: Impose a fluid pressure on the boundary.
Include as many cards as needed. This input ends at the next keyword ("*") card.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | PID | LCID | SF | DEATH | BIRTH |  |  |  |
| Type | I | I | F | F | F |  |  |  |
| Default | none | none | 1. | $1 . E+28$ | 0.0 |  |  |  |

## VARIABLE

PID
LCID

SF $\quad$ Load curve scale factor. $($ default $=1.0)$

Time at which the imposed motion/constraint is removed:
EQ.0.0: default set to 10E28

BIRTH Time at which the imposed pressure is activated starting from the initial abscissa value of the curve
*ICFD_BOUNDARY_PRESCRIBED_SPTRANSP_CONC
Purpose: Specify the concentration of the transported species at the boundaries.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | PID | LCID |  |  |  |  |  |  |
| Type | 1 | 1 |  |  |  |  |  |  |
| Default | none | none |  |  |  |  |  |  |

## VARIABLE

PID
LCID

## DESCRIPTION

Part ID of the boundary with the concentration
Load curve ID for the curve giving the concentration at the boundary as a function of time

## *ICFD_BOUNDARY_PRESCRIBED_TEMP

Purpose: Impose a fluid temperature on the boundary.
Include as many cards as needed. This input ends at the next keyword ("*") card.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | PID | LCID | SF | DEATH | BIRTH |  |  |  |
| Type | I | I | F | F | F |  |  |  |
| Default | none | none | 1. | $1 . E+28$ | 0.0 |  |  |  |

## VARIABLE

PID
LCID

SF $\quad$ Load curve scale factor. $($ default $=1.0)$

## DESCRIPTION

PID for a fluid surface. rameters are allowed: $f(x, y, z, v x, v y, v z$, temp, pres, time $)$.

Load curve ID to describe the temperature value versus time; see *DEFINE_CURVE, *DEFINE_CURVE_FUNCTION or *DEFINE_FUNCTION. . If a DEFINE_FUNCTION is used, the following pa-

DEATH
Time at which the imposed temperature is removed:
EQ.0.0: default set to 10E28
BIRTH Time at which the imposed temperature is activated starting from the initial abscissa value of the curve

## *ICFD_BOUNDARY_PRESCRIBED_TURBULENCE

Purpose: Optional keyword for strongly imposing turbulence quantities when you select a RANS turbulence model. See *ICFD_CONTROL_TURBULENCE. This keyword is intended for modifying the default boundary conditions at the inlet.

Include as many cards as needed. This input ends at the next keyword ("*") card.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | PID | VTYPE | IMP | LCID | KS | CS |  |  |
| Type | I | I | I | I | F | F |  |  |
| Default | none | none | 0 | none | 0. | 0. |  |  |

## VARIABLE

PID
VTYPE

IMP

LCID

## DESCRIPTION

PID for a fluid surface
Variable type:
EQ.1: Turbulence kinetic energy (see Remark 1)
EQ.2: Turbulence dissipation rate (see Remark 2)
EQ.3: Specific dissipation rate (see Remark 3)
EQ.4: Modified turbulence viscosity (see Remark 4)
Imposition method:
EQ.0: Direct imposition through value specified by LCID
EQ.1: Using turbulence intensity specified by LCID if VTYPE $=1$ (see Remark 1). Using turbulence length scale specified by LCID if VTYPE $=2,3$, or 4 (see Remarks 2, 3, and 4).

EQ.2: Using turbulence viscosity ratio specified by LCID. Only available for VTYPE $=2$ and 3 . See Remarks 2 and 3 .

Load curve ID to describe the variable value as a function of time; see *DEFINE_CURVE, *DEFINE_CURVE_FUNCTION or *DEFINE_FUNCTION. If a *DEFINE_FUNCTION is used, the following parameters are allowed: $f(x, y, z, v x, v y, v z$, temp, pres, time, $k$, e, mut).

KS/CS

> Roughness physical height and roughness constant. When defined, the global values of *ICFD_CONTROL_TURBULENCE are replaced for this surface part.

## Remarks:

1. Turbulence Kinetic Energy. At the inlet, the relationship between the turbulence kinetic energy, $k$, and the turbulence intensity, $I$, is given by:

$$
k=\frac{3}{2}\left(U_{\mathrm{avg}}^{2} I^{2}\right) .
$$

By default, the solver uses an inlet intensity of 0.05 ( $5 \%$ ).
2. Turbulence Dissipation Rate. At the inlet, if you specify the turbulent dissipation rate using a length scale, $l$, the following relationship will be used:

$$
\epsilon=C_{\mu}^{3 / 4} \frac{k^{3 / 2}}{l} .
$$

By default, the solver estimates a length scale based on the total height of the channel.

If you specify the turbulent viscosity ratio, $r=\mu_{t} / \mu$, the following relationship will be used:

$$
\epsilon=\rho C_{\mu} \frac{k^{2}}{\mu r} .
$$

3. Specific Dissipation Rate. At the inlet, if you specify the specific dissipation rate using a length scale, $l$, the following relationship will be used:

$$
\omega=C_{\mu}{ }^{-1 / 4} \frac{k^{1 / 2}}{l} .
$$

By default, the solver estimates a length scale based on the total height of the channel.

If you specify the turbulence viscosity ratio, $r=\mu_{t} / \mu$, the following relationship will be used:

$$
\omega=\rho \frac{k}{\mu r}
$$

4. Modified Turbulent Viscosity. At the inlet, the relationship between the modified turbulent viscosity, $\tilde{v}$, and the length scale, $l$, is given by:

$$
\tilde{v}=0.05 \sqrt{\frac{3}{2}}\left(U_{\mathrm{avg}} l\right) .
$$

## *ICFD_BOUNDARY_PRESCRIBED_VEL

Purpose: Impose the fluid velocity on the boundary.
Include as many cards as needed. This input ends at the next keyword ("*") card.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | PID | DOF | VAD | LCID | SF | VID | DEATH | BIRTH |
| Type | I | I | 1 | I | F | I | F | F |
| Default | none | none | 1 | none | 1. | 0 | $10^{28}$ | 0.0 |

## VARIABLE

## DESCRIPTION

PID
DOF

VAD Velocity flag:
EQ.1: Linear velocity
EQ.2: Angular velocity
EQ.3: Parabolic velocity profile
EQ.4: Activates synthetic turbulent field on part. See *ICFD_CONTROL_TURB_SYNTHESIS.

LCID Load curve ID used to describe motion value versus time, see *DEFINE_CURVE, *DEFINE_CURVE_FUNCTION, or *DEFINE_FUNCTION. If a DEFINE_FUNCTION is used, the following parameters are allowed: $f(x, y, z, v x, v y, v z$, temp, pres, time). For steady state the motion value is a function of the number of iterations instead of time.

Load curve scale factor. $($ default $=1.0)$

## VARIABLE

VID

DEATH Time at which the imposed motion/constraint is removed:
EQ.O.O: default set to $10^{28}$

BIRTH Time at which the imposed motion/constraint is activated starting from the initial abscissa value of the curve

## *ICFD_BOUNDARY_WEAKVEL

Purpose: Specify the fluid boundary with a non-slip boundary condition which is imposed in a weak form.

Include as many cards as needed. This input ends at the next keyword ("*") card.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | PID |  |  |  |  |  |  |  |
| Type | I |  |  |  |  |  |  |  |
| Default | none |  |  |  |  |  |  |  |

VARIABLE
PID

## DESCRIPTION

Part ID of the fluid surface where a weak non-slip boundary condition is applied

## *ICFD_BOUNDARY_WINDKESSEL

Purpose: Impose the pressure function with circuit parameters where an analogy is made between the pressure and scalar potential and between the flux and the current intensity. Such conditions are frequently encountered in hemodynamics.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | PID | WTYPE | R1 | C1 | R2 | L1 |  |  |
| Type | I | I | F | F | F | F |  |  |
| Default | none | none | 0. | 0. | 0. | 0. |  |  |

Optional CV Circuit Card. This card is read if WTYPE $=3$ or 4. It is optional.

| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | P2LCID | C2 | R3 |  |  |  |  |  |
| Type | I | F | F |  |  |  |  |  |
| Default | 0 | 0. | 0. |  |  |  |  |  |

## VARIABLE

PID
WTYPE

## DESCRIPTION

PID for a fluid surface
Circuit type (see Figures 7-1 and 7-2):
EQ.1: Windkessel circuit
EQ.2: Windkessel circuit with inverted flux
EQ.3: CV-type circuit
EQ.4: CV-type circuit with inverted flux

R1/C1/L1/R
2/C2/R3
P2LCID

Parameters (resistances, inductances, capacities) for the different circuits

ID of load curve describing the behavior of $P_{2}(t)$ function of time for CV-type circuit.


Figure 7-1. Windkessel circuit


Figure 7-2. CV-type circuit

## *ICFD_CONTROL_ADAPT

Purpose: This keyword will activate the adaptive mesh refinement feature. The solver will use an a-posteriori error estimator to compute a new mesh size bounded by the user to satisfy a maximum perceptual global error.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | MINH | MAXH | ERR | MTH | NIT | VAR |  | KIS |
| Type | F | F | F | I | I | 1 |  | 1 |
| Default | none | none | 1. | 0 | 0 | 0 |  | 0 |

## VARIABLE

MINH

MAXH
ERR

MTH

NIT

VAR

## DESCRIPTION

Minimum mesh size allowed to the mesh generator. The resulting mesh will not have an element smaller than MINH even if the minimum size does not satisfy the maximum error.

Maximum mesh size
Maximum perceptual error allowed in the whole domain
Specify if the mesh size is computed based on function error or gradient error:

EQ.0: Function error
EQ.1: Gradient error
Number of iterations before a remeshing is forced:
GT.0: Number of iterations before a forced remeshing.
EQ.0: Do not remesh.
LT.O: $|\mathrm{NIT}|$ is a load curve ID giving the number iterations before a remeshing as a function of time.

Specify which variable is taken into account for the error calculation:

EQ.O: Velocity, pressure and levelset function are taken into account.

EQ.1: Remove the levelset function from the error calculation.

VARIABLE
DESCRIPTION
EQ.2: Remove the pressure from the error calculation.
EQ.3: Remove both pressure and levelset function from the error calculation. Only the fluid velocity will, therefore, remain.

KIS Keep initial mesh size:
EQ.0: Turned off: The remeshing process will ignore the initial mesh size in the volume.

EQ.1: Turned on: Whenever a remeshing occurs, the new local mesh size will not be allowed to be substantially coarser than the one from the previous mesh. The object is to diminish the excessive coarsening that can occur between two remeshes.
*ICFD_CONTROL_ADAPT_SIZE
Purpose: This keyword controls the re-meshing of elements taking into account the element quality and distortion in contrast to the default algorithm which only checks for inverted elements.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | ASIZE | NIT | KIS |  |  |  |  |  |
| Type | 1 | 1 | 1 |  |  |  |  |  |
| Default | 0 | none | 0 |  |  |  |  |  |

## VARIABLE

ASIZE

NIT Number of iterations before a re-meshing is forced. If a negative integer is entered, then a load curve function of time will be used to define NIT.

KIS

Keep initial mesh size:
EQ.0: Turned Off: The remeshing process will ignore the initial mesh size in the volume.

EQ.1: Turned on: Whenever a remeshing occurs, the new local mesh size will not be allowed to be substantially coarser than the one from the previous mesh. The object is to diminish the excessive coarsening that can occur between two remeshes.

## *ICFD_CONTROL_BACKFLOW

Purpose: Modify default values for backflow stabilization.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | BFOR | SF |  |  |  |  |  |  |
| Type | I | F |  |  |  |  |  |  |
| Default | 0 | 1.0 |  |  |  |  |  |  |

## VARIABLE

BFOR

DESCRIPTION
Set the backflow stabilization formulation:
EQ.0: Default stabilization dependent on spatial velocity gradients.

EQ.1: The stabilization adds a temporal velocity gradient which could be necessary for added stabilization.

SF Scale factor to increase the stabilization if needed. A very small value larger than zero minimizes the effect.

## *ICFD_CONTROL_CONJ

Purpose: This keyword allows to pick between the different coupling methods for conjugate heat transfer applications

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | CTYPE |  |  |  |  |  |  | TSF |
| Type | 1 |  |  |  |  |  |  |  |
| Default | 0 |  |  |  |  |  |  | none |

VARIABLE
CTYPE

TSF

## DESCRIPTION

Indicates the thermal coupling type.
EQ.0: Robust and accurate monolithic coupling where the temperature field are solved simultaneously between the fluid and the structure.

EQ.1: Weak thermal coupling. The fluid passes the heat flux to the solid at the fluid-structure interface and the solid returns the temperature which is applied as a Dirichlet condition.

Thermal Speedup Factor. This factor multiplies all thermal parameters present in the heat equation with units of time in the denominator (e.g., thermal conductivity, convection heat transfer coefficients). It is used to artificially time scale the thermal problem. A negative value will refer to a time dependent load curve.

## Remarks:

1.The keyword ICFD_BOUNDARY_CONJ_HEAT is ignored if CTYPE = 1 but the keyword ICFD_BOUNDARY_FSI is needed in all thermal coupling cases.
*ICFD_CONTROL_DEM_COUPLING
Purpose: Activate coupling between the ICFD and DEM solvers.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | CTYPE | BT | DT | SF | MAXVEL | DTYPE | SFF | FORM |
| Type | I | F | F | F | F | 1 | $F$ | 1 |
| Default | 0 | 0. | $10^{28}$ | 1. | none | 0 | 1.0 | 0 |

VARIABLE
CTYPE

BT Birth time for the DEM coupling
DT Death time for the DEM coupling
SF Scale factor applied to the force transmitted by the fluid to the structure

MAXVEL Maximal fluid velocity that can be used for the calculation of the fluid force passed on to the DEM particle. This is to avoid having spurious velocities in the fluid causing very high and unrealistic forces on the DEM particles which may lead to a crash.

DTYPE Drag calculation type:
EQ.O: Constant $C_{d}$ value 0.5 scaled by SF
EQ.1: Formula for $C_{d}$ calculation from Cheng 2009 based on the local Reynolds number value scaled by SF. See Remark 1.

SFF Scale factor applied to the force transmitted by the structure to the fluid

## VARIABLE

FORM Type of formulation used in the coupling:
EQ.O: The force at the particle is based on a velocity drag value.
EQ.1: The force is computed using the fluid pressure gradient.

## Remarks:

1. Coefficient of Drag by Cheng 2009. The calculation for $C_{d}$ is:

$$
C_{d}=\frac{24}{\operatorname{Re}} \times(1+0.27 \times \operatorname{Re})^{0.43}+0.47 \times\left(1-\exp \left(-0.04 \times \operatorname{Re}^{0.38}\right)\right)
$$

See Cheng 2009 for details.

## References:

[1] Cheng, N.-S., "Comparison of formulas for drag coefficient and settling velocity of spherical particles," Powder Technology, 189, 395-398 (2009).

## *ICFD_CONTROL_EMBEDSHELL

Purpose: This keyword allows the user to control specific options related to the use of the keyword MESH_EMBEDSHELL.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | GTYPE | DIST | TPS |  |  |  |  |  |
| Type | 1 | F | 1 |  |  |  |  |  |
| Default | 0 | 0.1 | 0 |  |  |  |  |  |

DIST Distance value if GTYPE $=1$ or scale factor value if GTYPE $=0$.

## VARIABLE

GTYPE

TPS

## DESCRIPTION

Gap type. Defines the criteria for selecting a distance to build the gap between the embedded nodes and the newly generated :

EQ.O: Automatic and based on the surface mesh size multiplied by a scale factor given by DIST. Default method.

EQ.1: Specific gap size given by the user and defined by DIST.

Triple Point Seal. Allows to control the fluid escape through triple points

EQ.0: Off.
EQ.1: On. The triple points of embedded shells in contact to walls or among each other are sealed and no flow goes through them.

## *ICFD_CONTROL_FSI

Purpose: This keyword modifies default values for the fluid-structure interaction coupling algorithm.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | OWC | BT | DT | IDC | LDICSF | XPROJ |  |  |
| Type | I | F | F | F | I | 1 |  |  |
| Default | 0 | 0 | $10^{28}$ | 0.25 | 0 | 0 |  |  |


| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | NSUB |  |  |  |  |  |  |  |
| Type | I |  |  |  |  |  |  |  |
| Default | none |  |  |  |  |  |  |  |

VARIABLE
OWC

## DESCRIPTION

Indicates the coupling direction to the solver:
EQ.O: Two-way coupling. Loads and displacements are transferred across the FSI interface, and the full non-linear problem is solved. It gives weak FSI coupling when coupled to explicit mechanical solver and strong FSI coupling when coupled to implicit mechanical solver.
EQ.1: One-way coupling. The solid mechanics solver transfers displacements to the fluid solver.

EQ.2: One-way coupling. The fluid solver transfers stresses to the solid mechanics solver.

EQ.3: Two-way coupling. It causes weak coupling (no sub-stepping) with the implicit mechanical solver.

## VARIABLE

BT

DT

IDC
LCIDSF Optional load curve ID to apply a scaling factor on the forces transferred to the solid:

GT.0: Load curve ID for scale factor as a function of iterations
LT.0: |LCIDSF| is a load curve ID for scale factor as a function of time.

XPROJ Projection of the nodes of the CFD domain that are at the FSI interface onto the structural mesh (see Remark 2):

## EQ.0: No projection

EQ.1: Projection
Optional limit on the number of FSI fluid subiterations. This avoids the sometimes unneeded excessive number of FSI subiterations when the fluid and very light structures (like parachutes) develop a resonance-like mode inside the FSI subiterations (coupling iterations).


Figure 0-1. Geometry of FSI contact.

## Remarks:

1. Detecting Fluid-Solid Interaction. One of the criteria to automatically detect the fluid and solid surfaces that will interact in FSI problems is the distance $d$ between a fluid (solid) node and a solid (fluid) element, respectively:

$$
d \leq \operatorname{IDC} \times \min (h, H)
$$

where $h$ is the size of the fluid mesh, $H$ is the size of the solid mechanics mesh, and IDC is a detection coefficient criteria with IDC $=0.25$ by default. In the majority of cases, this default value is sufficient to ensure FSI interaction. However, it can happen in special cases that the fluid and solid geometries have curvatures that differ too much (such as pipe flows in conjugate heat transfer applications). In such cases, a bigger IDC value may be needed. This flag should be handled with care.
2. Rotation and Projection of Nodes. $\mathrm{XPROJ}=1$ is recommended for cases with rotation.

## *ICFD_CONTROL_GAP

Purpose: Activate the gap closure treatment that deals with flow blockage when surfaces come into contact. In ICFD, surface meshes are not allowed to collide or penetrate each other. Rather, activation of the gap closure feature triggers a flow blockage between two surfaces in close proximity based on a user defined contact distance.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | HGAP | MTYPE | DVCL | RDVCL |  |  |  |  |
| Type | F | 1 | 1 | 1 |  |  |  |  |
| Default | none | 0 | 0 | 0 |  |  |  |  |


| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | PID1 | PID2 | PID3 | PID4 | PID5 | PID6 | PID7 | PID8 |
| Type | 1 | 1 | 1 | 1 | 1 | 1 | 1 | 1 |
| Default | none | none | none | none | none | none | none | none |

## VARIABLE

HGAP

PIDn Part IDs of the surfaces involved in the gap closure treatment
*ICFD_CONTROL_GENERAL
Purpose: Specify the type of CFD analysis.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | ATYPE | MTYPE | DVCL | RDVCL |  |  |  |  |
| Type | 1 | 1 | 1 | 1 |  |  |  |  |
| Default | 0 | 0 | 0 | 0 |  |  |  |  |

VARIABLE
ATYPE

MTYPE

DVCL

RDVCL

Divergence cleaning flag:
EQ.O: Initialize the solution with divergence cleaning (default)
EQ.1: No divergence cleaning
EQ.2: Initial divergence cleaning using potential flow
EQ.4: Initial divergence cleaning using steady state solver

RDVCL Remeshing divergence cleaning:
EQ.0: No divergence cleaning after remesh (default)
EQ.1: Divergence cleaning after each remeshing step

## *ICFD_CONTROL_IMPOSED_MOVE

Purpose: Impose a velocity on specific ICFD parts or on the whole volume mesh. Global translation, global rotation, and local rotation components can be defined and combined. This keyword can be used to save calculation time in certain applications, such as sloshing, where the modeling of the whole fluid box and the solving of the consequent FSI problem are not necessarily needed.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | PID | LCVX | LCVY | LCVZ | VADT |  |  |  |
| Type | 1 | 1 | 1 | 1 | 1 |  |  |  |
| Default | none | none | none | none | 0 |  |  |  |

Optional Card. Rotational velocity components using Euler angles (See Remark 1).

| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | ALPHAL | BETAL | GAMMAL | ALPHAG | BETAG | GAMMAG | VADR |  |
| Type | 1 | 1 | 1 | 1 | 1 | 1 | 1 |  |
| Default | 0 | 0 | 0 | 0 | 0 | 0 | 0 |  |

Optional Card. Local reference frame definition if ALPHAL, BETAL or GAMMAL used.

| Card 3 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | PTID | X 1 | Y 1 | Z 1 | X 2 | Y 2 | $\mathrm{Z2}$ |  |
| Type | I | F | F | F | F | F | F |  |
| Default | 0 | 1. | 0. | 0. | 0. | 1. | 0. |  |

Optional Card. When defined, Cards 2 and 3 are ignored. With this card, rotation is imposed around a point using the velocity of a second point.

| Card 4 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | PTIDO | AXE | NID |  |  |  |  |  |
| Type | 1 | 1 | 1 |  |  |  |  |  |
| Default | 0 | 0 | 0 |  |  |  |  |  |

## VARIABLE

PID

LCVX, LCVY, LCVZ

VADT

ALPHAL, BETAL, GAMMAL

ALPHAG, BETAG, GAMMAG

VADR

PTID

## DESCRIPTION

Part ID. This can be any part ID referenced in *ICFD_PART or *ICFD_PART_VOL. If PID $=0$, then the whole volume mesh will be used.

Load curve IDs for the velocity/displacements in the three global directions ( $X, Y, Z$ ). To use a *DEFINE_FUNCTION, see Remark 4.

Velocity/displacements flag for translation components:
EQ.O: Prescribe velocity
EQ.1: Prescribe displacements
Load curves IDs for the three Euler angle rotational velocities/displacements in the local reference frame (see Remarks 1 and 2). To use a *DEFINE_FUNCTION, see Remark 4.

Load curve IDs for the three Euler angle rotational velocities/displacements in the global reference frame (see Remarks 1 and 2). To use a *DEFINE_FUNCTION, see Remark 4.

Velocity / displacements flag for rotation components:
EQ.O: Prescribe velocity
EQ.1: Prescribe displacements
Point ID for the origin of the local reference frame. If not defined, the barycenter of the volume mesh will be used


Figure 7-3. A rotation represented by Euler angles $(\alpha, \beta, \gamma)$ using $\mathbf{Z}(\alpha) \mathbf{X}(\beta) \mathbf{Z}(\gamma)$ intrinsic rotations.

## VARIABLE

X1, Y1, Z1

X2, Y2, Z2

PTIDO

AXE

NID

DESCRIPTION
Three components of the local reference X1 axis. If not defined, the global $X$ axis will be used. See Remark 2.

Three components of the local reference X 2 axis. If not defined, the global $Y$ axis will be used. See Remark 2.

Point ID (See *ICFD_DEFINE_POINT) for the center of rotation.

Rotation axis:
EQ.1: X-axis
EQ.2: $Y$-axis
EQ.3: Z-axis

ICFD Surface Node ID for the rotational velocity. If the node is static, no rotation will occur. See Remark 3.

## Remarks:

1. Rotations. Any target orientation can be reached starting from a known reference orientation using a specific sequence of intrinsic rotations whose magnitudes are the Euler angles $(\alpha, \beta, \gamma)$. Equivalently, any rotation matrix $\mathbf{R}$ can be decomposed as a product of three elemental rotation matrices. For instance:

$$
\mathbf{R}=\mathbf{X}(\alpha) \mathbf{Y}(\beta) \mathbf{Z}(\gamma)
$$

However, different definition of the elemental rotation matrices $(x, y, z)$ and their multiplication order can be adopted. The ICFD solver uses the following approach and rotation matrix:

$$
\mathbf{Z}(\alpha) \mathbf{X}(\beta) \mathbf{Z}(\gamma)=\left[\begin{array}{ccc}
c_{\alpha} c_{\gamma}-c_{\beta} s_{\alpha} s_{\gamma} & -c_{\beta} c_{\gamma} s_{\alpha}-c_{\alpha} s_{\gamma} & s_{\alpha} s_{\beta} \\
c_{\gamma} s_{\alpha}+c_{\alpha} c_{\beta} s_{\gamma} & c_{\alpha} c_{\beta} c_{\gamma}-s_{\alpha} s_{\gamma} & -c_{\alpha} s_{\beta} \\
s_{\beta} s_{\gamma} & c_{\gamma} s_{\beta} & c_{\beta}
\end{array}\right]
$$

where $\boldsymbol{X}(\alpha), \boldsymbol{Y}(\beta)$, and $\boldsymbol{Z}(\gamma)$ are the matrices representing the elemental rotations about the axes $(x, y, z), s_{\alpha}=\sin (\alpha)$, and $c_{\beta}=\cos (\beta)$.
2. Local Coordinate Systems. It is possible to have the ICFD parts or ICFD_PART_VOLs rotate around the global reference frame but also to define and use a local reference frame by defining its point of origin and two of its vectors $\mathbf{v}_{1}=$ ( $\mathrm{X} 1, \mathrm{Y} 1, \mathrm{Z} 1$ ) and $\mathrm{v}_{2}=(\mathrm{X} 2, \mathrm{Y} 2, \mathrm{Z} 2)$ (note that $\mathbf{v}_{1}$ and $\mathbf{v}_{2}$ should be orthogonal). The third vector is, then, in the direction of $\mathbf{v}_{1} \times \mathbf{v}_{\mathbf{2}}$. See Figure 7-3.
3. Purpose of NID. We developed this feature for rotating problems involving FSI and sliding mesh. For example, the airflow can push the blades of a wind turbine and the rotation of the sliding mesh can be prescribed as function of the blade rotation speed.
4. *DEFINE_FUNCTION. For each of the load curves, a *DEFINE_FUNCTION can be used. If a *DEFINE_FUNCTION is used, the following parameters are allowed: $f(x, y, z, v x, v y, v z$, temp, visc, pres, time, $d t)$.

## *ICFD_CONTROL_LEVELSET

Purpose: Modify default values for the level set solver.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | LSRST | LSINL | LSMTH |  |  |  |  |  |
| Type | 1 | 1 | F |  |  |  |  |  |
| Default | 20 | 0 | 0.0 |  |  |  |  |  |

## VARIABLE

LSRST

LSINL

LSMTH

## DESCRIPTION

This parameter specifies how often the level set distance function is re-initialized. The default value is every 20 time steps.

Set the level set to be positive at a velocity inlet:
EQ.0: Default is let the level set algorithm compute the value.
EQ.1: Force a positive level set value at the inlet.
Scale factor for level set smoothness.
GT.0.0: Add smoothness to the free surface. Small values like 0.1 are reasonable. The optimal value could be problem dependent.

## *ICFD_CONTROL_LOAD

Purpose: This keyword resets the body load in the ICFD solver to zero, while leaving the body load unchanged for the solid mechanics solver. It is useful in problems where the gravity acceleration may be neglected for the fluid problem, but not for the solid mechanics problem.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | ABL |  |  |  |  |  |  |  |
| Type | 1 |  |  |  |  |  |  |  |
| Default | 1 |  |  |  |  |  |  |  |

VARIABLE
ABL

## DESCRIPTION

EQ.0: the body load provided in *LOAD_BODY is reset to zero only for the fluid analysis.

## *ICFD_CONTROL_MESH

Purpose: Modify default values for automatic volume mesh generation.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | MGSF |  | MSTRAT | 2DSTRUC | NRMSH |  |  |  |
| Type | F |  | 1 | 1 | 1 |  |  |  |
| Default | 1.41 |  | 0 | 0 | 0 |  |  |  |

Optional card. This card is optional.

| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | AVER | SFBL |  |  |  |  |  |  |
| Type | I | F |  |  |  |  |  |  |
| Default | 14 | 1.0 |  |  |  |  |  |  |

## VARIABLE

MGSF

MSTRAT

2DSTRUC

## DESCRIPTION

Mesh Growth Scale Factor. It specifies the maximum mesh size that the volume mesher is allowed to use when generating the volume mesh based on the mesh surface element sizes defined in *MESH_SURFACE_ELEMENT. See Remark 1.

Mesh generation strategy (see Remark 2):
EQ.O: Mesh generation based on Delaunay criteria
EQ.1: Mesh generation based on octree

Flag to decide between an unstructured mesh generation strategy in 2D or a structured mesh strategy:

EQ.O: Structured mesh
EQ.1: Unstructured mesh

## VARIABLE

NRMSH

AVER

SFBL

Scale factor that controls the speed of boundary layer inflation. If the boundary layer is much larger in size than the surface mesh, we recommend a value less than one.

## Remarks:

1. MGSF. For MGSF, values between 1 and 2 are allowed. Values closer to 1 will result in a finer volume mesh ( 1 means the volume mesh is not allowed to be coarser than the element size from the closest surface meshes) and values closer to 2 will result in a coarser volume mesh ( 2 means the volume can use elements as much as twice as coarse as those from the closest surface mesh). MGSF has a fixed value of 1 in 2D.
2. Mesh Generation Strategy. The default mesh generation strategy (based on Delaunay criteria) yields a linear interpolation of the mesh size between two surfaces facing each other whereas the octree-based generation strategy allows for the sizes of the elements to remain close to the element surface mesh size over a longer distance. This octree strategy can be useful for creating a smoother transition in configurations where two surface meshes facing each other have very distinct sizes.
3. NRMSH. If you know in advance that no remeshing will occur during the analysis, then setting NRMSH to 1 may be useful as it will free up space used to back up the mesh and consequently lower memory consumption.
4. Version. Version 14 is the default version used for the ICFD solver automatic volume mesher. Version 16 is now supported and available as option. In some cases, it can yield an approximatively $20 \%$ mesh generation speed gain.

## *ICFD_CONTROL_MESH_MOV

Purpose: Choose the type of algorithm for mesh movement.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | MMSH | LIM_ITER | RELTOL |  |  |  |  |  |
| Type | 1 | 1 | F |  |  |  |  |  |
| Default | 2 | 100 | $10^{-3}$ |  |  |  |  |  |

## VARIABLE

MMSH

## DESCRIPTION

Mesh motion selector:
EQ.-1: Completely shuts off any mesh movement
EQ.1: Mesh moves based on the distance to moving walls.
EQ.2: Mesh moves by solving a linear elasticity problem using the element sizes as stiffness (default).

EQ.3: Mesh uses a Laplacian smoothing with stiffness on edges and from node to opposite faces. Very robust, but costly.

EQ.4: Full Lagrangian. The mesh moves with the velocity of the flow.

EQ.11: Mesh moves using an implicit ball-vertex spring method.

EQ.22: Mesh moves by solving a linear elasticity problem using a constant size. This can be useful to avoid large distortions in rotating problems that involve large discrepancies in mesh sizes (typically in cases involving boundary layer mesh).

LIM_ITER Maximum number of linear solver iterations for the ball-vertex linear system

RELTOL
Relative tolerance to use as a stopping criterion for the ball-vertex method iterative linear solver (conjugate gradient solver with diagonal scaling preconditioner)

## *ICFD_CONTROL_MONOLITHIC

Purpose: This keyword allows to choose between the Fractional Step Solver and the Monolithic Solver.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | SID |  |  |  |  |  |  |  |
| Type | 1 |  |  |  |  |  |  |  |
| Default | 0 |  |  |  |  |  |  |  |

## VARIABLE

SID

## DESCRIPTION

Solver ID :
EQ.0: Fractional Step Solver. Default.
EQ.1: Monolithic Solver.

## *ICFD_CONTROL_OUTPUT

Purpose: This keyword modifies default values for screen and file outputs related to this fluid solver only.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | MSGL | OUTL | DTOUT | LSPPOUT |  | ITOUT |  |  |
| Type | I | I | F | I |  | 1 |  |  |
| Default | 0 | 0 | 0 | 0 |  | 0 |  |  |

## Optional Card.

| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | PITOUT |  |  |  |  |  |  |  |
| Type | 1 |  |  |  |  |  |  |  |
| Default | none |  |  |  |  |  |  |  |

## VARIABLE

MSGL

## DESCRIPTION

Message level.
EQ.O: only time step information is output.
EQ.1: first level solver information.
EQ.2: full output information with details about linear algebra and convergence steps.
EQ.4: full output information is also copied to the messag file.

Output the fluid results in other file formats apart from d3plot.
EQ.0: only d3plot output
EQ.2: output a file with mesh statistics and the fluid results in OpenDX format. A directory named dx will be created in the work directory where the output files will be written.
EQ.6: output a file with mesh statistics and the fluid results in VTK format readable by Paraview. A directory named vtk will be created in the work directory where the output files will be written.

EQ.7: output a file with mesh statistic and the fluid results in VTU format readable by Paraview. A directory named vtk will be created in the work directory where the output files will be written.

EQ.10: output a file with mesh statistic and the fluid results in Fieldview ASCII format. . A directory named fv will be created in the work directory where the output files will be written. Only available in 3D.

EQ.11: output a file with mesh statistic and the fluid results in Fieldview binary format. . A directory named fv will be created in the work directory where the output files will be written. Only available in 3D.

DTOUT Time interval to print the output when OUTL is different than 0.

LSPPOUT

ITOUT

EQ.0: no LSPP output is produced.
EQ.1: outputs a file with the automatically created fluid volume mesh in a format compatible for LSPP at each remesh. Also outputs the fluid volume mesh in a format compatible with a subsequent ICFD analysis.
EQ.3: Outputs the fluid volume mesh in a format compatible with a subsequent ICFD analysis at each DTOUT.

Iteration interval to print the output, including the d3plot files when the steady state solver is selected (See ICFD_CONTROL_GENER$A L)$.

## VARIABLE

PITOUT

## DESCRIPTION

Pressure iteration limit output. If the number of pressure iterations in the fractional step solve goes above PITOUT, an extra d3plot will be dumped. This is mainly a debugging feature which can help the user identify problematic areas in the model which often precede a divergence.

## *ICFD_CONTROL_OUTPUT_SUBDOM

Purpose: Defines a specific zone that should be output in the format specified by the ICFD_CONTROL_OUTPUT card rather than the whole domain.

Shape Control. First card specifies the shape of the output sub domain.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | SNAME |  |  |  |  |  |  |  |
| Type | A |  |  |  |  |  |  |  |
| Default | none |  |  |  |  |  |  |  |

Box Case. Card 2 for Sname = box

| Cards 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | PMINX | PMINY | PMINZ | PMAXX | PMAXY | PMAXZ |  |  |
| Type | F | F | F | F | F | F |  |  |
| Default | none | none | none | none | none | none |  |  |

Sphere Case. Card 2 for Sname $=$ sphere

| Cards 3 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | RADIUS | CENTERX | CENTERY | CENTERZ |  |  |  |  |
| Type | F | F | F | F |  |  |  |  |
| Default | none | none | none | none |  |  |  |  |

Cylinder Case. Card 2 for Sname = cylinder

| Cards 4 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | Radius | PMINX | PMINY | PMAXZ | PMAXX | PMAXY | PMAXZ |  |
| Type | F | F | F | F | F | F | F |  |
| Default | none | none | none | none | none | none | none |  |

## VARIABLE

SNAME
PMINX, Y, Z]
PMAX[X, Y, Z]
CEN-
TER[X, Y, Z]
RADIUS Radius of the sphere if SNAME is sphere or of the cross section disk if SNAME is cylinder.

## *ICFD_CONTROL_OUTPUT_VAR

Purpose: This keyword allows the user to turn off the output of certain CFD variables to reduce the size of the d3plot files.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | VEL | AVGVEL | VORT |  |  |  |  |  |
| Type | 1 | 1 | 1 |  |  |  |  |  |
| Default | 0 | 0 | 0 |  |  |  |  |  |


| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | PRE | PREAVG | LSET | QC | CFL |  |  |  |
| Type | 1 | 1 | 1 | 1 | 1 |  |  |  |
| Default | 0 | 0 | 0 | 0 | 0 |  |  |  |


| Card 3 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | TEMP | TEMPAVG |  |  |  |  |  |  |
| Type | 1 | 1 |  |  |  |  |  |  |
| Default | 0 | 0 |  |  |  |  |  |  |


| Card 4 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | KP | EP | MUT | INT | CMU |  |  |  |
| Type | I | 1 | 1 | 1 | 1 |  |  |  |
| Default | 0 | 0 | 0 | 0 | 0 |  |  |  |

VARIABLE
VEL/AVGVEL/ VORT

Velocity, average velocity, vorticity :
EQ.0: Is output.
EQ.1: Is not output.

PRE/PREAVG/
LSET/QC/CFL

TEMP/
TEMPAVG

KP/EP/MUT /INT/CMU

Pressure, average pressure, levelset, Q criterion, CFL number :
EQ.O: Is output.
EQ.1: Is not output.
Temperature, average temperature :
EQ.O: Is output.
EQ.1: Is not output.
RANS output variables, kinetic energy, diffusion, turbulent viscosity, turbulent intensity, Cmu variable :

EQ.O: Is output.
EQ.1: Is not output.

## *ICFD_CONTROL_PARTITION

Purpose: This keyword changes the default option for the partition in MPP, thus it is only valid in MPP.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | PTECH |  |  |  |  |  |  |  |
| Type | 1 |  |  |  |  |  |  |  |
| Default | 1 |  |  |  |  |  |  |  |

## VARIABLE

## DESCRIPTION

Indicates the type of partition:
EQ.1: the library Metis is used.
EQ.2: partition along the axis with higher aspect ratio
EQ.3: partition along $X$-axis
EQ.4: partition along $Y$-axis
EQ.5: partition along Z-axis

## *ICFD_CONTROL_POROUS

Purpose: This keyword modifies the porous media solve.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | PMSTYPE | VELMETH |  |  |  |  |  |  |
| Type | 1 | 1 |  |  |  |  |  |  |
| Default | 0 | 0 |  |  |  |  |  |  |

VARIABLE
PMSTYPE

VELMETH

## DESCRIPTION

Indicates the porous media solve type.
EQ.0: Anisotropic Generalized Navier-Stokes model for porous media (See *ICFD_MODEL_POROUS) using Fractional step method.
EQ.1: Anisotropic Darcy-Forcheimer model using a Monolithic approach for the solve. This method is better suited for very low Reynolds flows through porous media (frequently encountered in Resin Transfer Molding (RTM) applications). See Remark 1.

Method for determining advection velocity:
EQ.0: Uses FEM approximation for advection velocity
EQ.1: Uses PFEM2 for advection velocity

## Remarks:

1. Anisotropic Darcy-Forcheimer Model. When using the Anisotropic DarcyForcheimer model, the convective term in the Navier Stokes formulation is neglected.
*ICFD_CONTROL_STEADY
Purpose: This keyword allows to specify convergence options for the steady state solver.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | ITS | TOL1 | TOL2 | TOL3 | REL1 | REL2 | UREL | ORDER |
| Type | I | F | F | F | F | F | F | I |
| Default | $1 e 6$ | $1 . e-3$ | $1 . e-3$ | $1 . e-3$ | 0.3 | 0.7 | 1. | 0 |

UREL Under relaxation parameter. Lowering this value may improve the

## VARIABLE

ITS

TOL1/2/3

REL1/2

ORDER

## DESCRIPTION

Maximum number of iterations to reach convergence.
Tolerance limits for the momentum pressure and temperature equations respectfully.

Relaxation parameters for the velocity and pressure respectfully. Decreasing those values may add stability but more iterations may be needed to reach convergence. final accuracy of the solution but more iterations may be needed to achieve convergence.

Analysis order :
EQ.0: Second order. More accurate but more time consuming.
EQ.1: First order: More stable and faster but may be less accurate.

## *ICFD_CONTROL_SURFMESH

Purpose: This keyword enables automatic surface re-meshing. The objective of the remeshing is to improve the mesh quality on the boundaries. It should not be used on a regular basis.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | RSRF | SADAPT |  |  |  |  |  |  |
| Type | 1 | 1 |  |  |  |  |  |  |
| Default | 0 | 0 |  |  |  |  |  |  |

VARIABLE
RSRF

## DESCRIPTION

Indicates whether or not to perform a surface re-meshing.
EQ.0: no re-meshing is applied.
EQ.1: Laplacian smoothing surface remeshing
EQ.2: Curvature preserving surface remeshing
SADAPT Indicates whether or not to trigger adaptive surface remeshing.
EQ.0: no adaptive surface re-meshing is applied.
EQ.1: automatic surface remeshing when quality deteriorates (3D only).

## *ICFD_CONTROL_TAVERAGE

Purpose: This keyword controls the restarting time for computing the time average values. By default, there is no restarting and the average quantities are given starting from $t=0$. This keyword can be useful in turbulent problems that admit a steady state.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | DT |  |  |  |  |  |  |  |
| Type | F |  |  |  |  |  |  |  |
| Default | none |  |  |  |  |  |  |  |

## VARIABLE

DT

## DESCRIPTION

Over each DT time interval, the average quantities are reset.

## *ICFD_CONTROL_TIME

Purpose: Change the default values related to time parameters for the fluid problem.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | TTM | DT | CFL | LCIDSF | DTMIN | DTMAX | DTINIT | TDEATH |
| Type | F | F | F | I | F | F | F | F |
| Default | $10^{28}$ | 0.0 | 1.0 | 0 | $10^{-9}$ | $10^{28}$ | $\downarrow$ | $10^{28}$ |

Optional card

| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | DTT |  |  |  |  |  |  |  |
| Type | F |  |  |  |  |  |  |  |
| Default | Rem 1 |  |  |  |  |  |  |  |

Optional card

| Card 3 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | BTBL |  |  |  |  |  |  |  |
| Type | 1 |  |  |  |  |  |  |  |
| Default | 0 |  |  |  |  |  |  |  |

## VARIABLE

TTM
DT

## DESCRIPTION

Total time of simulation for the fluid problem
Time step for the fluid problem. If nonzero, the time step will be constant and equal to this value. If set to 0.0 , then the time step is automatically computed based on the CFL condition.

## VARIABLE

CFL

LCIDSF

DTMIN

DTMAX

DTINIT

TDEATH

DTT
DTBL Flag to include boundary layer elements in the automatic time step calculation.

EQ.0: The boundary layer elements are excluded (default).
EQ.1: The boundary layer elements are included.

## Remarks:

1. Thermal Time Step. By default, the heat equation is solved using the same time step as that for the velocity/pressure system. This option allows you to assign a specific time step for the thermal solve. It can be useful in cases where the time scales are very different between the two domains. When defined, we recommend DTT to always be greater than or equal to the regular CFD time step.

## *ICFD_CONTROL_TRANSIENT

Purpose: This keyword allows to specify different integration scheme options for the transient solver.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | TORD | FSORD |  |  |  |  |  |  |
| Type | 1 | 1 |  |  |  |  |  |  |
| Default | 0 | 0 |  |  |  |  |  |  |

## VARIABLE

TORD

FSORD Fractional step integration order :
EQ.0: Second order.
EQ.1: First order.

## *ICFD_CONTROL_TURBULENCE

Purpose: Modify the default values for the turbulence model.

## Card Summary:

Card 1. This card is required.

| TMOD | SUBMOD | WLAW | KS | CS |  | TWLAW | TYPLUS |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |

Card 2a. This card is read if $\mathrm{TMOD}=1$. It is optional.

| CE1 | CE2 | SIGMAEPS | SIGMAK | CMU | CCUT |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |

Card $2 b$. This card is read if $\mathrm{TMOD}=2$ or 3 . It is optional.

| Cs |  |  |  |  |  |  |  |
| :---: | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

Card 2c.1. This card is read if $\mathrm{TMOD}=4$. It is optional.

| GAMMA | BETA01 | SIGMAW1 | SIGMAK1 | BETA0ST | CCUT |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |

Card 2c.2. This card is read if $\mathrm{TMOD}=4$. It is optional.

| A1 | BETA02 | SIGMAW2 | SIGMAK2 | CL |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |

Card 2d. This card is read if $\mathrm{TMOD}=5$. It is optional.

| CB1 | CB2 | SIGMANU | CNU1 | CW1 | CW2 |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

## Data Card Definitions:

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | TMOD | SUBMOD | WLAW | KS | CS |  | TWLAW | TYPLUS |
| Type | I | I | 1 | F | F |  | 1 | F |
| Default | 0 | 1 | 1 | 0. | 0. |  | none | none |

## VARIABLE

## DESCRIPTION

TMOD

> Indicates what turbulence model will be used.

SUBMOD Turbulence sub-model.

KS

CS
TWLAW

For RANS $k-\varepsilon$ approach (TMOD $=1$ ):
EQ.1: Standard model
EQ.2: Realizable model
For LES Smagorinsky or dynamic sub-grid model (TMOD = 2):
EQ.1: Smagorinsky model (see Remark 6)
EQ.2: Dynamic model (see Remark 7).
For RANS $k-\omega$ approach (TMOD $=4$ ):
EQ.1: Standard Wilcox 98 model.
EQ.2: Standard Wilcox 06 model.
EQ.3: SST Menter 2003.
WLAW Law of the wall ID if a RANS turbulence model is selected (see Remark 4):

EQ.1: Standard classic law of the wall (default for TMOD = 1)
EQ.2: Standard Launder and Spalding law of the wall
EQ.4: Nonequilibrium Launder and Spalding law of the wall
EQ.5: Automatic classic law of the wall

## DESCRIPTION

EQ.O: Turbulence model based on a variational multiscale approach is used by default.
EQ.1: RANS $k-\varepsilon$ approach (see Remark 1)
EQ.2: LES Smagorinsky or dynamic sub-grid scale model
EQ.3: LES Wall adapting local eddy-viscosity (WALE) model
EQ.4: RANS $k-w$ approach (see Remark 2)
EQ.5: RANS Spalart-Allmaras approach

Roughness physical height, only used for RANS turbulence models.

Roughness constant, only used for RANS turbulence models.
Thermal law of the wall flag (see Remark 8):
EQ.O: No thermal law of the wall activated.

## VARIABLE

## DESCRIPTION

EQ.1: Thermal law of the wall
TYPLUS

Thermal $Y+$ value $\left(Y_{+t}\right)$. If $Y_{+t}$ is not defined, the solver will automatically estimate its value using $Y_{+t c}=Y_{+c} / \operatorname{Pr}^{1 . / 3}$ with $Y_{+c}=$ 11.225 the critical $Y_{+}$value and $\operatorname{Pr}$ the Prandtl number.

RANS $k-\varepsilon$ Card. Optional card if TMOD $=$ 1. Optional card read if $\mathrm{TMOD}=1$. See Remark 1.

| Card 2a | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | CE1 | CE2 | SIGMAE | SIGMAK | CMU | CCUT |  |  |
| Type | F | F | F | F | F | F |  |  |
| Default | 1.44 | 1.92 | 1.3 | 1.0 | 0.09 | -1. |  |  |


| VARIABLE | DES |
| :---: | :---: |
| CEPS1 | $k-\varepsilon$ model constant, $C_{1 \varepsilon}$ |
| CEPS2 | $k-\varepsilon$ model constant, $C_{2 \varepsilon}$ |
| SIGMAEPS | $k-\varepsilon$ model constant, $\sigma_{\varepsilon}$ |
| SIGMAK | $k-\varepsilon$ model constant, $\sigma_{k}$ |
| CMU | $k-\varepsilon$ model constant, $C_{\mu}$ |
| CCUT | $k-\varepsilon$ model constant, $C_{\text {cut }}$ |

LES Card. Optional card read if TMOD $=2$ or 3 .

| Card 2b | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | Cs |  |  |  |  |  |  |  |
| Type | F |  |  |  |  |  |  |  |
| Default | 0.18 |  |  |  |  |  |  |  |

## VARIABLE

Cs

## DESCRIPTION

Smagorinsky constant if TMOD $=2$ and $\operatorname{SUBMOD}=1$ or WALE constant if TMOD $=3$

RANS $k-\omega$ Card. Optional card read if TMOD $=4$. See Remark 2 .

| Card 2c.1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | GAMMA | BETA01 | SIGMAW1 | SIGMAK1 | BETAOST | CCUT |  |  |
| Type | F | F | F | F | F | F |  |  |
| Default | 1.44 | 0.072 | 2. | 2. | 0.09 | -1. |  |  |

RANS $k-\omega$ Card. Optional card read if TMOD $=4$. See Remark 2.

| Card 2c.2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | A1 | BETA02 | SIGMAW2 | SIGMAK2 | CL |  |  |  |
| Type | F | F | F | F | F |  |  |  |
| Default | 0.31 | 0.0828 | 2 | 2 | 0.875 |  |  |  |


| VARIABLE |  | DESCRIPTION |
| :---: | :--- | :--- |
| GAMMA | $k-\omega$ model constant, $\gamma$ |  |
| BETA01 | $k-\omega$ model constant, $\beta_{01}$ |  |
| SIGMAW1 | $k-\omega$ model constant, $\sigma_{\omega 1}$ |  |
| SIGMAK1 | $k-\omega$ model constant, $\sigma_{k 1}$ |  |
| BETA0ST | $k-\omega$ model constant, $\beta_{0}^{*}$ |  |
| CCUT | $k-\omega$ model constant, $C_{\text {cut }}$ |  |
| A1 | $k-\omega$ model constant, $a_{1}$ |  |
| BETA02 | $k-\omega$ model constant, $\beta_{02}$ |  |
| SIGMAW2 | $k-\omega$ model constant, $\sigma_{\omega 2}$ |  |


| VARIABLE |  |
| :---: | :--- |
| SIGMAK2 |  |
| CL | $k-\omega$ model constant, $\sigma_{k 2}$ |
|  |  |

RANS Spalart-Allmaras Card. Optional card read if TMOD $=5$.

| Card 2d | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | CB1 | CB2 | SIGMANU | CNU1 | CW1 | CW2 |  |  |
| Type | F | F | F | F | F | F |  |  |
| Default | 0.1355 | 0.622 | 0.66 | 7.2 | 0.3 | 2.0 |  |  |

## VARIABLE

## DESCRIPTION

CB1
Spalart-Allmaras constant, $C_{b 1}$
CB2 Spalart-Allmaras constant, $C_{b 2}$
SIGMANU
CNU1

CW1
CW2
Spalart-Allmaras constant, $\sigma_{v}$
Spalart-Allmaras constant, $C_{\nu 1}$
Spalart-Allmaras constant, $C_{w 1}$
Spalart-Allmaras constant, $C_{w 2}$

## Remarks:

1. $\quad k-\varepsilon$ Model (TMOD = 1). For the Standard $k-\varepsilon$ model, the following two equations are solved for the turbulent kinetic energy $(k)$ and the turbulent dissipation ( $\varepsilon$ ):

$$
\begin{aligned}
& \frac{\partial k}{\partial t}+\frac{\partial\left(k u_{i}\right)}{\partial x_{i}}=\frac{\partial}{\partial x_{j}}\left[\left(\frac{\mu}{\rho}+\frac{\mu_{t}}{\rho \sigma_{k}}\right) \frac{\partial k}{\partial x_{j}}\right]+P_{k}+P_{b}-\varepsilon+S_{k} \\
& \frac{\partial \varepsilon}{\partial t}+\frac{\partial\left(\varepsilon u_{i}\right)}{\partial x_{i}}=\frac{\partial}{\partial x_{j}}\left[\left(\frac{\mu}{\rho}+\frac{\mu_{t}}{\rho \sigma_{\varepsilon}}\right) \frac{\partial \varepsilon}{\partial x_{j}}\right]+C_{1 \varepsilon} \frac{\varepsilon}{k} P_{k}-C_{2 \varepsilon} \frac{\varepsilon^{2}}{k}+S_{e}
\end{aligned}
$$

Here $P_{k}$ is the $k$ production term (see Remark 3), $P_{b}$ is the production term due to buoyancy and $S_{k}$ and $S_{e}$ are the user defined source terms. $P_{k}$ and $P_{b}$ are expressed as:

$$
\begin{aligned}
P_{k} & =\frac{\mu_{t}}{\rho} S^{2} \\
P_{b} & =\frac{\beta \mu_{t}}{\rho \operatorname{Pr}_{t}} g_{i} \frac{\partial T}{\partial x_{i}}
\end{aligned}
$$

where $S$ is the modulus of the mean rate of strain tensor $\left(S^{2}=2 S_{i j} S_{i j}\right), \beta$ is the coefficient of thermal expansion, and $\operatorname{Pr}_{t}$ is the turbulent Prandtl number. The turbulent viscosity is then expressed as:

$$
\mu_{t}=\rho C_{\mu} \frac{k^{2}}{\varepsilon}
$$

For the realizable $k-\varepsilon$ model, the equation for the turbulent kinetic energy does not change, but the equation for the turbulent dissipation is now expressed as:

$$
\frac{\partial \varepsilon}{\partial t}+\frac{\partial\left(\varepsilon u_{i}\right)}{\partial x_{i}}=\frac{\partial}{\partial x_{j}}\left[\left(\frac{\mu}{\rho}+\frac{\mu_{t}}{\rho \sigma_{\varepsilon}}\right) \frac{\partial \varepsilon}{\partial x_{j}}\right]+C_{1} S \varepsilon-C_{2 \varepsilon} \frac{\varepsilon^{2}}{k+\sqrt{\frac{\mu}{\rho} \varepsilon}}-\varepsilon+S_{e}
$$

Here $C_{1}=\max \left[0.43, \frac{\eta}{\eta+5}\right]$ with $\eta=S \frac{k}{\varepsilon}$.
Furthermore, while the turbulent viscosity is still expressed the same way, $C_{\mu}$ is no longer a constant:

$$
C_{\mu}=\frac{1}{A_{0}+A_{s} k \frac{U^{*}}{\epsilon}}
$$

In the above,

$$
\begin{aligned}
U^{*} & =\sqrt{\Omega_{i j} \Omega_{i j}+S_{i j} S_{i j}} \\
A_{0} & =4.04 \\
A_{s} & =\sqrt{6} \cos \left(\frac{1}{3} \cos ^{-1}\left(\sqrt{6} \frac{S_{i j} S_{j k} S_{k i}}{\left(S_{i j} S_{i j}\right)^{3 / 2}}\right)\right)
\end{aligned}
$$

Note that in this case, the constant value $C_{\mu}$ that can be input by you serves as the limiting value that $C_{\mu}$ can take. By default, $C_{\mu}=0.09$ so:

$$
0.0009<C_{\mu}<0.09
$$

2. $k-\omega$ Model (TMOD = 4). For the Standard Wilcox $06 k-\omega$ model, the following two equations are solved for the turbulent kinetic energy and the specific turbulent dissipation rate, respectively $k$ and $\omega$ :

$$
\begin{aligned}
\frac{\partial k}{\partial t}+\frac{\partial\left(k u_{i}\right)}{\partial x_{i}} & =\frac{\partial}{\partial x_{j}}\left[\left(\frac{\mu}{\rho}+\frac{\mu_{t}}{\rho \sigma_{k 1}}\right) \frac{\partial k}{\partial x_{j}}\right]+P_{k}-\beta^{*} k \omega+S_{k} \\
\frac{\partial \omega}{\partial t}+\frac{\partial\left(\omega u_{i}\right)}{\partial x_{i}} & =\frac{\partial}{\partial x_{j}}\left[\left(\frac{\mu}{\rho}+\frac{\mu_{t}}{\rho \sigma_{\omega 1}}\right) \frac{\partial \varepsilon}{\partial x_{j}}\right]+\gamma \frac{\omega}{k} P_{k}-\beta \omega^{2}+\sigma_{d} X_{k} \omega^{2}+S_{\omega}
\end{aligned}
$$

Here $P_{k}$ is the $k$ production term (see Remark 3) and $S_{k}$ and $S_{\omega}$ are the user defined source terms. $P_{k}, \beta^{*}, \beta$ and $\sigma_{d}$ are expressed as:

$$
\begin{aligned}
P_{k} & =\frac{\mu_{t}}{\rho} S^{2} \\
\beta^{*} & =\beta_{0}^{*} f_{\beta *} \\
\beta & =\beta_{01} f_{\beta} \\
\sigma_{d} & = \begin{cases}0 . & X_{k} \leq 0 . \\
1 / 8 & X_{k}>0 .\end{cases}
\end{aligned}
$$

where

$$
\begin{aligned}
f_{\beta} & =\frac{1+85 X_{\omega}}{1+100 X_{\omega}} \\
f_{\beta *} & =1 . \\
X_{k} & =\frac{1}{\omega} \frac{\partial k}{3} \frac{\partial \omega}{\partial x_{j}} \frac{\partial \omega}{\partial x_{j}} \\
X_{\omega} & =\left|\frac{\Omega_{i j} \Omega_{j k} S_{k i}}{\left(\beta_{0}^{*} \omega\right)^{3}}\right|
\end{aligned}
$$

The turbulent viscosity is then:

$$
\mu_{t}=\rho \frac{k}{\max \left[\omega, C_{l} \sqrt{\frac{2 S_{i j} S_{i j}}{\beta_{0}^{*}}}\right]}
$$

For the Standard Wilcox 98 model, the following terms are modified:

$$
\begin{aligned}
f_{\beta} & =\frac{1+70 X_{\omega}}{1+80 X_{\omega}} \\
f_{\beta *} & =\left\{\begin{array}{cl}
1 & \text { if } X_{k} \leq 0 . \\
1+400 X_{k}^{2}
\end{array}\right. \\
\sigma_{d} & \text { if } X_{k}>0 .
\end{aligned}
$$

The turbulent viscosity is then:

$$
\mu_{t}=\rho \frac{k}{\omega}
$$

For the Menter SST 2003 model, the following equations are solved:

$$
\begin{aligned}
& \frac{\partial k}{\partial t}+\frac{\partial\left(k u_{i}\right)}{\partial x_{i}}=\frac{\partial}{\partial x_{j}}\left[\left(\frac{\mu}{\rho}+\frac{\mu_{t}}{\rho \sigma_{k}}\right) \frac{\partial k}{\partial x_{j}}\right]+P_{k}-\beta_{0}^{*} k \omega+S_{k} \\
& \frac{\partial \omega}{\partial t}+\frac{\partial\left(\omega u_{i}\right)}{\partial x_{i}}=\frac{\partial}{\partial x_{j}}\left[\left(\frac{\mu}{\rho}+\frac{\mu_{t}}{\rho \sigma_{\omega}}\right) \frac{\partial \varepsilon}{\partial x_{j}}\right]+\frac{\gamma}{\mu_{t}} P_{k}-\beta \omega^{2}+2\left(1-F_{1}\right) \sigma_{\omega 2} X_{k} \omega^{2}+S_{\omega}
\end{aligned}
$$

Each of the constants, $\gamma, \beta, \sigma_{k}$, and $\sigma_{\omega}$ are now computed by a blend of two constants with a blending function through:

$$
\alpha=\alpha_{1} F_{1}+\alpha_{2}\left(1-F_{1}\right)
$$

The blending function $F_{1}$ is defined by:

$$
F_{1}=\tanh \left\langle\left[\min \left(\max \left(\frac{\sqrt{k}}{\beta_{0}^{*} \omega y}, \frac{500 v}{y^{2} \omega}\right), \frac{4 \rho \sigma_{\omega 2} k}{\mathrm{CD} \times y^{2}}\right)\right]^{4}\right\rangle
$$

With $y$ the distance to the nearest wall and:

$$
\mathrm{CD}=\max \left(2 \rho \sigma_{\omega 2} X_{k} \omega^{2}, 10^{-10}\right)
$$

The turbulent viscosity is then:

$$
\mu_{t}=\rho \frac{a_{1} k}{\max \left(a_{1} \omega, S F_{2}\right)}
$$

with:

$$
F_{2}=\tanh \left[\left(\max \left(\frac{2 \sqrt{k}}{\beta_{0}^{*} \omega y}, \frac{500 v}{y^{2} \omega}\right)\right)^{2}\right]
$$

3. Production Term. You can activate a limiter on the production term, $P_{k}$, for TMOD $=1$ and 4 . If $C_{\text {cut }} \geq 0$ (CCUT), then:

$$
P_{k}= \begin{cases}\min \left(P_{k}, C_{\mathrm{cut}} \varepsilon\right) & \text { if TMOD }=1 \\ \min \left(P_{k}, C_{\mathrm{cut}} \beta_{0}^{*} k \omega\right) & \text { if TMOD }=4\end{cases}
$$

This is especially common when using the Menter SST 2003 model.
4. Laws of the Wall for RANS Models. For RANS models, the following laws of the wall are available:
a) Standard Classic $(W L A W=1)$.

$$
\begin{aligned}
& U^{+}= \begin{cases}\frac{1}{\kappa} \ln \left(E Y^{+}\right) & \text {if } Y^{+}>11.225 \\
Y^{+} & \text {otherwise }\end{cases} \\
& Y^{+}=\frac{\rho y U_{\tau}}{\mu} \\
& U^{+}=\frac{U}{U_{\tau}} \\
& U_{\tau}=\sqrt{\frac{\tau_{w}}{\rho}}
\end{aligned}
$$

This is the default for $\mathrm{TMOD}=1$.
b) Standard Launder and Spalding $(W L A W=2)$.

$$
U^{*}= \begin{cases}\frac{1}{\mathcal{K}} \ln \left(E Y^{*}\right) & \text { if } Y^{*}>11.225 \\ Y^{*} & \text { otherwise }\end{cases}
$$

$$
\begin{aligned}
& Y^{*}=\frac{\rho C_{\mu}{ }^{1 / 4} k^{1 / 2} y}{\mu} \\
& U^{*}=\frac{U C_{\mu}{ }^{1 / 4} k^{1 / 2}}{U_{\tau}{ }^{2}} \\
& U_{\tau}=\sqrt{\frac{\tau_{w}}{\rho}}
\end{aligned}
$$

c) Nonequilibrium Launder and Spalding $(W L A W=3)$. The nonequilibrium laws of the wall modify the expression of the velocity at the wall making it sensitive to the pressure gradient :

$$
\widetilde{u}=U-\frac{1}{2} \frac{d p}{d x}\left[\frac{y_{v}}{\rho \kappa \sqrt{k}} \ln \left(\frac{y}{y_{v}}\right)+\frac{y-y_{v}}{\rho \kappa \sqrt{k}}+\frac{y_{v}{ }^{2}}{\mu}\right]
$$

with:

$$
y_{v}=\frac{11.225}{y^{*}} y
$$

This law is recommended with TMOD $=1$ and in cases of complex flows involving separation, reattachment and recirculation.
d) Automatic Classic ( $\mathrm{WLAW}=4$ ). The automatic wall law attempts to blend the viscous and $\log$ layers to better account for the transition zone. In the buffer region, we have :

$$
\begin{aligned}
& U^{+}=\frac{U}{U_{\tau}} \\
& U_{\tau}=\sqrt[4]{\left(\frac{U}{y^{+}}\right)^{4}+\left(\frac{U}{\frac{1}{\bar{\kappa}} \ln \left(E y^{+}\right)}\right)^{4}}
\end{aligned}
$$

This is the recommended approach for TMOD $=4$.
5. RANS Turbulence Model with Roughness Included. When a RANS turbulence model is selected, it is possible to define extra parameters to account for roughness effects. In such cases, an extra term will be added to the logarithmic part of the different laws of the wall:

$$
U^{+}=\frac{1}{\kappa} \ln \left(E Y^{+}\right)-\Delta B
$$

If we introduce the non-dimensional roughness height:

$$
K^{+}=\frac{\rho K_{s} C_{\mu}^{1 / 4} k^{1 / 2}}{\mu},
$$

we have:

$$
\Delta B= \begin{cases}0 & \text { for } K^{+} \leq 2.25 \\ \frac{1}{\kappa} \ln \left(\frac{K^{ \pm 2.25}}{87.75}+C_{s} K^{+}\right) \times \sin \left(0.4258\left(\ln K^{+}-0.811\right)\right) & \text { for } 2.25<K^{+} \leq 90.0 \\ \frac{1}{\kappa} \ln \left(1+C_{s} K^{+}\right) & \text {for } 90 .<K^{+}\end{cases}
$$

6. LES Smagorinksy. The LES Smagorinsky turbulence model uses the Van Driest damping function close to the wall:

$$
f_{v}=1-e^{-\frac{y^{+}}{A^{+}}}
$$

7. LES Dynamic Model. The LES dynamic model is based on the model originally proposed by Germano et. al. (1991) and improved by Lilly (1992), with localization on Cs by Piomelli and Liu (1995).
8. Thermal Law of the Wall. When the thermal law of the wall is activated, the turbulent heat flux will be calculated as an additional output variable:

$$
\begin{aligned}
Q_{t} & =\rho C_{p} \frac{U_{\tau}}{T_{+}}\left(T_{s}-T_{c}\right) \\
T_{+} & = \begin{cases}\operatorname{Pr}_{t} Y_{+} & \text {if } Y_{+t} \leq Y_{+t c} \\
\frac{\operatorname{Pr}_{t}}{\vartheta} \log \left(Y_{+}\right)+\left(3.85 \operatorname{Pr}_{t}^{1 . \overline{3}}-1.3\right)^{2}+2.12 \log \left(\operatorname{Pr}_{t}\right) & \text { otherwise }\end{cases}
\end{aligned}
$$

Purpose: This keyword enables the user impose a divergence-free turbulent field on inlets.

Card must be used jointly with VAD $=4$ of keyword *ICFD_BOUNDARY_PRESCRIBED_VEL.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | PID | IU | IV | IW | LS |  |  |  |
| Type | I | F | F | F | F |  |  |  |
| Default | 0 | $10^{-3}$ | $10^{-3}$ | $10^{-3}$ | $h_{\text {min }}$ |  |  |  |

VARIABLE
PID
IU, IV, IW
Intensity of field fluctuations over $x, y$, and $z$ directions,

$$
\mathrm{IU}=\frac{u^{\prime}}{u_{\mathrm{avg}}} .
$$

LS Integral length scale of turbulence

## Remarks:

1. If this card is not defined but a turbulent field inlet has been activated. See VAD $=4$ of *ICFD_BOUNDARY_PRESCRIBED_VEL, the default parameters will be used.

## *ICFD_DATABASE_AVERAGE

Purpose: This keyword enables the computation of time average variables at given time intervals.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | DT |  |  |  |  |  |  |  |
| Type | F |  |  |  |  |  |  |  |
| Default | none |  |  |  |  |  |  |  |

## VARIABLE

DT

## DESCRIPTION

Over each DT time interval, an average of the different fluid variables will be calculated and then reset when moving to the next DT interval.

## Remarks:

1. The file name for this database is icfdavg. ${ }^{*}$. dat with the different averaged variable values copied in a ASCII format.

## *ICFD_DATABASE_DRAG_\{OPTION\}

Available options include
VOL
Purpose: This keyword enables the computation of drag forces over given surface parts of the model. If multiple keywords are given, the forces over the PID surfaces are given in separate files and are also added and output in a separate file.

For the VOL option, drag calculation can also be applied on a volume defined by ICFD_PART_VOL. This is mostly useful in porous media applications to output the pressure drag of the porous media domain.

Surface Drag Cards. Include one card for each surface on which drag is applied. This input ends at the next keyword ("*") card.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | PID | CPID | DTOUT | PEROUT | DIVI | ELOUT | SSOUT |  |
| Type | I | I | F | 1 | 1 | 1 | 1 |  |
| Default | none | none | 0. | 0 | 10 | 0 | 0 |  |

## VARIABLE

PID

DIVI

ELOUT
SSOUT

CPID Center point ID used for the calculation of the force's moment. By default the reference frame center is used is $\mathbf{0}=(0,0,0)$.

DTOUT Time interval to print the output. If DTOUT is equal to 0.0 , then the ICFD timestep will be used.

PEROUT Outputs the contribution of the different elements on the total drag Outputs the contribution of the different ele
in fractions of the total drag in the d3plots.

## DESCRIPTION

Part ID of the surface where the drag force will be computed.

Number of drag divisions for PEROUT. Default is 10 which means the contributions will be grouped in 10 deciles.

Outputs the drag value of each element in the d3plots.
Outputs the pressure loads caused by the fluid on each solid segment set in keyword format. FSI needs to be activated.

## Remarks:

1. The file name for this database is icfdragi for instantaneous drag and icfdraga for the drag computed using average values of pressure and velocities.
2. The output contains:
a) "Fpx", "Fpy", and "Fpz" refer to the three components of the pressure drag force

$$
\mathbf{F}_{p}=\int P d A
$$

where $P$ is the pressure and $A$ the surface area.
b) "Fvx", "Fxy", and "Fvz" refer to the three components of the viscous drag force

$$
\mathbf{F}_{v}=\int \mu \frac{\partial \mathrm{u}}{\partial \mathrm{y}} \mathrm{~d} A
$$

where $\frac{\partial \mathrm{u}}{\partial \mathrm{y}}$ is the shear velocity at the wall, $\mu$ is the viscosity and $A$ is the surface area.
c) "Mpx", "Mpy", "Mpz", "Mvx", "Mvy", and "Mvz" refer to the three components of the pressure and viscous force moments respectively.

## *ICFD_DATABASE_FLUX

Purpose: This keyword enables the computation of the flow rate and average pressure over given parts of the model. If multiple keywords are given, separate files are output.

Include as many cards as needed. This input ends at the next keyword ("*") card.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | PID | DTOUT |  |  |  |  |  |  |
| Type | I | F |  |  |  |  |  |  |
| Default | none | $\downarrow$ |  |  |  |  |  |  |

VARIABLE
PID
DTOUT Output frequency. Default is at every fluid timestep.

## Remarks:

1. Database Name. The file name for this database is icfd_flux.dat.
2. Database Components. The flux database contains the flow rate through a section, called "output flux,"

$$
\Phi=\sum_{i}\left(\mathbf{V}_{i} \cdot \mathbf{n}_{i}\right) A_{i}
$$

the average pressure, called "Pre-avg,"

$$
P_{\mathrm{avg}}=\frac{\sum_{i} P_{i} A_{i}}{\sum_{i} A_{i}}
$$

and the total area, called "Areatot."

## *ICFD_DATABASE_FLUX_SURF

Purpose: Enable the computation of the flow rate and average pressure over a given surface (icfd part) of the model which is not necessary part of the analysis but lies in the fluid volume by mapping and interpolating volume results on the given surface.

Include as many cards as needed. This input ends at the next keyword ("*") card.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | PID | DTOUT |  |  |  |  |  |  |
| Type | I | F |  |  |  |  |  |  |
| Default | none | $\downarrow$ |  |  |  |  |  |  |

## VARIABLE

PID
DTOUT

## DESCRIPTION

Part ID of the surface where the flow rates will be computed Output frequency. Default is at every fluid timestep.

## Remarks:

1. Database Name. The file name for this database is icfd_fluxsurf.dat.

## *ICFD_DATABASE_FORCE_DEM

Purpose: Enable the computation of the total fluid force that is transferred to the DEM particles over time. The output is in icfd_force_dem.dat.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | OUT |  |  |  |  |  |  |  |
| Type | 1 |  |  |  |  |  |  |  |
| Default | 0 |  |  |  |  |  |  |  |
| VARIABLE |  |  |  |  |  |  |  |  |

OUT
Flag to enable computing the fluid forces and generating output:
EQ.0: No output is generated.
EQ.1: Output is generated.

## *ICFD_DATABASE_HTC

Purpose: This keyword allows the user to trigger the calculation of the Heat transfer coefficient using different methods and to control the output options.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | OUT | HTC | TB |  |  |  |  | OUTDT |
| Type | I | I | F |  |  |  |  | F |
| Default | 0 | 0. | 0. |  |  |  |  | 0. |

## VARIABLE

OUT

HTC Determines how the HTC is calculated.
EQ.O: Automatically calculated by the solver based on the average temperature flowing through the pipe section (See Remark 1).

EQ.1: User imposed value (See Remark 2).
Value of the bulk temperature if $\mathrm{HTC}=1$.
OUTDT Output frequency of the HTC in the various ASCII files. If left to 0 ., the solver will output the HTC at every timestep.

## Remarks:

1. The heat transfer coefficient is frequently used in thermal applications to estimate the effect of the fluid cooling and it derived from a CFD calculation.
2. The heat transfer coefficient is defined as follows:

$$
h=\frac{q}{T_{s}-T_{b}}
$$

with $q$ the heat flux, $T_{s}$ the surface temperature and $T_{b}$ the so called "bulk" temperature. For external aerodynamic applications, this bulk temperature is often defined as a constant (ambient or far field conditions, HTC =1). However, for internal aerodynamic application, this temperature is often defined as an average temperature flowing through the pipe section with the flow velocity being used as a weighting factor $(\mathrm{HTC}=0)$.

## *ICFD_DATABASE_NODEAVG

Purpose: This keyword enables the computation of the average quantities on surface nodes defined in *ICFD_DATABASE_NODOUT.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | 0N |  |  |  |  |  |  |  |
| Type | 1 |  |  |  |  |  |  |  |
| Default | 0 |  |  |  |  |  |  |  |

VARIABLE
ON

## DESCRIPTION

If equal to 1 , the average quantities will be computed.

## Remarks:

1. The file name for this database is icfd_nodeavg.dat.

## *ICFD_DATABASE_NODOUT

Purpose: This keyword enables the output of ICFD data on surface nodes. For data in the fluid volume, it is advised to use points or tracers (See *ICFD_DATABASE_POINTOUT).

Output Options Card.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | OUTLV | DTOUT |  |  |  |  |  |  |
| Type | I | F |  |  |  |  |  |  |
| Default | 0 | 0. |  |  |  |  |  |  |

Include as many cards as needed. This input ends at the next keyword ("*") card.

| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | NID1 | NID2 | NID3 | NID4 | NID5 | NID6 | NID7 | NID8 |
| Type | 1 | 1 | 1 | 1 | 1 | 1 | 1 | 1 |
| Default | none | none | none | none | none | none | none | none |

VARIABLE
OUTLV

## DESCRIPTION

Determines if the output file should be dumped.
EQ.O: No output file is generated.
EQ.1: The output file is generated.
DTOUT Time interval to print the output. If DTOUT is equal to 0.0 , then the ICFD timestep will be used.

NID.. Node IDs.

## Remarks:

1. The file name for this database is icfd_nodout.dat.

## *ICFD_DATABASE_NTEMPOUT

Purpose: Output the temperature at individual nodes in a format consistent with *ICFD_INITIAL_TEMPNODE to initialize a subsequent ICFD problem.

Include as many cards as needed. This input ends at the next keyword ("*") card.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | NID | DTOUT |  |  |  |  |  |  |
| Type | I | F |  |  |  |  |  |  |
| Default | none | 0.0 |  |  |  |  |  |  |

VARIABLE
NID Internal ICFD node ID
DTOUT Output frequency
EQ.0.0: The ICFD timestep will be used.

## *ICFD_DATABASE_POINTAVG

Purpose: This keyword enables the computation of the average quantities on point sets using the parameters defined in *ICFD_DATABASE_POINTOUT.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | 0N |  |  |  |  |  |  |  |
| Type | I |  |  |  |  |  |  |  |
| Default | 0 |  |  |  |  |  |  |  |

VARIABLE
ON

## DESCRIPTION

If equal to 1 , the average quantities will be computed.

## Remarks:

1. The file name for this database is icfd_psavg.dat.

## *ICFD_DATABASE_POINTOUT

Purpose: This keyword enables the output of ICFD data on points.
Output Options Card.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | PSID | DTOUT | PSTYPE | VX | VY | VZ |  |  |
| Type | I | F | I | F | F | F |  |  |
| Default | 0 | 0. | 0 | 0. | 0. | 0. |  |  |

Include as many cards as needed. This input ends at the next keyword ("*") card.

| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | PID | X | Y | Z |  |  |  |  |
| Type | 1 | F | F | F |  |  |  |  |
| Default | none | none | none | none |  |  |  |  |
| VARIABLE |  | DESCRIPTION |  |  |  |  |  |  |
| PSID |  | Point Set ID. |  |  |  |  |  |  |
| DTOUT |  | Time interval to print the output. If DTOUT is equal to 0.0 , then the ICFD timestep will be used. |  |  |  |  |  |  |
| PSTYPE |  | Point Set type |  |  |  |  |  |  |
|  |  | EQ.O: Fixed points. |  |  |  |  |  |  |
|  |  | EQ.1: Tracer points using prescribed velocity. |  |  |  |  |  |  |
|  |  | EQ.2: Tracer points using fluid velocity. |  |  |  |  |  |  |
|  |  | EQ.3: Tracer points using mesh velocity.. |  |  |  |  |  |  |
| VX, VY, VZ |  | Constant velocities to be used when PSTYPE $=1$ |  |  |  |  |  |  |
| PID |  | Point ID |  |  |  |  |  |  |

## VARIABLE

## DESCRIPTION

$\mathrm{X}, \mathrm{Y}, \mathrm{Z} \quad$ Point initial coordinates

## Remarks:

1. The file name for this database is icfd_pointout.dat.

## *ICFD_DATABASE_RESIDUALS

Purpose: This keyword allows the user to output the residuals of the various systems.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | RLVL |  |  |  |  |  |  |  |
| Type | 1 |  |  |  |  |  |  |  |
| Default | 0 |  |  |  |  |  |  |  |

## VARIABLE

## DESCRIPTION

RLVL
Residual output level :
EQ.0: No output.
EQ.1: Only outputs the number of iterations needed for solving the pressure Poisson equation.
EQ.2: Outputs the number of iterations for the momentum, pressure, mesh movement and temperature equations.
EQ.3: Also gives the residual for each iteration during the solve of the momentum, pressure, mesh movement and temperature equations.

## Remarks:

1. The file names for the momentum, pressure, mesh movement and temperature equations are called icfd_residuals.moms.dat, icfd_residuals.pres.dat, icfd_residuals.mmov.dat, and icfd_residuals.temp.dat respectively.

## *ICFD_DATABASE_SSOUT

Purpose: Output the pressure load on a structure from the fluid. It can be useful for linear FSI applications, where the structure is made static, and the loads applied by the fluid are retrieved.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | OUT | OUTDT | LCIDSF |  |  |  |  | POFF |
| Type | 1 | 1 | 1 |  |  |  |  | F |
| Default | 0 | 0. | 0 |  |  |  |  | 0. |

## VARIABLE

OUT

OUTDT

LCIDSF Optional load curve ID to apply a scale factor on the fluid pressure output

POFF Optional pressure offset on the fluid pressure output

## *ICFD_DATABASE_SSOUT_EXCLUDE

Purpose: This keyword defines which segment set IDs are excluded from the SSOUT search. No forces coming from the fluid will be transmitted on those segment sets for output (see ICFD_DATABASE_SSOUT).

Include as many cards as needed. This input ends at the next keyword ("*") card.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | SSOUTID |  |  |  |  |  |  |  |
| Type | 1 |  |  |  |  |  |  |  |
| Default | none |  |  |  |  |  |  |  |

VARIABLE
SSOUTID

## DESCRIPTION

Segment Set ID of the solid mechanics problem which is to be excluded from the output of the fluid forces on the solid boundaries.

## *ICFD_DATABASE_TEMP

Purpose: This keyword enables the computation of the average temperature and the heat flux over given parts of the model. If multiple keywords are given, separate files are output.

Include as many cards as needed. This input ends at the next keyword ("*") card.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | PID | DTOUT |  |  |  |  |  |  |
| Type | I | F |  |  |  |  |  |  |
| Default | none | none |  |  |  |  |  |  |

## VARIABLE

PID

## DESCRIPTION

Part ID of the surface where the average temperature and heat flux will be computed.

DTOUT Output frequency. Default is at every fluid timestep.

## Remarks:

1. The file name for this database is icfd_thermal.dat.
2. Two average temperature are given in the icfd_thermal.dat file: "Temp-avg" and "Temp-sum". The average temperature is calculated using the local node area as weighting factor,

$$
T_{\mathrm{avg}}=\frac{\sum_{i}^{N} T_{i} A_{i}}{\sum_{i}^{N} A_{i}}
$$

whereas, the sum is not weighted by area

$$
T_{\text {sum }}=\frac{\sum_{i}^{N} T_{i}}{N}
$$

If the mesh is regular, the two values will be of similar value. The icfd_thermal.dat output file also includes the average heat flux, the total surface area, and the average heat transfer coefficients (See *ICFD_DATABASE_HTC).
*ICFD_DATABASE_TIMESTEP
Purpose: This keyword enables the output of ICFD data regarding the ICFD timestep.
Output Options Card.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | OUTLV |  |  |  |  |  |  |  |
| Type | 1 |  |  |  |  |  |  |  |
| Default | 0 |  |  |  |  |  |  |  |

## VARIABLE

DESCRIPTION
OUTLV Determines if the output file should be dumped.
EQ.O: No output file is generated.
EQ.1: The output file is generated.

## Remarks:

1. The file name for this database is icfd_tsout.dat.
2. Outputs the run's ICFD timestep versus the timestep calculated using the ICFD CFL condition as criteria (autotimestep). This can be useful in cases using a fixed timestep where big mesh deformations and/or big fluid velocity changes occur in order to track how that fixed timestep value compares to the reference autotimestep.

## *ICFD_DATABASE_UINDEX

Purpose: This keyword allows the user to have the solver calculate the uniformity index (See Remark 1).

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | OUT |  |  |  |  |  |  |  |
| Type | 1 |  |  |  |  |  |  |  |
| Default | 0 |  |  |  |  |  |  |  |

VARIABLE
OUT

## DESCRIPTION

Determines if the solver should calculate the uniformity index.
EQ.0: Off.
EQ.1: On.

## Remarks:

1. Uniformity Index. The uniformity index is a post treatment quantity which measures how uniform the flow is through a given section. It is especially useful in internal aerodynamics cases. It is expressed as :

$$
\gamma=1-\frac{1}{2 n A} \sum_{i=1}^{n}\left[\frac{\sqrt{\left(u_{i}-\bar{u}\right)^{2}}}{\bar{u}} A_{i}\right]
$$

with $A_{i}$, the local cell area, $A$ the total section area, $u_{i}$ the local velocity, $\bar{u}$ the average velocity through the section, and $n$ the number of cells.

Values close to 0 means that the flow is very unevenly distributed. This can be used to identify bends, corners or turbulent effects. Values close to 1 imply smooth or equally distributed flow through the surface.

## *ICFD_DATABASE_WETNESS

Purpose: For free surface problems, output the wetted area as a function of time for a given surface part. It is output to an ASCII file beginning with icfd_wetness.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | SPID | DTOUT |  |  |  |  |  |  |
| Type | I | F |  |  |  |  |  |  |
| Default | none | 0. |  |  |  |  |  |  |

VARIABLE
SPID
DTOUT

## DESCRIPTION

Part ID of the surface where the wetted area will be computed
Time interval to print the output. If DTOUT is equal to 0.0 , then the ICFD time step will be used.

## Remarks:

1. Wetted Area. The ICFD solver identifies a surface as wet if its level set value is positive.

## *ICFD_DEFINE_HEATSOURCE

Purpose: This keyword defines a volumetric heat source for the heat equation solve.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | HSID | LCID | SHAPE | R | PTID1 | PTID2 |  |  |
| Type | I | I | I | F | I | I |  |  |
| Default | none | none | none | none | none | none |  |  |

VARIABLE
HSID
LCID

SHAPE

R

PTID1

PTID2

DESCRIPTION
Heat source ID.
Load curve ID specifying the evolution of the heat source term function of time for the $\mathrm{X}, \mathrm{Y}$ and Z dofs, see *DEFINE_CURVE,*DEFINE_CURVE_FUNCTION or ${ }^{*}$ DEFINE_FUNCTION. If a DEFINE_FUNCTION is used, the following parameters are allowed: $f(x, y, z, v x, v y, v z$, temp, pres, time $)$.

Shape of the volumetric heat source:
EQ.1: Box shape
EQ. 2 : Cylinder shape
EQ. 3 : Sphere shape
Radius of the sphere is SHAPE $=3$
ID of point (See ICFD_DEFINE_POINT) of minimum coordinates if $\mathrm{SHAPE}=1$, tail point if $\mathrm{SHAPE}=2$, origin if SHAPE $=3$.

ID of point of maximum coordinates if $\mathrm{SHAPE}=2$, head point if $\mathrm{SHAPE}=2$.

## *ICFD_DEFINE_RESIDENCETIMESOURCE

Purpose: Specify a volumetric residence time (RT) source.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | RTSID | ISHAPE | R | PTID1 | PTID2 | MASSDIFF | DEATHT | IRTOPBC |
| Type | I | I | F | I | i | F | F | I |
| Default | none | none | none | none | none | $10^{-6}$ | $\downarrow$ | 0 |

## VARIABLE

RTSID
ISHAPE

R Radius of the cylinder if ISHAPE $=2$ or radius of the sphere if

PTID1

PTID2

MASSDIFF Mass diffusion for the transport equation
DEATHT

IRT0PBC

## DESCRIPTION

RT source ID
Shape of the volumetric RT source:
EQ.1: Box
EQ.2: Cylinder
EQ.3: Sphere ISHAPE $=3$ if ISHAPE $=2$, or the origin of the sphere if ISHAPE $=3$ ISHAPE $=1$ or the head point of the cylinder if ISHAPE $=2$.

End time for the source.
EQ.O.O: End time of the simulation
Flag for which prescribed boundaries $\mathrm{RT}=0$ is imposed:

ID of a point (see *ICFD_DEFINE_POINT) giving the minimum coordinate of the box if ISHAPE $=1$, the tail point for the cylinder

ID of a point giving the maximum coordinate of the box if

EQ.O: Imposed only on boundaries with prescribed velocity
EQ.1: Imposed on boundaries with either prescribed velocity or prescribed pressure
*ICFD_DEFINE_SOURCE
Purpose: Define a volumetric external force for the momentum equation solve.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | SID | LCIDX | LCIDY | LCIDZ | SHAPE | R | PTID1 | PTID2 |
| Type | 1 | 1 | 1 | 1 | 1 | F | 1 | 1 |
| Default | none | none | none | none | none | none | none | none |

VARIABLE
SID
LCIDX/Y/Z

SHAPE

R
PTID1 Point ID (see *ICFD_DEFINE_POINT) whose meaning depends on SHAPE:

SHAPE.EQ.1: Minimum coordinates of the box
SHAPE.EQ.2: Tail point of the cylinder
SHAPE.EQ.3: Origin of the sphere
PTID2 Point ID (see *ICFD_DEFINE_POINT) whose meaning depends on SHAPE:

SHAPE.EQ.1: Maximum coordinates of the box
SHAPE.EQ.2: Head point of the cylinder

## *ICFD_DEFINE_SPTRANSPSOURCE

Purpose: Specify a volumetric species source for the species transport solver.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | SPTRSID | LCID | ISHAPE | R | PTID1 | PTID2 | MASSDIFF | DEATHT |
| Type | I | I | I | F | I | i | F | F |
| Default | none | none | none | none | none | none | $10^{-6}$ | $\downarrow$ |

## VARIABLE

SPTRSID
LCID

ISHAPE

R

PTID1 ID of a point (see *ICFD_DEFINE_POINT) giving the minimum coordinate of the box if ISHAPE $=1$, the tail point for the cylinder if ISHAPE $=2$, or the origin of the sphere if ISHAPE $=3$

PTID2 ID of a point giving the maximum coordinate of the box if ISHAPE $=1$ or the head point of the cylinder if ISHAPE $=2$.

MASSDIFF
DEATHT End time for the source.
EQ.O.O: End time of the simulation

## *ICFD_DEFINE_TURBSOURCE

Purpose: This keyword defines a external source for the RANS turbulent equations.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | TSID | LCIDK | LCIDEP | LCIDNU | SHAPE | R | PTID1 | PTID2 |
| Type | I | I | I | I | I | F | I | 1 |
| Default | none | none | none | none | none | none | none | None |

VARIABLE
TSID
LCIDK

LCIDEP

LCIDNU

DESCRIPTION
Turbulent external source ID.
Load curve ID specifying the evolution of the external source term function of time for the turbulent kinetic energy $k$ equation, see ${ }^{*}$ DEFINE_CURVE,*DEFINE_CURVE_FUNCTION or *DEFINE_FUNCTION. If a DEFINE_FUNCTION is used, the following parameters are allowed: $f(x, y, z, v x, v y, v z$, temp, pres, time $)$.

Load curve ID specifying the evolution of the external source term function of time for the turbulent diffusion $\varepsilon$ or specific rate of dissipation $w$ equation, see *DEFINE_CURVE,*DEFINE_CURVE_FUNCTION or *DEFINE_FUNCTION. If a DEFINE_FUNCTION is used, the following parameters are allowed: $f(x, y, z, v x, v y, v z$, temp, pres, time $)$.

Load curve ID specifying the evolution of the external source term function of time for the kinematic eddy turbulent viscosity equation used in the Spalart-Allmaras model, see *DEFINE_CURVE,*DEFINE_CURVE_FUNCTION or *DEFINE_FUNCTION. If a DEFINE_FUNCTION is used, the following parameters are allowed: $f(x, y, z, v x, v y, v z$, temp, pres, time $)$.

VARIABLE
SHAPE

R
PTID1

PTID2

## DESCRIPTION

Shape of the external source:
EQ. 1 : Box shape
EQ. 2 : Cylinder shape
EQ. 3 : Sphere shape
Radius of the sphere is SHAPE $=3$
ID of point (See ICFD_DEFINE_POINT) of minimum coordinates if $\mathrm{SHAPE}=1$, tail point if $\mathrm{SHAPE}=2$, origin if $\mathrm{SHAPE}=3$.

ID of point of maximum coordinates if $\mathrm{SHAPE}=2$, head point if $\mathrm{SHAPE}=2$.

## *ICFD_DEFINE_POINT

Purpose: This keyword defines a point in space that could be used for multiple purposes.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | POID | X | Y | Z | CONSTPID |  |  |  |
| Type | I | F | F | F | I |  |  |  |
| Default | none | none | none | none | none |  |  |  |

Optional Card 2. Load curve IDS specifying velocity components of translating point

| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | LCIDX | LCIDY | LCIDZ |  |  |  |  |  |
| Type | 1 | 1 | 1 |  |  |  |  |  |
| Default | 0 | 0 | 0 |  |  |  |  |  |

Optional Card 3. Load curve IDS and rotation axis of rotating point

| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | LCIDW | XT | YT | ZT | XH | YH | ZH |  |
| Type | I | F | F | F | F | F | F |  |
| Default | 0 | none | none | none | none | none | none |  |

VARIABLE
POID
$X / Y / Z$

## DESCRIPTION

Point ID.
$x, y, z$ coordinates for the point.

VARIABLE
CONSTPID

LCIDX/LCIDY /LCIDZ

LCIDW

XT/YT/ZT
XH/YH/ZH

DESCRIPTION
Surface Part ID to which the point is constrained. This means that if the selected surface moves, then the localization of the point will update as well.

The point can be made to translate. Those are the three load curve IDs for the three translation components.

The point can also be made to rotate. This load curve specifies the angular velocity.

Rotation axis tail point coordinates.
Rotation axis head point coordinates.

## *ICFD_DEFINE_NONINERTIAL

Purpose: Define a non-inertial reference frame to avoid heavy mesh distortions and remeshing associated with large-scale rotations. This frame helps when modeling spinning cylinders, wind turbines, and turbomachinery.

Include as many cards as needed. This input ends at the next keyword ("*") card.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | W1 | W2 | W3 | R | PTID | L | LCID | RELV |
| Type | F | F | F | F | I | F | I | I |
| Default | none | none | none | none | none | none | none | 0 |

## VARIABLE

W1, W2, W3
R

## DESCRIPTION

Rotational velocity, $\boldsymbol{\omega}$, along the $X, Y$, and $Z$-axes
Radius of the rotating reference frame.
GT.0.0: Constant value
LT.0.0: $|\mathrm{R}|$ refers to a *DEFINE_FUNCTION ID. The function supports the following arguments: $f(x, y, z, v x, v y, v z$, temp, pres, time).


Figure 7-4. Non Inertial Reference Frame Example

## VARIABLE

PTID

L Length of the rotating reference frame
LCID Load curve for scaling factor of $\omega$.
GT.0: Load curve ID (see *DEFINE_CURVE) for the curve giving the scale factor as a function of time

LT.O: |LCID| refers to a *DEFINE_FUNCTION ID. The function supports the following arguments: $f(x, y, z, v x, v y, v z$, temp, pres, time).

RELV Velocities computed and displayed:
EQ.O: Relative velocity, only the non-rotating components of the velocity are used and displayed.
EQ.1: Absolute velocity. All the components of the velocity are used. Useful in cases where several or at least one noninertial reference frame is combined with an inertial "classical" reference frame.

## *ICFD_DEFINE_WAVE_DAMPING

Purpose: This keyword defines a damping zone for free surface waves.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | PID | NID | L | F1 | F2 | N | LCID |  |
| Type | I | I | F | F | F | I | I |  |
| Default | none | none |  | 10 | 10 | 1 | none |  |

VARIABLE
PID
NID

L

F1/F2
N
LCID

DESCRIPTION
Point ID defining the start of the damping layer.
Normal ID defined using ICFD_DEFINE_POINT and pointing to the outgoing direction of the damping layer.

Length of damping layer. If no is value specified, the damping layer will have a length corresponding to five element lengths.

Linear and quadratic damping factor terms.
Damping term factor.
Load curve ID acting as temporal scale factor on damping term.

## Remarks:

1. The damping is achieved by adding a source term to the momentum equations :

$$
s^{d}=w\left(f_{1}+f_{2}|u|\right) u
$$

with $w$ the weight function :

$$
w=\frac{e^{\gamma}-1}{e-1}
$$

and $\gamma$ the blending function which allows a smooth insertion of the source term in the damping layer :

$$
\gamma=\left(\frac{x-x_{s d}}{x_{e d}-x_{s d}}\right)^{n}
$$

$x_{s d}$ and $x_{e d}$ representing the start and end coordinates of the damping zone.
*ICFD_INITIAL
Purpose: Simple initialization of velocity and temperature within a volume.
Include as many cards as needed. This input ends at the next keyword ("*") card.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | PID | Vx | Vy | Vz | T | P |  | DFUNC |
| Type | I | F | F | F | F | F |  | 1 |
| Default | none | none | none | none | none | none |  | 0 |

## VARIABLE

## DESCRIPTION

Part ID for the volume elements or the surface elements where the values are initialized (see *ICFD_PART_VOL and *ICFD_PART). PID $=0$ to assign the initial condition to all nodes at once.

Vx $\quad \mathrm{x}$ coordinate for the velocity.
Vy y coordinate for the velocity.
$\mathrm{Vz} \quad \mathrm{z}$ coordinate for the velocity.
T Initial temperature.
P Initial Pressure.
DFUNC Option to define initial conditions using *DEFINE_FUNCTION
EQ.O: Turned off.
EQ.1: Turned on. All previous flags for initial velocity, pressure and temperature now refer to *DEFINE_FUNCTION IDs. The following parameters are allowed : $f(x, y, z)$, allowing to define initial profiles function of coordinates.

## *ICFD_INITIAL_LEVELSET

Purpose: Define an initial level set surface instead of a multi-fluid domain (replaces the need for *MESH_INTERF).

Include as many cards as needed. This input ends at the next keyword ("*") card.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | STYPE | NX | NY | NZ | X | Y | Z | INVERT |
| Type | I | F | F | F | F | F | F | I |
| Default | none | none | none | none | none | none | none | 0 |

## VARIABLE

STYPE

NX, NY, NZ $\quad X, Y$ and $Z$ components of the section plane normal if STYPE $=1$. Minimum coordinates of the box, $P_{\text {min }}$, if STYPE $=2$. NX is the sphere/cylinder radius if STYPE $=3$ and STYPE $=4$. NY is the cylinder length if STYPE $=4$. NZ is the global axis if STYPE $=4$ ( $\mathrm{NZ}=1$, 2, or 3 means $X$-axis, $Y$-axis, or $Z$-axis, respectively).
$\mathrm{X}, \mathrm{Y}, \mathrm{Z} \quad X, Y$ and Z components of the section plane origin point if STYPE $=1$ and 4 . Maximum coordinates of the box, $P_{\max }$ if STYPE $=2$. Coordinates of the sphere origin point if $S T Y P E=3$.

INVERT Inversion of initial level set:
EQ.0: No inversion. Positive level set values are assigned to nodes contained within the volume defined by STYPE.
EQ.1: The sign of the initial level set values is reversed.

## Remarks:

1. Box Adjacent to Fluid Boundaries. When STYPE $=2$ is used and the box is adjacent to the fluid boundaries such as during a dam break simulation, the distance from any point in the fluid to the fluid boundary must remain smaller than the distance to the defined box. Therefore, the $P_{\min }$ coordinates need to be defined far outside the initial fluid domain.
2. Multiple Keyword Definitions. Multiple definitions of this keyword are possible, but the different shapes generated must not intersect for a correct initialization.


Figure 7-5. 2D Dam breaking example with initial levelset surface defined using STYPE $=2 . P_{\min }$ is defined sufficiently far away from the fluid surface boundaries.

## *ICFD_INITIAL_SPTRANSP

Purpose: Initialize the concentration of the species being transported within a volume.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | PID | CONC | IDFUNC |  |  |  |  |  |
| Type | I | F | 1 |  |  |  |  |  |
| Default | 0 | none | 0 |  |  |  |  |  |

CONC Initial concentration. CONC is *DEFINE_FUNCTION ID if ID-

## VARIABLE

PID

IDFUNC

## DESCRIPTION

Part ID for the volume elements or surface elements where the values are initialized.

EQ.O: Assign the initial condition to all nodes at once. FUNC = 1 .

Flag to define initial conditions using a *DEFINE_FUNCTION:
EQ.0: Turned off.
EQ.1: Turned on. CONC is a *DEFINE_FUNCTION ID. The following parameters are allowed: $f(x, y, z)$. This allows for defining initial profiles as a function of coordinates.
*ICFD_INITIAL_TEMPNODE
Purpose: Allow the solver to initialize the temperature at individual nodes.
Include as many cards as needed. This input ends at the next keyword ("*") card.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | NID | TEMP |  |  |  |  |  |  |
| Type | I | F |  |  |  |  |  |  |
| Default | none | none |  |  |  |  |  |  |

VARIABLE
NID
TEMP Initial temperature value

## *ICFD_INITIAL_TURBULENCE

Purpose: Modify the default initial values of the turbulence quantities for a RANS turbulence model.

Include as many cards as needed. This input ends at the next keyword ("*") card.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | PID | I | R | K | EW |  |  |  |
| Type | I | F | F | F | F |  |  |  |
| Default | none | none | none | optional | optional |  |  |  |

## VARIABLE

PID

I
$\mathrm{R} \quad$ Initial turbulence viscosity to laminar viscosity ratio $\left(r=\mu_{\mathrm{turb}} / \mu\right)$.
K Initial kinetic energy. When defined, it replaces the choice of I.
LT.0.0: $|\mathrm{K}|$ refers to a *DEFINE_FUNCTION ID. The following parameters are allowed: $f(x, y, z)$, allowing you to define initial profiles as a function of coordinates.

EW Initial turbulence specific dissipation rate or dissipation rate depending on the choice of turbulence model. When defined, it replaces the choice of $R$.

LT.O.O: |EW| refers to a *DEFINE_FUNCTION ID. The following parameters are allowed: $f(x, y, z)$, allowing you to define initial profiles as a function of coordinates.

## Remarks:

1. Default Initial Conditions. If no initial conditions have been assigned to a specific PID, the solver will automatically pick $\mathrm{I}=0.05$ (5\%) and $\mathrm{R}=10000$.
*ICFD_MAT_\{OPTION\}
Available options include

## TITLE

Purpose: Specify physical properties for the fluid material.

## Fluid Material Card Sets:

The Material Fluid Parameters Card is required. If a second card is given, it must be a Thermal Fluid Parameters Card. If the fluid thermal properties are not needed, the second card can be a blank card. With the third card, you can associate the fluid material to a non-Newtonian model, to a porous media model, and/or to a species transport model (see *ICFD_MODEL_NONNEWT, *ICFD_MODEL_POROUS, and *ICFD_MODEL_SPECIES_TRANSPORT).

Material Fluid Parameters Card.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | MID | FLG | R0 | VIS | ST | STSFLCID | CA |  |
| Type | I | I | F | F | F | I | F |  |
| Default | none | 1 | 0. | 0. | 0. | none | 0. |  |

Thermal Fluid Parameters Card. Only to be defined if the thermal problem is solved. Otherwise include a blank card.

| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | HC | TC | BETA | PRT | HCSFLCID | TCSFLCID |  |  |
| Type | F | F | F | F | I | I |  |  |
| Default | 0. | 0. | 0. | 0.85 | none | none |  |  |

Additional fluid models. Only to be defined if the fluid is non-Newtonian, there is a porous media, and/or a species is being transported.

| Card 3 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | NNMOID | PMMOID |  | SPTRID |  |  |  |  |
| Type | 1 | 1 |  | 1 |  |  |  |  |
| Default | optional | optional |  | optional |  |  |  |  |

## VARIABLE

MID
FLG

RO Flow density
VIS Dynamic viscosity
ST Surface tension coefficient
STSFLCID Load curve ID for scale factor applied on ST as a function of time. See *DEFINE_CURVE, *DEFINE_CURVE_FUNCTION, or *DEFINE_FUNCTION. If a *DEFINE_FUNCTION is used, the following parameters are allowed: $f(x, y, z, v x, v y, v z$, temp, pres, time).

CA Contact angle
HC Heat capacity
TC Thermal conductivity
BETA Thermal expansion coefficient used in the Boussinesq approximation for buoyancy

PRT Turbulent Prandlt number. Only used if K-Epsilon turbulence model selected.

HCSFLCID

TCSFLCID

NNMOID

PMMOID Porous media model ID. This refers to a porous media model defined using *ICFD_MODEL_POROUS.

SPTRID Species transport model ID. This refers to a species transport model defined using *ICFD_MODEL_SPECIES_TRANSPORT.

## *ICFD_MODEL_NONNEWT

Purpose: Specify a non-newtonian model or a viscosity law that can associated to a fluid material.

Non-Newtonian Model ID and type.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | NNMOID | NNID |  |  |  |  |  |  |
| Type | 1 | 1 |  |  |  |  |  |  |
| Default | none | none |  |  |  |  |  |  |

Non-Newtonian Fluid Parameters Card.

| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | K | N | MUMIN | LAMBDA | ALPHA | TALPHA |  |  |
| Type | F | F | F | F | F | F |  |  |
| Default | 0.0 | 0.0 | 0.0 | $1 . \mathrm{e} 30$ | 0.0 | 0.0 |  |  |

## VARIABLE

NNMOID Non-Newtonian Model ID.
NNID Non-Newtonian fluid model type :
EQ. 1 : Power-Law model
EQ. 2 : Carreau model
EQ. 3 : Cross model
EQ. 4 : Herschel-Bulkley model
EQ. 5 : Cross II model
EQ. 6 : Sutherland formula for temperature dependent viscosity
EQ. 7 : Power-Law for temperature dependent viscosity
EQ.8 : Viscosity defined by Load Curve ID or Function ID

| VARIABLE | DESCRIPTION |
| :---: | :---: |
| K | Consistency index if NNID $=1$ and 4. Zero shear Viscosity if NNID $=2,3$ and 5.Reference viscosity if NNID $=6$ and NNID $=7$. Load curve ID or function ID if NNID $=8$. |
| N | Measure of the deviation of the fluid from Newtonian (Power Law index) for NNID $=1,2,3,4,5,7$. Not used for NNID $=6$ and 8 . |
| MUMIN | Minimum acceptable viscosity value if NNID $=1$. Infinite Shear Viscosity if NNID $=2,5$. Yielding viscosity if NNID $=4$. Not used if NNID $=3,6,7,8$. |
| LAMBDA | Maximum acceptable viscosity value if NNID $=1$. Time constant if NNID $=2,3,5$. Yield Stress Threshold if NNID $=4$.Sutherland constant if NNID $=6$. Not used if NNID $=7,8$. |
| ALPHA | Activation energy if NNID $=1,2$. Not used if NNID $=3,4,5,6,7,8$. |
| TALPHA | Reference temperature if NNID=2. Not used if NNID $=1,3,4,5,6,7,8$ |

## Remarks:

1. For the Non-Newtonian models, the viscosity is expressed as :
a) POWER-LAW :

$$
\begin{aligned}
& \mu=k \dot{\gamma}^{n-1} e^{\alpha T_{0} / T} \\
& \mu_{\min }<\mu<\mu_{\max }
\end{aligned}
$$

With $k$ the consistency index, $n$ the power law index, $\alpha$ the activation energy, $T_{0}$ the initial temperature, $T$ the temperature at any given time $t, \mu_{\text {min }}$ the minimum acceptable viscosity and $\mu_{\max }$ the maximum acceptable viscosity.
b) CARREAU :

$$
\begin{gathered}
\mu=\mu_{\infty}+\left(\mu_{0}-\mu_{\infty}\right)\left[1+(H(T) \dot{\gamma} \lambda)^{2}\right]^{(n-1) / 2} \\
H(T)=\exp \left[\alpha\left(\frac{1}{T-T_{0}}-\frac{1}{T_{\alpha}-T_{0}}\right)\right]
\end{gathered}
$$

With $\mu_{\infty}$ the infinite shear viscosity, $\mu_{0}$ the zero shear viscosity, $n$ the power law index, $\lambda$ a time constant, $\alpha$ the activation energy, $T_{0}$ the initial temperature, $T$
the temperature at any given time $t$ and $T_{\alpha}$ the reference temperature at which $H(T)=1$.
c) CROSS :

$$
\mu=\frac{\mu_{0}}{1+(\lambda \dot{\gamma})^{1-n}}
$$

With $\mu_{0}$ the zero shear viscosity, $n$ the power law index and $\lambda$ a time constant.
d) HERSCHEL-BULKLEY :

$$
\begin{gathered}
\mu=\mu_{0} \text { if }\left(\dot{\gamma}<\tau_{0} / \mu_{0}\right) \\
\mu=\frac{\tau_{0}+k\left[\dot{\gamma}^{n}-\left(\tau_{0} / \mu_{0}\right)^{n}\right]}{\dot{\gamma}}
\end{gathered}
$$

With $k$ the consistency index, $\tau_{0}$ the Yield stress threshold, $\mu_{0}$ the yielding viscosity and $n$ the power law index.
e) CROSS II :

$$
\mu=\mu_{\infty}+\frac{\mu_{0}-\mu_{\infty}}{1+(\lambda \dot{\gamma})^{n}}
$$

With $\mu_{0}$ the zero shear viscosity, $\mu_{\infty}$ the infinite shear viscosity, $n$ the power law index and $\lambda$ a time constant.
2. For the temperature dependent viscosity models, the viscosity is expressed as :
a) SUTHERLAND's LAW :

$$
\mu=\mu_{0}\left(\frac{T}{T_{0}}\right)^{3 / 2} \frac{T_{0}+S}{T+S}
$$

With $\mu_{0}$ a reference viscosity, $T_{0}$ the initial temperature (which therefore must not be 0 .), $T$ the temperature at any given time $t$ and $S$ Sutherland's constant.
b) POWER LAW :

$$
\mu=\mu_{0}\left(\frac{T}{T_{0}}\right)^{n}
$$

With $\mu_{0}$ a reference viscosity, $T_{0}$ the initial temperature (which therefore must not be 0 .), $T$ the temperature at any given time $t$ and $n$ the power law index.
3. For NNID $=8$, a load curve function of time, a curve function or a function can be used. If it references a DEFINE_FUNCTION, the following arguments are allowed $f(x, y, z, v x, v y, v z$, temp , pres, shear, time $)$.

## *ICFD_MODEL_POROUS

Purpose: Specify a porous media model.

## Card Summary:

Card 1. This card is required.

| PMMOID | PMMTYPE | FORM | RHOCP | KAPPA |  |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

Card 2a. This card is included if PMMTYPE $=1,2$, or 8.

| POR | PER | FF |  | PSFLCID |  |  |  |
| :---: | :---: | :---: | :--- | :--- | :--- | :--- | :--- |

Card 2 b. This card is included if PMMTYPE $=3$ or 10 .

| POR | TH |  | FABTH | PVLCID |  |  |  |
| :---: | :---: | :--- | :--- | :--- | :--- | :--- | :--- |

Card 2c. This card is included if PMMTYPE $=4,6$, or 7 .

| POR |  |  |  |  |  |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

Card 2d. This card is included if PMMTYPE $=5$.

| POR | THX | THY | THZ | PVLCIDX | PVLCIDY | PVLCIDZ |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |

Card 2e. This card is included if $\mathrm{PMMTYPE}=11$.

| POR | ALPHA | BETA |  |  |  |  |  |
| :---: | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

Card 3. This card is included if PMMTYPE $=4,5,6$, or 7

| KXP | KYP | KZP |  |  |  |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

Card 4a. This card is included if PMMTYPE $=4$ or 6 .

| PROJXPX | PROJXPY | PROJXPZ | PROJYPX | PROJYPY | PROJYPZ |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

Card 4b. This card is included if PMMTYPE $=5$ or 7 .

| PID1REF | PID2REF |  |  |  |  |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

## Data Card Definitions:

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | PMMOID | PMMTYPE | FORM | RHOCP | KAPPA |  |  |  |
| Type | I | I | I | F | F |  |  |  |
| Default | none | none | 0 | 0.0 | 0.0 |  |  |  |

VARIABLE
PMMOID Porous media model ID
PMMTYPE Porous media model type:
EQ.1: Isotropic porous media - Ergun correlation
EQ.2: Isotropic porous media - Darcy-Forchheimer model
EQ.3: Isotropic porous media - permeability defined through pressure-velocity data
EQ.4: Anisotropic porous media. Fixed local reference frame (see Figure 7-6).

EQ.5: Anisotropic porous media model - moving local reference frame and permeability vector in local reference frame $\left(x^{\prime}, y^{\prime}, z^{\prime}\right)$ defined by three pressure-velocity curves.

EQ.6: Anisotropic porous media model - moving local reference frame and permeability vector constant.

EQ.7: Anisotropic porous media model - moving local reference frame and permeability vector constant. This model differs from PMMTYPE $=6$ in the way the local reference frame is moved.

EQ.8: Main parachute model to be used jointly with *MESH_EMBEDSHELL for the parachute surface. Similar to PMMTYPE $=2$.

EQ.10: Parachute model to be used jointly with *MESH_EMBEDSHELL where the fabric permeability and Forchheimer factor are computed from the pressure-velocity curves of experimental data given by a *LOAD_CURVE. Similar to PMMTYPE $=3$.

## DESCRIPTION

EQ.11: Parachute model similar to $\mathrm{PMMTYPE}=8$, but pressure gradient is directly defined by coefficients $\alpha$ and $\beta$ as:

$$
\frac{\Delta P\left(u_{x}\right)}{\Delta x}=\alpha u_{x}+\beta u_{x}^{2} .
$$

FORM Porous media formulation:
EQ.0: Classical (default)
EQ.2: Interstitial velocity
RHOCP Density of the structure multiplied by the specific heat of the structure

KAPPA Thermal conductivity of the structure
Porous Media Parameters Card (PMMTYPE = 1, 2, and 8). This card is included PMMTYPE =1, 2, or 8 .

| Card 2a | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | POR | PER | FF |  | PSFLCID |  |  |  |
| Type | F | F | F |  | 1 |  |  |  |
| Default | 0. | 0. | 0. |  | optional |  |  |  |

VARIABLE
POR
PER
FF
PSFLCID

DESCRIPTION
Porosity, $\varepsilon$
Permeability, $\kappa$
Forchheimer factor to be defined if PMMTYPE $=2$ or 8.
Optional permeability scale factor load curve ID, *DEFINE_CURVE_FUNCTION ID or *DEFINE_FUNCTION ID. If a *DEFINE_FUNCTION is used, the following parameters are allowed: $f(x, y, z, v x, v y, v z$, temp, pres, time).

Porous Media Parameters Card (PMMTYPE = 3 and 10). This card is included if PMMTYPE $=3$ or 10 .

| Card 2b | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | POR | TH |  | FABTH | PVLCID |  |  |  |
| Type | F | F |  | F | I |  |  |  |
| Default | 0. | 0. |  | 0. | none |  |  |  |

## VARIABLE

POR Porosity, $\varepsilon$

PVLCID

TH $\quad$ Probe thickness if PMMTYPE $=3$
FABTH $\quad$ Fabric thickness if PMMTYPE $=10$
DESCRIPTION

Pressure as a function of velocity load curve ID

Porous Media Parameters Card (PMMTYPE = 4, 6, and 7). This card is included if PMMTYPE $=4,6$, or 7 .

| Card 2c | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | POR |  |  |  |  |  |  |  |
| Type | F |  |  |  |  |  |  |  |
| Default | 0. |  |  |  |  |  |  |  |

VARIABLE
POR

DESCRIPTION
Porosity, $\varepsilon$

Porous Media Parameters Card (PMMTYPE = 5). This card is included if PMMTYPE $=5$.

| Card 2d | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | POR | THX | THY | THZ | PVLCIDX | PVLCIDY | PVLCIDZ |  |
| Type | F | F | F | F | I | I | I |  |
| Default | 0. | 0. | 0. | 0. | none | none | none |  |

## VARIABLE

POR
THX
THY
THZ
PVLCIDX

PVLCIDY

PVLCIDZ

## DESCRIPTION

Porosity, $\varepsilon$
Probe thickness, $\Delta x$
Probe thickness, $\Delta y$
Probe thickness, $\Delta z$
Load curve ID for pressure as a function of velocity in the global X-direction

Load curve ID for pressure as a function of velocity in the global $Y$-direction

Load curve ID for pressure as a function of velocity in the global Z-direction

Porous Media Parameters Card (PMMTYPE = 11). This card is included if PMMTYPE $=11$.

| Card 2e | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | POR | ALPHA | BETA |  |  |  |  |  |
| Type | F | F | F |  |  |  |  |  |
| Default | 0. | 0. | 0. |  |  |  |  |  |

## VARIABLE

POR
ALPHA
BETA

## DESCRIPTION

Porosity, $\varepsilon$
Coefficient, $\alpha$
Coefficient, $\beta$

Permeability Vector Card in local reference frame. Only to be defined if the porous media is anisotropic (PMMTYPE $=4,5,6,7$ ).

| Card 3 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | KXP | KYP | KZP |  |  |  |  |  |
| Type | F | F | F |  |  |  |  |  |
| Default | 0. | 0. | 0. |  |  |  |  |  |

## VARIABLE

KXP, KYP,
KZP

## DESCRIPTION

Permeability vector in local reference frame $\left(x^{\prime}, y^{\prime}, z^{\prime}\right)$. Those values become scale factors if PMMTYPE $=5$.

Projection of Local Vectors in Global Reference Frame. This card is defined if PMMTYPE $=4$ or 6 .

| Card 4a | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | PROJXPX | PROJXPY | PROJXPZ | PROJYPX | PROJYPY | PROJYPZ |  |  |
| Type | F/I | F/I | F/I | F/I | F/I | F/I |  |  |
| Default | $0 . / 0$ | $0 . / 0$ | $0 . / 0$ | $0 . / 0$ | $0 . / 0$ | $0 . / 0$ |  |  |

## VARIABLE

PROJXPX, PROJXPY, PROJXPZ

## DESCRIPTION

Projection of the local permeability vector, $\mathbf{x}^{\prime}$, in the global reference frame, $(x, y, z)$. If PMMTYPE $=6$, PROJXPX, PROJXPY, and PRPJXPZ become load curve IDs, so the coordinates of the local $\mathbf{x}^{\prime}$ vector can change in time.

VARIABLE
PROJYPX, PROJYPY, PROJYPZ

## DESCRIPTION

Projection of the local permeability vector, $\mathbf{y}^{\prime}$, in the global reference frame, $(x, y, z)$. If PMMTYPE $=6$, PROJYPX, PROJYPY, and PRPJYPZ become load curve IDs, so the coordinates of the local $\mathbf{y}^{\prime}$ vector can change in time.

Local Reference Frame Vectors. This card is defined if PMMTYPE $=5$ or 7 .

| Card 4b | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | PID1REF | PID2REF |  |  |  |  |  |  |
| Type | 1 | 1 |  |  |  |  |  |  |
| Default | 0 | 0. |  |  |  |  |  |  |

## VARIABLE

PID1REF,
PID2REF

## DESCRIPTION

Two local reference frame vectors are defined by the coordinates of the two-point IDs defined by PID1REF and PID2REF. (See *ICFD_DEFINE_POINT). Since those points can be made to move, it is therefore possible to define a moving reference frame for the anisotropic porous media domain.

## Remarks:

1. Generalized Flow Equations in a Porous Media. Let $\varepsilon$ be the porosity and $\kappa$ be the permeability of the porous media. Then,

$$
\varepsilon=\frac{\text { void volume }}{\text { total volume }} \text {. }
$$

$u_{i}$, the volume averaged velocity field, can then be defined in terms of the fluid velocity field, $u_{i f}$, as:

$$
u_{i}=\varepsilon u_{i f} .
$$

The generalized flow equations of momentum and mass conservation can be expressed as:

$$
\begin{aligned}
\frac{\partial u_{i}}{\partial x_{i}} & =0 \\
\frac{\rho}{\varepsilon}\left[\frac{\partial u_{i}}{\partial t}+\frac{\partial}{\partial x_{j}}\left(\frac{u_{i} u_{j}}{\varepsilon}\right)\right] & =-\frac{1}{\varepsilon} \frac{\partial(P \varepsilon)}{\partial x_{i}}+\frac{\mu}{\varepsilon}\left(\frac{\partial^{2} u_{i}}{\partial x_{j} \partial x_{j}}\right)+\rho g_{i}-D_{i}
\end{aligned}
$$



Figure 7-6. Anisotropic porous media vectors definition (PMMTYPE $=4,5,6$, and 7). The vectors $\mathbf{X}$ and $\mathbf{Y}$ are the global axes; $\mathbf{x}^{\prime}$ and $\mathbf{y}^{\prime}$ define the system for the primed coordinate $\left(x^{\prime}, y^{\prime}, z^{\prime}\right)$.
where $D_{i}$ are the forces exerted on the fluid by the porous matrix (see Remarks 2 and 3 ).
2. Porous Forces for Isotropic Models. For the isotropic model, the porous forces are a function of the matrix porosity and its permeability. For the isotropic case, three models are available:
a) Model 1 (Ergun correlation).

$$
D_{i}=\frac{\mu u_{i}}{\kappa}+\frac{1.75 \rho|U|}{\sqrt{150} \sqrt{\kappa} \varepsilon^{3 / 2}} u_{i}
$$

b) Model 2 (Darcy-Forcheimer).

$$
D_{i}=\frac{\mu u_{i}}{\kappa}+\frac{F \varepsilon \rho|U|}{\sqrt{\kappa}} u_{i}
$$

c) Model 3. Using the $\Delta P-V$ experimental data. In this case, it is assumed that the pressure-velocity curve was obtained by applying a pressure difference or pressure drop on both ends of a porous slab of thickness $\Delta x$ with porous properties $\kappa$ and $\varepsilon$. It then becomes possible for the solver to fit that experimental curve with a quadratic polynomial of the form $\Delta P\left(u_{x}\right)=\alpha u_{x}{ }^{2}+\beta u_{x}$. Once $\alpha$ and $\beta$ are known, it is possible to estimate $D_{i}$.
3. Anisotropic Darcy-Forcheimer Term. The anisotropic (see Figure 7-6) version of the Darcy-Forcheimer term can be written as:

$$
\begin{aligned}
D_{i} & =\mu B_{i j} u_{j}+F \varepsilon|U| C_{i j} u_{j} \\
B_{i j} & =\left(K_{i j}\right)^{-1} \\
C_{i j} & =\left(K_{i j}\right)^{-1 / 2}
\end{aligned}
$$

Here $K_{i j}$ is the anisotropic permeability tensor.
*ICFD_MODEL_SPECIES_TRANSPORT
Purpose: Specify a species transport model that can be associated with a fluid material.

Species transport model ID and type.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | SPTRID | SPTRTYPE |  |  |  |  |  |  |
| Type | 1 | 1 |  |  |  |  |  |  |
| Default | none | none |  |  |  |  |  |  |

VARIABLE
SPTRID
SPTRTYPE

DESCRIPTION
Species transport model ID
Species transport model type:
EQ.1: Passive species transport
*ICFD_PART_\{OPTION\}
Available options include

## TITLE

Purpose: Define parts for this incompressible flow solver.
The TITLE option allows the user to define an additional optional line with a HEADING in order to associate a name to the part.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | HEADING |  |  |  |  |  |  |  |
| Type | A |  |  |  |  |  |  |  |
| Default | none |  |  |  |  |  |  |  |

Part Material Card. Include as many cards as needed. This input ends at the next keyword ("*") card.

| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | PID | SECID | MID |  |  |  |  |  |
| Type | 1 | 1 | 1 |  |  |  |  |  |
| Default | none | none | none |  |  |  |  |  |

VARIABLE
PID
SECID
MID Material identifier defined with the *ICFD_MAT card.
*ICFD_PART_VOL_\{OPTION\}
Available options include

## TITLE

Purpose: This keyword assigns material properties to the nodes enclosed by surface ICFD parts.

The TITLE option allows the user to define an additional optional line with a HEADING in order to associate a name to the part.

| Title | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |
| Variable | HEADING |  |  |  |  |  |  |  |
| Type | A |  |  |  |  |  |  |  |
| Default | none |  |  |  |  |  |  |  |


| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | PID | SECID | MID |  |  |  |  |  |
| Type | 1 | 1 | 1 |  |  |  |  |  |
| Default | none | none | none |  |  |  |  |  |

Provide as many cards as necessary. This input ends at the next keyword ("**) card

| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | SPID1 | SPID2 | SPID3 | SPID4 | SPID5 | SPID6 | SPID7 | SPID8 |
| Type | 1 | 1 | 1 | 1 | 1 | 1 | 1 | 1 |
| Default | none | none | none | none | none | none | none | none |

## VARIABLE

PID
SECID
MID
SPID1, ...

## DESCRIPTION

Part identifier for fluid volumes.
Section identifier defined by the *ICFD_SECTION card.
Material identifier.

Part IDs for the surface elements that define the volume mesh.


```
$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$
$$$$ *ICFD_PART VOI
$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$
$
$ PART ID 5 is defined by the surfaces that enclose it.
$
*ICFD_PART_VOL
$
```



```
$ pid secid mid
```



Purpose: Define a section for the incompressible flow solver.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | SID |  |  |  |  |  |  |  |
| Type | I |  |  |  |  |  |  |  |
| Default | none |  |  |  |  |  |  |  |

VARIABLE DESCRIPTION

Section identifier.

## *ICFD_SET_NODE_LIST

Purpose: Only used in cases where the mesh is specified by the user (See *MESH_VOLUME_ELEMENT). Defines a set of nodes associated with a part ID on which boundary conditions can be applied.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | SID | PID |  |  |  |  |  |  |
| Type | I | 1 |  |  |  |  |  |  |
| Default | none | none |  |  |  |  |  |  |

Node List Card. Provide as many cards as necessary. This input ends at the next keyword ("*") card

| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | NID1 | NID2 | NID3 | NID4 | NID5 | NID6 | NID7 | NID8 |
| Type | 1 | 1 | 1 | 1 | 1 | 1 | 1 | 1 |
| Default | none | none | none | none | none | none | none | none |

VARIABLE
SID
PID
NID1, ...

## DESCRIPTION

Set ID
Associated Part ID.
Node IDs

## Remarks:

1. The convention is the similar to the one used by the keyword *SET_NODE_LIST and serves a similar purpose.

## *ICFD_SOLVER_SPLIT

Purpose: This keyword provides an option to trigger an iterative procedure on the fluid system. This procedure aims to bring more precision to the final pressure and velocity values but is often very time consuming. It must therefore be used with caution. It is intended only for special cases. For stability purposes, this method is automatically used for the first ICFD time step.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | NIT | TOL |  |  |  |  |  |  |
| Type | I | F |  |  |  |  |  |  |
| Default | 1 | $10^{-3}$ |  |  |  |  |  |  |

VARIABLE
NIT

TOL

DESCRIPTION
Maximum Number of iterations of the system for each fluid time step. If TOL criteria is not reached after NIT iterations, the run will proceed.

Tolerance Criteria for the pressure residual during the fluid system solve.

## *ICFD_SOLVER_TOL_FSI

Purpose: This keyword allows the user to change the default tolerance values for the Newton Raphson loop in the strong FSI analysis. Care should be taken when deviating from the default values.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | ATOL | RTOL |  | MAXIT |  |  |  |  |
| Type | F | F |  | 1 |  |  |  |  |
| Default | $10^{-5}$ | $10^{-5}$ |  | 1000 |  |  |  |  |

## VARIABLE

ATOL

RTOL

MAXIT Maximum number of iterations allowed to achieve convergence. If a negative integer is entered, then that value will be used as a load curve ID for MAXIT.

## *ICFD_SOLVER_TOL_LSET

Purpose: This keyword allows the user to change the default tolerance values for the advection equation for levelset. Care should be taken when deviating from the default values.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | ATOL | RTOL |  | MAXIT |  |  |  |  |
| Type | F | F |  | 1 |  |  |  |  |
| Default | $10^{-8}$ | $10^{-8}$ |  | 1000 |  |  |  |  |

VARIABLE
ATOL

RTOL

MAXIT

## DESCRIPTION

Absolute convergence criteria. Convergence is achieved when Residual $_{i+1}-$ Residual $_{i} \leq$ ATOL. If a negative integer is entered, then that value will be used as a load curve ID for ATOL.

Relative convergence criteria. Convergence is achieved when $\left(\operatorname{Residual}_{i+1}-\operatorname{Residual}_{i}\right) /$ Residual $_{\text {initial }} \leq R T O L$. If a negative integer is entered, then that value will be used as a load curve ID for RTOL.

Maximum number of iterations allowed to achieve convergence. If a negative integer is entered, then that value will be used as a load curve ID for MAXIT.

## *ICFD_SOLVER_TOL_MMOV

Purpose: This keyword allows the user to change the default tolerance values for the mesh movement algorithm. Care should be taken when deviating from the default values.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | ATOL | RTOL |  | MAXIT |  | DISPTOL |  |  |
| Type | F | F |  | 1 |  | $F$ |  |  |
| Default | $1 e-8$ | $1 e-8$ |  | 1000 |  | 0. |  |  |

VARIABLE
ATOL

RTOL Relative convergence criteria. Convergence is achieved when $\left(\right.$ Residual $\left._{i+1}-\operatorname{Residual}_{i}\right) /$ Residual $_{\text {initial }} \leq$ RTOL. If a negative integer is entered, then that value will be used as a load curve ID for RTOL.

MAXIT Maximum number of iterations allowed to achieve convergence. If a negative integer is entered, then that value will be used as a load curve ID for MAXIT.

DISPTOL Element deformation tolerance before a matrix reassembly is triggered. Default is 0 . which means any element deformation detected will automatically trigger a matrix reassembly. Higher values will potentially save calculation times at the expense of accuracy.

## *ICFD_SOLVER_TOL_MOM

Purpose: This keyword allows the user to change the default tolerance values for the momentum equation solve. Care should be taken when deviating from the default values.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | ATOL | RTOL |  | MAXIT |  |  |  |  |
| Type | F | F |  | 1 |  |  |  |  |
| Default | $10^{-8}$ | $10^{-8}$ |  | 1000 |  |  |  |  |

VARIABLE
ATOL

RTOL

MAXIT

## DESCRIPTION

Absolute convergence criteria. Convergence is achieved when Residual $_{i+1}-$ Residual $_{i} \leq$ ATOL. If a negative integer is entered, then that value will be used as a load curve ID for ATOL.

Relative convergence criteria. Convergence is achieved when $\left(\right.$ Residual $\left._{i+1}-\operatorname{Residual}_{i}\right) /$ Residual $_{\text {initial }} \leq$ RTOL. If a negative integer is entered, then that value will be used as a load curve ID for RTOL.

Maximum number of iterations allowed to achieve convergence. If a negative integer is entered, then that value will be used as a load curve ID for MAXIT.

## *ICFD_SOLVER_TOL_MONOLITHIC

Purpose: This keyword allows the user to change the default tolerance values for the monolithic solver. Care should be taken when deviating from the default values.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | ATOL | RTOL |  | MAXIT |  |  |  |  |
| Type | F | F |  | 1 |  |  |  |  |
| Default | $10^{-8}$ | $10^{-8}$ |  | 1000 |  |  |  |  |

VARIABLE
ATOL

RTOL

MAXIT

## DESCRIPTION

Absolute convergence criteria. Convergence is achieved when Residual $_{i+1}-$ Residual $_{i} \leq$ ATOL. If a negative integer is entered, then that value will be used as a load curve ID for ATOL.

Relative convergence criteria. Convergence is achieved when $\left(\right.$ Residual $\left._{i+1}-\operatorname{Residual}_{i}\right) /$ Residual $_{\text {initial }} \leq$ RTOL. If a negative integer is entered, then that value will be used as a load curve ID for RTOL.

Maximum number of iterations allowed to achieve convergence. If a negative integer is entered, then that value will be used as a load curve ID for MAXIT.

## *ICFD_SOLVER_TOL_PRE

Purpose: Change the default tolerance values for the Poisson equation for pressure. Care should be taken when deviating from the default values.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | ATOL | RTOL |  | MAXIT | PREID | PTOL |  |  |
| Type | F | F |  | I | I | F |  |  |
| Default | $10^{-8}$ | $10^{-8}$ |  | 1000 | 2 | $10^{-3}$ |  |  |

VARIABLE
ATOL

RTOL

MAXIT Maximum number of iterations allowed to achieve convergence. If a negative integer is entered, then that value will be used as a load curve ID for MAXIT.

PREID Choice of Preconditioner for the Conjugate Gradient Solve:
EQ.1: Diagonal Preconditioner
EQ.2: Incomplete LU factorization
EQ.5: Global MUMPS factorization

PTOL Preconditioner tolerance (a.k.a Drop Tolerance if PREID = 2)

## *ICFD_SOLVER_TOL_TEMP

Purpose: This keyword allows the user to change the default tolerance values for the heat equation. To be handled with great care.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | ATOL | RTOL |  | MAXIT |  |  |  |  |
| Type | F | F |  | 1 |  |  |  |  |
| Default | $1 e-8$ | $1 e-8$ |  | 1000 |  |  |  |  |

## VARIABLE

ATOL

RTOL

MAXIT

## DESCRIPTION

Absolute convergence criteria. Convergence is achieved when Residual $_{i+1}-$ Residual $_{i} \leq A T O L$. If a negative integer is entered, then that value will be used as a load curve ID for ATOL.

Relative convergence criteria. Convergence is achieved when $\left(\right.$ Residual $_{i+1}-$ Residual $\left._{i}\right) /$ Residual $_{\text {initial }} \leq R T O L$. If a negative integer is entered, then that value will be used as a load curve ID for RTOL.

Maximum number of iterations allowed to achieve convergence. If a negative integer is entered, then that value will be used as a load curve ID for MAXIT.

## *MESH

The keyword *MESH is used to create a mesh that will be used in the analysis. So far only tetrahedral (or triangular in 2-d) elements can be generated. The keyword cards in this section are defined in alphabetical order:

*MESH_BL<br>*MESH_BL_SYM<br>*MESH_EMBEDSHELL<br>*MESH_INTERF<br>*MESH_NODE<br>*MESH_SIZE_<br>*MESH_SIZE_SHAPE<br>*MESH_SURFACE_ELEMENT<br>*MESH_SURFACE_NODE<br>*MESH_VOLUME<br>*MESH_VOLUME_ELEMENT<br>*MESH_VOLUME_NODE<br>*MESH_VOLUME_PART

## *MESH_BL

Purpose: Define a boundary-layer mesh as a refinement on the volume-mesh. The boundary layer mesh is constructed by subdividing elements near the surface.

Boundary Layer Cards. Define as many cards as are necessary. The next keyword ("*") card terminates this input.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | PID | NELTH | BLTH | BLFE | BLST |  |  |  |
| Type | I | I | F | F | I |  |  |  |
| Default | none | none | 0. | 0. | 0 |  |  |  |

## VARIABLE

PID
NELTH

BLTH Boundary layer mesh thickness if BLST = 1 or 2 . Growth scale factor if BLST $=3$. Ignored if $\mathrm{BLST}=0$.

BLFE Distance between the wall and the first volume mesh node if $\mathrm{BLST}=3$. Scaling coefficient if BLST $=1$ or 2 . Ignored if BLST $=0$.

BLST Boundary layer mesh generation strategy:
EQ.0: $2^{\text {NELTH+1 }}$ subdivisions based on surface mesh size (default). See Remark 1.

EQ.1: Power law using BLTH and NELTH with BLFE as a scale factor. See Remark 2 and Figure 8-1.

EQ.2: Geometric series based on BLTH and BLFE. See Remark 3 and Figure 8-2.

EQ.3: Repartition following a growth scale factor (BLTH). See Remark 4 and Figure 8-3.

## Remarks:

1. $\quad \mathbf{B L S T}=\mathbf{0}$. For $\operatorname{BLST}=0$, for every additional NELTH, the automatic volume mesher will divide the elements closest to the surface by two so that the smallest element in the boundary layer mesh will have an aspect ratio of $2^{\mathrm{NELTH}+1}$. A default boundary layer mesh thickness based on the surface mesh size will be chosen.
2. $\mathbf{B L S T}=1$. For a constant repartition of the nodes in the boundary layer, use $\operatorname{BLST}=1$ with $\operatorname{BLFE}=1$. For BLST = 1, starting from the wall, the position of node $n$ in the normal direction is given by:

$$
X_{n}=\left(\frac{n}{\mathrm{NELTH}+1}\right)^{[5 \times(1-\mathrm{BLFE})]} \frac{\mathrm{BLTH}}{\sum_{i=1}^{\mathrm{NELTH}+1}\left[i /(\mathrm{NELTH}+1)^{]^{[5 \times(1-\mathrm{BLFE})]}}\right.}
$$

3. $\quad \mathbf{B L S T}=$ 2. Setting $\mathrm{BLFE}=1$ makes BLST $=2$ equivalent to BLST $=0$ except that BLST $=2$ allows you to specify the boundary layer thickness instead of automatically using the local surface mesh size. For BLST $=2$, starting from BLTH from the wall, each newly inserted node will have its location closer to the wall, following this law:

$$
X_{n}=(0.5 \times \mathrm{BLFE})^{n} \times \operatorname{BLTH}(1-0.5 \times \mathrm{BLFE})
$$

4. $\mathbf{B L S T}=3$. For $\operatorname{BLST}=3$, starting from the wall, the position of node $n$ in the normal direction is given by:

$$
X_{n}=\sum_{i=0}^{n} \text { BLFE } \times \text { BLTH }^{i} \text { with } 0 \leq n \leq \text { NELTH }
$$



Figure 8-1. $\mathrm{BLST}=1$ example


Figure 8-2. BLST $=2$ example


Figure 8-3. BLST $=3$ example

## *MESH_BL_SYM

Purpose: Specify the part IDs that will have symmetry conditions for the boundary layer. On these surfaces, the boundary layer mesh follows the surface tangent.

Boundary Layer with Symmetry Condition Cards. Define as many cards as necessary. The next "*" card terminates the input.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | PID1 | PID2 | PID3 | PID4 | PID5 | PID6 | PID7 | PID8 |
| Type | 1 | 1 | 1 | 1 | 1 | 1 | 1 | 1 |
| Default | none | none | none | none | none | none | none | none |

## VARIABLE

PID1, ...

## DESCRIPTION

Part identifiers for the surface element. This is the surface with symmetry.

## *MESH_EMBEDSHELL

Purpose: Define surfaces that the mesher will embed inside the volume mesh. These surfaces will have no thickness and will conform to the rest of the volume mesh having matching nodes on the interface.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | VOLID |  |  |  |  |  |  |  |
| Type | I |  |  |  |  |  |  |  |
| Default | none |  |  |  |  |  |  |  |

Define as many cards as are necessary based on the number of PIDs (the next "*" card terminates the input.)

| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | PID1 | PID2 | PID3 | PID4 | PID5 | PID6 | PID7 | PID8 |
| Type | 1 | 1 | 1 | 1 | 1 | 1 | 1 | 1 |
| Default | none | none | none | none | none | none | none | none |

## VARIABLE

VOLID

PIDn

## DESCRIPTION

ID assigned to the new volume in the keyword *MESH_VOLUME. The surface mesh size will be applied to this volume.

Part IDs for the surface elements that will be embedded in the volume mesh.

## *MESH_INTERF

Purpose: Define the surfaces that will be used by the mesher to specify fluid interfaces in multi-fluid simulations.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | VOLID |  |  |  |  |  |  |  |
| Type | I |  |  |  |  |  |  |  |
| Default | none |  |  |  |  |  |  |  |

Define as many cards as are necessary based on the number of PIDs. This input ends at the next keyword ("*") card.

| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | PID1 | PID2 | PID3 | PID4 | PID5 | PID6 | PID7 | PID8 |
| Type | 1 | 1 | 1 | 1 | 1 | 1 | 1 | 1 |
| Default | none | none | none | none | none | none | none | none |

## VARIABLE

VOLID ID assigned to the new volume in the keyword *MESH_VOLUME. The interface meshes will be applied to this volume.

PIDn Part IDs for the surface elements.

## *MESH_NODE

Purpose: Define a fluid node and its coordinates. These nodes are used in the mesh generation process by the *MESH_SURFACE_ELEMENT keyword, or as user defined volume nodes by the *MESH_VOLUME_ELEMENT keyword.

Node Cards. Include one additional card for each node. This input ends at the next keyword ("*") card.

| Card 1 | 1 | 2 |  | 3 | 4 |  | 5 | 6 |  | 7 | 8 | 9 | 10 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | NID | X |  |  | Y |  |  | z |  |  |  |  |  |
| Type | 1 | F |  |  | F |  |  | F |  |  |  |  |  |
| Default | none | 0 |  |  |  | 0 |  |  | 0 |  |  |  |  |

## VARIABLE

NID
X
$Y \quad y$ coordinate.
$Z \quad z$ coordinate.

## Remarks:

1. The data card format for the *MESH_NODE keyword is identical to *NODE.
2. The *MESH_NODE keyword supersedes *MESH_SURFACE_NODE, which was for surfaces nodes as well as *MESH_VOLUME_NODE for, which was for volume nodes in user defined.

## *MESH_SIZE

Purpose: Define the surfaces that will be used by the mesher to specify a local mesh size inside the volume. If no internal mesh is used to specify the size, the mesher will use a linear interpolation of the surface sizes that define the volume enclosure.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | VOLID |  |  |  |  |  |  |  |
| Type | I |  |  |  |  |  |  |  |
| Default | none |  |  |  |  |  |  |  |

Define as many cards as are necessary based on the number of PIDs (the next "*" card terminates the input.).

| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | PID1 | PID2 | PID3 | PID4 | PID5 | PID6 | PID7 | PID8 |
| Type | 1 | 1 | 1 | 1 | 1 | 1 | 1 | 1 |
| Default | none | none | none | none | none | none | none | none |

## VARIABLE

VOLID

PIDn Part IDs for the surface elements that are used to define the mesh size next to the surface mesh.

## *MESH_SIZE_SHAPE

Purpose: Defines a local mesh size in specific zones corresponding to given geometrical shapes (box, sphere, cylinder and polynomial). The solver will automatically apply the conditions specified during the generation of the volume mesh. This zone does not need to be entirely defined in the volume mesh. In the polynomial case, it is recommended to define several zones for a better mesh size control.

## Remeshing Control Card sets:

Add as many remeshing control cards paired with a case card as desired. The input of such pairs ends at the next keyword "*" card.

Remeshing Control. First card specifies whether to maintain this mesh sizing criterion through a remesh operation.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | SNAME | FORCE | METHOD | BT | DT |  |  |  |
| Type | A | I | I | F | F |  |  |  |
| Default | none | 0 | 0 | 0. | $1 . E 12$ |  |  |  |

Box Case. Card 2 for SNAME = "box" and METHOD = 0

| Cards 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | MSIZE | PMINX | PMINY | PMINZ | PMAXX | PMAXY | PMAXZ |  |
| Type | F | F | F | F | F | F | F |  |
| Default | none | none | none | none | none | none | none |  |

Sphere Case. Card 2 for SNAME = "sphere" and METHOD $=0$

| Cards 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | MSIZE | RADIUS | CENTERX | CENTERY | CENTERZ |  |  |  |
| Type | F | F | F | F | F |  |  |  |
| Default | none | none | none | none | none |  |  |  |

Cylinder Case. Card 2 for SNAME = "cylinder" and METHOD = 0

| Cards 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | MSIZE | RADIUS | PMINX | PMINY | PMINZ | PMAXX | PMAXY | PMAXZ |
| Type | F | F | F | F | F | F | F | F |
| Default | none | none | none | none | none | none | none | none |

Polynomial Case. Card 2 for SNAME $=$ "pol" and METHOD $=0$

| Cards 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | MSIZE | X | Y | Z | NX | NY | NZ |  |
| Type | F | F | F | F | F | F | F |  |
| Default | none | none | none | none | none | none | none |  |

Card 2 for METHOD = 1

| Cards 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | MSIZE | RADIUS | PTID1 | PTID2 |  |  |  |  |
| Type | F | F | 1 | I |  |  |  |  |
| Default | none | none | none | none |  |  |  |  |

VARIABLE
SNAME

FORCE

METHOD

BT/DT

MSIZE Mesh size that needs to be applied in the zone of the shape defined by SNAME

PMIN[X, Y, $\quad x, y$, or $z$ value for the point of minimum coordinates Z]
$\operatorname{PMAX}[X, Y, \quad x, y$, or $z$ value for the point of maximum coordinates Z]

CEN- Coordinates of the sphere center in cases where SNAME is sphere TER[X, Y, Z]

RADIUS Radius of the sphere if SNAME is Sphere or of the cross section disk if SNAME is Cylinder.

## VARIABLE

X/Y/Z
NX/NY/NZ

PTID1

PTID2

## DESCRIPTION

Coordinates of starting point in cases where SNAME is pol.
Direction in which mesh size will be applied in cases where SNAME is pol.

Point ID1 referring to ICFD_DEFINE_POINT. Replaces PMIN, X/Y/Z or CENTER for the various SNAME cases.

Point ID2. Not needed if SNAME is Sphere. Replaces PMAX or NX/NY/NZ for the various SNAME cases.

## *MESH_SURFACE_ELEMENT

Purpose: Specify a set of surface elements (quadrilateral or triangular in 3D and linear segments in 2D) that will be used by the mesher to construct a volume mesh. These surface elements may define the enclosed volume to be meshed, or alternatively they can be used to specify different mesh sizes inside the volume (see card *MESH_SIZE).

Each solver that supports the *MESH volume mesher for building its volume mesh uses the PID given for each surface element specified with this keyword differently.

1. For the *ICFD solver, the *ICFD_PART keyword references the surface mesh PIDs.
2. For the *DUALCESE solver, the MSPIDs in *DUALCESE_BOUNDARY $\qquad$ words cards reference the surface element PIDs. Note that when the dual CESE solver uses *MESH_SURFACE_ELEMENT, this card defines which boundary faces belong to each mesh surface PID. In this case, no other mechanism exists for defining these PIDs. Also, when the dual CESE solver is defined with *MESH cards, there should not be any *DUALCESE_SEGMENTSET cards related to the dual CESE mesh.

Surface Element Card. Define as many cards as necessary. The next keyword ("**) card terminates this input.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 | 9 | 10 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | EID | PID | N1 | N2 | N3 | N 4 |  |  |  |  |
| Type | 1 | I | 1 | 1 | 1 | 1 |  |  |  |  |
| Default | none | none | none | none | none | none |  |  |  |  |

## VARIABLE

EID

N1
N2

PID Mesh surface part ID. A unique identifier for the surface to which this mesh surface element belongs.

## DESCRIPTION

Element ID. A unique number with respect to all *MESH_SURFACE_ELEMENTS cards.

Nodal point 1
Nodal point 2

VARIABLE
N3 Nodal point 3
N4 Nodal point 4

## Remarks:

1. Defining Surface Elements. The convention for defining surface elements is the same as for *ELEMENT_SHELL. In the case of a triangular face, N3 $=$ N4. In 2 D N2 $=$ N3 $=$ N4. Note that the accepted card format is 6 i 8 (not 6i10).

## *MESH_SURFACE_NODE

Purpose: Define a node and its coordinates. These nodes will be used in the mesh generation process by the *MESH_SURFACE_ELEMENT keyword.
*MESH_NODE supersedes this card; so please use *MESH_NODE instead of this card.

Surface Node Cards. Include one card for each node. Include as many cards a necessary. This input ends at the next keyword ("*") card.

| Card 1 | 1 | 2 |  | 3 | 4 |  | 5 | 6 |  | 7 | 8 | 9 | 10 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | NID |  | X |  |  | Y |  |  | Z |  |  |  |  |
| Type | I |  | F |  |  | F |  |  | F |  |  |  |  |
| Default | none |  | 0 |  |  | 0 |  |  | 0 |  |  |  |  |

## VARIABLE

NID

X
$Y \quad y$ coordinate.
$\mathrm{Z} \quad z$ coordinate.

## *MESH_VOLUME

Purpose: This keyword defines the volume space that will be meshed. The boundaries of the volume are the surfaces defined by *MESH_SURFACE_ELEMENT. The surfaces listed have to be non-overlapping, and should not leave any gaps or open spaces between the surface boundaries. On the boundary between two neighbor surfaces, nodes have to be in common (no duplicate nodes) and should match exactly on the interface. They are defined by the keyword *MESH_SURFACE_NODE. This card will be ignored if the volume mesh is specified by the user and not generated automatically.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | VOLID |  |  |  |  |  |  |  |
| Type | 1 |  |  |  |  |  |  |  |
| Default | none |  |  |  |  |  |  |  |

Define as many cards as are necessary based on the number of PIDs (the next "*" card terminates the input.)

| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | PID1 | PID2 | PID3 | PID4 | PID5 | PID6 | PID7 | PID8 |
| Type | 1 | 1 | 1 | 1 | 1 | 1 | 1 | 1 |
| Default | none | none | none | none | none | none | none | none |

VARIABLE

## DESCRIPTION

VOLID ID assigned to the new volume.
PIDn Part IDs for the surface elements that are used to define the volume.

## *MESH_VOLUME_ELEMENT

Purpose: Specify a set of volume elements for the fluid volume mesh in cases where the volume mesh is specified by the user and not generated automatically. The nodal point are specified in the *MESH_VOLUME_NODE keyword. Only tetrahedral elements are supported (triangles in 2D).

Volume Element Card. Define as many cards as necessary. The next "*" card terminates the input.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 | 9 | 10 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | EID | PID | N1 | N2 | N3 | N4 |  |  |  |  |
| Type | 1 | 1 | 1 | 1 | 1 | 1 |  |  |  |  |
| Default | none | none | none | none | none | none |  |  |  |  |

## VARIABLE

EID Element ID. A unique number with respect to all *MESH_VOLUME_ELEMENTS cards.

PID
N1 Nodal point 1.
N2 Nodal point 2.
N3 Nodal point 3.
N4 Nodal point 4.

## Remarks:

1. The convention is the same used by the keyword *ELEMENT_SOLID.

## *MESH_VOLUME_NODE

Purpose: Define a node and its coordinates. This keyword is only used in cases where the fluid volume mesh is provided by the user and is not automatically generated. It serves the same purpose as the *NODE keyword for solid mechanics. Only tetrahedral elements are supported.
*MESH_NODE supersedes this card; so please use *MESH_NODE instead of this card.
Volume Node Cards. Include as many cards in the following format as desired. This input ends at the next keyword ("**") card.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 | 9 | 10 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | NID | X |  | Y |  | Z |  |  |  |  |
| Type | I | F | F |  | F |  |  |  |  |  |
| Default | none | 0 |  | 0 |  | 0 |  |  |  |  |

VARIABLE
NID Node ID. A unique number with respect to the other volume nodes.
$\mathrm{X} \quad x$ coordinate.
$Y \quad y$ coordinate.
$Z \quad z$ coordinate.

## *MESH_VOLUME_PART

Purpose: Associate a volume part number created by a *MESH_VOLUME card with the part number of a part card from a selected solver (designated by the SOLVER field).

Mesh Volume Part Card. Include as many cards in the following format as desired. This input ends at the next keyword ("*") card.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | VOLPRT | SOLPRT | SOLVER |  |  |  |  |  |
| Type | 1 | 1 | A |  |  |  |  |  |
| Default |  |  |  |  |  |  |  |  |

VARIABLE
VOLPRT Part ID of a volume part created by a *MESH_VOLUME card.
SOLPRT Part ID of a part created using SOLVER's part card.
SOLVER Name of a solver using a mesh created with *MESH cards.

## *STOCHASTIC

The keyword *STOCHASTIC is used to describe the particles and numerical details for solving a set of stochastic PDEs. Currently, there are two types of stochastic PDE models in the code: a spray model and a model of embedded particles in TBX explosives. The keyword cards for using these models are:
*STOCHASTIC_SPRAY_PARTICLES
*STOCHASTIC_TBX_PARTICLES
An additional option "_TITLE" may be appended to all *STOCHASTIC keywords. If this option is used, then an 80 character string is read as a title from the first card of that keyword's input. At present, LS-DYNA does not make use of the title. Inclusion of titles gives greater clarity to input decks.

## *STOCHASTIC_SPRAY_PARTICLES

Purpose: Specify particle and other model details for spray modeling using stochastic PDEs that approximate such processes. A pair of cards is required to specify the characteristics of each nozzle (cards 3 and 4 describe the first nozzle).

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | INJDIST | IBRKUP | ICOLLDE | IEVAP | IPULSE | LIMPR | IDFUEL |  |
| Type | I | I | I | I | I | I | 1 |  |
| Default | 1 | none | none | 0 | none | none | 1 |  |


| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | RHOP | TIP | PMASS | PRTRTE | STRINJ | DURINJ |  |  |
| Type | F | F | F | F | F | F |  |  |

Nozzle card 1: Provide as many pairs of nozzle cards 1 and 2 as necessary. This input ends at the next keyword ("*") card (following a nozzle card 2).

| Card 3 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | XORIG | YORIG | ZORIG | SMR | VELINJ | DRNOZ | DTHNOZ |  |
| Type | F | F | F | F | F | F | F |  |

Nozzle card 2: Provide as many pairs of nozzle cards 1 and 2 as necessary. This input ends at the next keyword ("*") card.

| Card 4 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :--- | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | TILTXY | TILTXZ | CONE | DCONE | ANOZ | AMPO |  |  |
| Type | F | F | F | F | F | F |  |  |

## VARIABLE

INJDIST

IBRKUP

ICOLLDE
IEVAP Evaporation flag:
EQ.0: off (no evaporation)
EQ.1: Turn evaporation on (see Remark 1)
IPULSE Type of injection:
EQ.O: continuous injection
EQ.1: sine wave
EQ.2: square wave
LIMPRT Upper limit on the number of parent particles modeled in this spray. This is not used with the continuous injection case (IPULSE $=0$ ).

IDFUEL

## DESCRIPTION

Selected spray liquid fuels:

| EQ.1: | (Default), $\mathrm{H}_{2} \mathrm{O}$ |
| :--- | :--- |
| EQ.2: | Benzene, $\mathrm{C}_{6} \mathrm{H}_{6}$ |
| EQ.3: | Diesel \# 2, $\mathrm{C}_{12} \mathrm{H}_{26}$ |
| EQ.4: | Diesel \# 2, $\mathrm{C}_{13} \mathrm{H}_{13}$ |
| EQ.5: | Ethanol, $\mathrm{C}_{2} \mathrm{H}_{5} \mathrm{OH}$ |
| EQ.6: | Gasoline, $\mathrm{C}_{8} \mathrm{H}_{18}$ |
| EQ.7: | Jet-A, $\mathrm{C}_{12} \mathrm{H}_{23}$ |
| EQ.8: | Kerosene, $\mathrm{C}_{12} \mathrm{H}_{23}$ |
| EQ.9: | Methanol, $\mathrm{CH}_{3} \mathrm{OH}$ |
| EQ.10: | N -dodecane, $\mathrm{C}_{12} \mathrm{H}_{26}$ |

RHOP Particle density
TIP Initial particle temperature.
PMASS Total particle mass
PRTRTE Number of particles injected per second for continuous injection.
STRINJ
DURINJ
Duration of injection(s)
XORIG X-coordinate of center of a nozzle's exit plane
YORIG Y-coordinate of center of a nozzle's exit plane
ZORIG Z-coordinate of center of a nozzle's exit plane
SMR Sauter mean radius
VELINJ Injection velocity
DRNOZ Nozzle radius
DTHNOZ Azimuthal angle (in degrees measured counterclockwise) of the injector nozzle from the $\mathrm{j}=1$ plane.

TILTXY

TILTXZ

CONE

DCONE

ANOZ

AMP0

## DESCRIPTION

Rotation angle (in degrees) of the injector in the $x-y$ plane, where 0.0 points towards the 3 o'clock position ( $\mathrm{j}=1$ line), and the angle increases counterclockwise from there.

Inclination angle (in degrees) of the injection in the $x-z$ plane, where 0.0 points straight down, $x>0.0$ points in the positive $x$ direction, and $x<0.0$ points in the negative $x$ direction.

Spray mean cone angle (in degrees) for hollow cone spray; spray cone angle (in degrees) for solid cone spray.

Injection liquid jet thickness in degrees.
Area of injector
Initial amplitude of droplet oscillation at injector

## Remarks:

1. When IEVAP $=1$, the keyword input file must be modified in a fashion similar to a chemistry problem. This is illustrated in a portion of an example keyword file below. That is, the following keywords need to be used, along with the inclusion of other chemistry-related files (i.e. evap.inp and the corresponding thermodynamics data file):
*CHEMISTRY_MODEL
*CHEMISTRY_COMPOSITION
*CHEMISTRY_CONTROL_FULL
*CESE_INITIAL_CHEMISTRY
```
$ Setup stochastic particles
$
*STOCHASTIC_SPRAY_PARTICLES
$ injdist - ibrk}up icollide
            3 rhop 1, 1
        1000.0 300. 0.01
    the next card is needed for fireball position and max. particle velocity:
        XORIG YORIG ZORIG SMR Velinj Drnoz Dthnoz
        0.005 0.005 1.0e-5 5.0e-6
        200.0 9.0e-5
        TILTXY TILTXZ CONE DCONE
        ANOZ AMPO
        2.5e-8 0.0
$
*CHEMISTRY_MODEL
$ model_id jacsel errlim
```

```
                            1 0
        evap.inp
    therm.dat
        tran.dat
$
*CHEMISTRY COMPOSITION
$ comp_id model_id
$ molefra Species
            1.0 O2
            3.76 N2
$
*CHEMISTRY_CONTROL_FULL
$ sol_id errlim
$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$$
$
$ Set global initial conditions for fluid
$
*CESE_INITIAL_CHEMISTRY
$ sol_id comp_id
$INITIAL CONDITIONS
$ uic vic wic ric pic tic hic
    0.0 0.0 0.0 1.2 101325. 300.0 0.0
```


## *STOCHASTIC_TBX_PARTICLES

Purpose: Specify particle and other model details for stochastic PDEs that model embedded particles in TBX explosives. Note that the components listed on the corresponding *CHEMISTRY_COMPOSITION card are in terms of molar concentrations of the species (in units of moles/[length] ${ }^{3}$, where "[length]" is the user's length unit).

For further information on the theory of the TBX model that has been implemented, a document on this topic can be found at this URL:
http:/ /www.lstc.com/applications/cese_cfd/documentation

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | PCOMB | NPRTCL | MXCNT | PMASS | SMR | RHOP | TICP | T_IGNIT |
| Type | I | I | I | F | F | F | F | F |
| Default | 0 | none | none | none | none | none | none | none |


| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | INITDST | AZIMTH | ALTITD | CPS/CVS | HVAP | EMISS | BOLTZ |  |
| Type | I | F | F | F | F | $F$ | $F$ |  |
| Default | 1 | none | none | none | none | none | none |  |
| Remarks |  |  |  |  |  | 1 | 1 |  |


| Card 3 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | XORIG | YORIG | ZORIG | XVEL | YVEL | ZVEL | FRADIUS |  |
| Type | F | F | F | F | F | $F$ | $F$ |  |
| Default | none | none | none | 0.0 | 0.0 | 0.0 | none |  |

VARIABLE
PCOMB

NPRTCL

MXCNT

PMASS
SMR
RHOP
TICP
T_IGNIT
INITDST

AZIMTH

ALTITD

CPS/CVS Heat coefficient
HVAP Latent heat of vaporization
EMISS Particle emissivity
BOLTZ Boltzmann coefficient
XORIG $\quad x$-coordinate of the origin of the initial reference frame of the TBX explosive

## VARIABLE

YORIG

ZORIG $\quad z$-coordinate of the origin of the initial reference frame of the TBX explosive

XVEL $\quad x$-component of the initial particle velocity the TBX explosive
YVEL $\quad y$-component of the initial particle velocity the TBX explosive
ZVEL $\quad z$-component of the initial particle velocity the TBX explosive
FRADIUS Radius of the explosive area.

## Remarks:

1. If radiation heat transfer is being modeled, then EMISS and BOLTZ are required.

## *LSO

These cards provide a general data output mechanism, causing the creation of a sequence of LSDA files. This facility is intended to allow several different time sequences of data to be output in the same simulation. In addition, any number of domains (and any number of variables on those domains) may be specified within each time sequence. The keyword cards in this section are defined in alphabetical order:
*LSO_DOMAIN
*LSO_ID_SET (not available in the single-precision version of LS-DYNA)
*LSO_POINT_SET
*LSO_TIME_SEQUENCE
*LSO_VARIABLE_GROUP
Note that only the mechanics solver is available in the single-precision version of LSDYNA, and therefore, only LSO mechanics variables are available for output from single precision LS-DYNA. These mechanics variables are listed by domain type in a separate document. This document (LSO_VARIABLES.TXT) is created by running the command: LS-DYNA print_lso_doc. Contrary to LSO_VARIABLES.TXT, element quantities such as stress are not available for output from the mechanics solver to the "lso" database.

An additional option "_TITLE" may be appended to all *LSO keywords. If this option is used, then an 80 character string is read as a title from the first card of that keyword's input. At present, LS-DYNA does not make use of the title. Inclusion of titles gives greater clarity to input decks.

## *LSO_DOMAIN

Purpose: This command provides a way to specify variables on a subset of the domain for a given solver. This domain can be a subset of the mesh used by that solver, a set of output points created with *LSO_POINT_SET, or a set of objects created with *LSO_ID_SET. The frequency and duration of the output for any given domain is determined by each *LSO_TIME_SEQUENCE card that references this *LSO_DOMAIN card. Note that for the single-precision version of LS-DYNA, the only allowed value of SOLVER_NAME $=\mathrm{MECH}$.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | DOMAIN_TYPE |  |  |  |  |  |  |  |
| Type |  |  |  |  |  |  |  |  |


| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | SOLVER_NAME |  |  |  |  |  |  |  |
| Type | A |  |  |  |  |  |  |  |

Special Domains Card. Card 3 when DOMAIN_TYPE is one of ROGO, CIRCUIT, THIST_POINT or TRACER_POINT.

| Card 3 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | OUTID | REFID |  | REDUCT |  |  |  |  |
| Type | 1 | 1 |  | 1 |  |  |  |  |
| Default | none | none |  | none |  |  |  |  |

Miscellaneous Domain Card. Card 3 when DOMAIN_TYPE is one of NODE, PART, SEGMENT, SURFACE_NODE, SURFACE_ELEMENT, VOLUME_ELEMENT, SURFACE_PART, VOLUME_PART.

| Card 3 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | OUTID | REFID | OVERRIDE | REDUCT |  |  |  |  |
| Type | 1 | 1 | 1 | 1 |  |  |  |  |
| Default | none | 0 | 0 | none |  |  |  |  |

Variable Name Card. Provide as many cards as necessary. This input ends at the next keyword ("*") card

| Card 4 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | VARIABLE_NAME |  |  |  |  |  |  |  |
| Type |  |  |  |  |  |  |  |  |

## VARIABLE

DOMAIN_TYPE
SOLVER_NAME

OUTID

REFID

OVERRIDE

## DESCRIPTION

The type of domain for which LSO output may be generated.
Selects the solver from which data is output on this domain. Accepted entries so far are "MECH", "EM", "CESE", and "ICFD".

LSO domain ID associated with this domain, and used by *LSO_TIME_SEQUENCE cards.

Support set ID. This can be a set defined by a *SET card, a *LSO_ID_SET, card, or a *LSO_POINT_SET card. Unless OVERRIDE is specified, this set must be of the same type as DOMAIN_TYPE.

If non-zero, then REFID is interpreted as:
EQ.1: a PART set for SOLVER_NAME
EQ.2: a PART set of volume parts created with a *LSO_ID_SET card (volume parts are defined with *MESH_VOLUME cards).

## VARIABLE

## DESCRIPTION

EQ.3: a PART set of surface parts created with a *LSO_ID_SET card (surface parts are defined with *MESH_SURFACE_ELEMENT cards).
EQ.4: a set of segment sets created with a *LSO_ID_SET card.

|  | EI |
| :---: | :---: |
|  | magneticF electricFí vecpotFie currentDen |
|  | ScalarPote |

REDUCT BLE_NAME

VARIA- Either the name of a single output variable or a variable group. See remarks.

## Remarks:

1. Supported choices for VARIABLE_NAME are listed by DOMAIN_TYPE for each SOLVER_NAME in a separate document. This document (LSO_VARIABLES.TXT) is created by running the command: LS-DYNA print_lso_doc. The following table shows a sample of the point output variables available when DOMAIN_TYPE = THIST_POINT:
*LSO_ID_SET
Purpose: Provides a way to create a set of existing sets (segment sets), or to define a set that is not available with other set-related keyword cards. These are then used in other *LSO cards to specify LSO output. This card is not available in the single precision version of LS-DYNA.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | SETID | TYPE | SOLVER |  |  |  |  |  |
| Type | I | A | A |  |  |  |  |  |
| Default | none | none | MECH |  |  |  |  |  |

Referenced IDs. Provide as many cards as necessary. This input ends at the next keyword ("*") card

| Card | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | ID1 | ID2 | ID3 | ID4 | ID5 | ID6 | ID7 | ID8 |
| Type | I | 1 | 1 | 1 | 1 | 1 | 1 | 1 |
| Default | none | none | none | none | none | none | none | none |

VARIABLE

## DESCRIPTION

SETID
Identifier for this ID set.

SOLVER Name of the solver (MECH, ICFD, CESE, EM, ...)

## VARIABLE

TYPE

ID1, ...

## DESCRIPTION

The kind of IDs in this set:
EQ.'SEG_SETS': Each ID is a segment set connected with SOLVER.

EQ.'CIRCUIT': Each ID is a circuit ID (from *EM cards)
EQ.'SURF_PARTS': Each ID is a surface part number (See *MESH_SURFACE_ELEMENT)

EQ.'VOL_PARTS': Each ID is a volume part number (See *MESH_VOLUME)

EQ.'SURF_ELES': Each ID is a surface element number (See *MESH_SURFACE_ELEMENT)

IDs of the TYPE kind.

## *LSO_POINT_SET

Purpose: Define a list of points used to sample variables in time. Of the different sampling methods, the most common one is to specify points for time history output.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | SETID | USE |  |  |  |  |  |  |
| Type | I | 1 |  |  |  |  |  |  |
| Default | none | 1 |  |  |  |  |  |  |
| Remarks |  | 1 |  |  |  |  |  |  |

Point Cards. Provide as many cards as necessary. This input ends at the next keyword ("*") card

| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | X | Y | Z |  |  |  |  |  |
| Type | F | F | F |  |  |  |  |  |
| Default | none | none | none |  |  |  |  |  |

$\mathrm{X}, \mathrm{Y}, \mathrm{Z} \quad$ Coordinates of a point. As many points as desired can be specified.

VARIABLE
SETID

USE

## DESCRIPTION

Identifier for this point set which is used by *LSO_DOMAIN
Points in this set are used as:
EQ.1: fixed time history points (default)
EQ.2: positions of tracer particles

## Remarks:

1. USE. For USE = 1, with the ICFD and CESE solvers, the fixed points must remain inside the fluid mesh or a zero result is returned, while for the EM solver, the points can be defined inside the conductors or in the air. In the latter case, the fields will be computed using a Biot-Savart type integration. For USE $=2$, a massless tracer particle is tracked for the ICFD and CESE solvers using their local velocity field to integrate the position of each particle in time.

## *LSO_TIME_SEQUENCE

Purpose: This command provides users with maximum flexibility in specifying exactly what they want to have appear in the output LSO binary database. Each instance of the *LSO_TIME_SEQUENCE command creates a new time sequence with an independent output frequency and duration. Furthermore, while the default domain for each output variable will be the entire mesh on which that variable is defined, at all selected snapshot times, the *LSO_DOMAIN keyword commands can be used to specify that output will only occur on a portion of SOLVER_NAME's mesh, and for a limited time interval, or that it will occur at a set of points (see *LSO_POINT_SET), or over a set of object IDs (see *LSO_ID_SET). Note that for the single-precision version of LS-DYNA, the only allowed value of SOLVER_NAME $=\mathrm{MECH}$.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :--- | :--- | :--- | :---: | :---: | :---: | :---: | :---: |
| Variable | SOLVER_NAME |  |  |  |  |  |  |  |
| Type |  |  |  |  |  |  |  |  |


| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | DT | LCDT | LCOPT | NPLTC | TBEG | TEND |  |  |
| Type | F | I | I | I | F | F |  |  |
| Default | 0.0 | 0 | 1 | 0 | 0.0 | 0.0 |  |  |
| Remarks | 1 | 1 | 1 | 1 |  |  |  |  |

Domain IDs. Provide as many cards as necessary. This input ends at the next keyword ("*") card, or when a global variable name card appears

| Card 3 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | DOMID1 | DOMID2 | DOMID3 | DOMID4 | DOMID5 | DOMID6 | DOMID7 | DOMID8 |
| Type | 1 | 1 | 1 | 1 | 1 | 1 | 1 | 1 |
| Default | none | none | none | none | none | none | none | none |

Global variable names. Provide as many cards as necessary. This input ends at the next keyword ("*") card

| Card 4 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | GLOBAL_VAR |  |  |  |  |  |  |  |
| Type |  |  |  |  |  |  |  |  |

## VARIABLE <br> SOLVER_NAM E

DT
LCDT

LCOPT

DESCRIPTION
Selects the solver from which data is output in this time sequence. Accepted entries so far are 'MECH', 'EM', 'CESE' and 'ICFD'

Time interval between outputs.
Optional load curve ID specifying the time interval between dumps.

Flag to govern behavior of plot frequency load curve:
EQ.1: At the time each plot is generated, the load curve value is added to the current time to determine the next plot time (this is the default behavior).
EQ.2: At the time each plot is generated, the next plot time T is computed so that $\mathrm{T}=$ the current time plus the load curve value at the time T.
EQ.3: A plot is generated for each ordinate point in the load curve definition. The actual value of the load curve is ignored.

VARIABLE
NPLTC
TBEG

TEND

DOMID1, ...

GLOBAL_VAR The name of a global output variable computed by SOLVER_NAME. This variable must have a single value (scalar, vector, or tensor), and therefore does not depend upon any DOMID. Any number of such variables may be specified with a given time sequence. These variables are listed as having "global" domain for SOLVER_NAME in a separate document. This document (LSO_VARIABLES.TXT) is created by running the command: LSDYNA print_lso_doc.

## Remarks:

1. If LCDT is nonzero, then it is used and DT and NPLTC are ignored. If LCDT is zero and NPLTC is non-zero, then NPLTC determines the snapshot time increment. If LCDT and NPLTC are both zero, then the minimum non-zero time increment specified by DT is used to determine the snapshot times.

## *LSO_VARIABLE_GROUP

Purpose: To provide a means of defining a shorthand name for a group of variables. That is, wherever the given group name is used, it is replaced by the list of variables given in this command. Note that for the single-precision version of LS-DYNA, the only allowed value of SOLVER_NAME $=\mathrm{MECH}$.

| Card 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | SOLVER_NAME |  |  |  |  |  |  |  |
| Type |  |  |  |  |  |  |  |  |


| Card 2 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | DOMAIN_TYPE |  |  |  |  |  |  |  |
| Type |  |  |  |  |  |  |  |  |


| Card 3 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | GROUP_NAME |  |  |  |  |  |  |  |
| Type | A |  |  |  |  |  |  |  |

List Of Variables In Group. Provide as many cards as necessary. This input ends at the next keyword ("*") card

| Card 4 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| ---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Variable | VAR_NAME |  |  |  |  |  |  |  |
| Type | A |  |  |  |  |  |  |  |

## VARIABLE

## DESCRIPTION

SOLVER_NAME Selects the solver for which data is output in a time sequence.

VARIABLE
DOMAIN_TYPE

GROUP_NAME

## DESCRIPTION

Name of the type of domain on which each VAR_NAME is defined.

Name of (or alias for) the group of names given by the listed VAR_NAMEs

VAR_NAME The name of an output variable computed by SOLVER_NAME

## Remarks:

1. Valid VAR_NAMEs depend both upon the SOLVER_NAME and the DOMAIN_TYPE. These variables are listed by DOMAIN_TYPE for each SOLVER_NAME in a separate document. This document (LSO_VARIABLES.TXT) is created by running the command: LS-DYNA print_lso_doc.
