

Numerical Heat Transfer

Chapter 13 Application examples of fluent for flow and heat transfer problem



Instructor Tao, Wen-Quan; Chen, Li

CFD-NHT-EHT Center

Key Laboratory of Thermo-Fluid Science & Engineering

Xi'an Jiaotong University

Xi'an, 2017-Dec.-20

数值传热学

第 13 章 求解流动换热问题的Fluent软件应用举例



主讲 陶文铨, 陈 黎

西安交通大学能源与动力工程学院
热流科学与工程教育部重点实验室
2017年12月20日, 西安

第 13 章 求解流动换热问题的Fluent软件应用举例

13.1 Heat transfer with source term

13.2 Unsteady cooling process of a steel ball

13.3 Lid-driven flow and heat transfer (Homework of Chapter 1)

13.4 Flow and heat transfer in a micro-channel (4-19)

13.5 Flow and heat transfer in chip cooling (4-19)

13.6 Flow and heat transfer in porous media

13.7 Flow and heat transfer in air film cooling

第 13 章 求解流动换热问题的Fluent软件应用举例

13.1 有内热源的导热问题

导热问题

13.2 非稳态圆球冷却问题

13.3 顶盖驱动流动换热问题

混合对流问题

13.4 微通道内流动换热问题

微通道问题

13.5 芯片冷却流动换热问题

13.6 多孔介质流动换热问题

多孔介质流动

13.7 气膜冷却流动换热问题

湍流

For each example, the general content of the lecture is as follows:

1: Using slides to explain in detail the general **10 steps for Fluent simulation! (PPT讲解)**

- | | |
|--------------------------|------------------------------|
| 1. Read mesh | 2. scale domain |
| 3. Choose model | 4. define material |
| 5. define zone condition | 6. define boundary condition |
| 7. Solution | 8. Initialization |
| 9. Run the simulation. | 10. Post-processing |

2: Operating the Fluent software to simulate the example and post-process the results. (运行软件)

3: Drawing inferences for each example (举一反三) _{5/73}

13.1 Heat transfer with source term

Known: Steady heat conduction in a three-layer solid region with source term in the middle layer. The outer two layers are Aluminum(铝), while the middle layer is nuclear fuel zone. The domain and size is shown in Fig. 1. The boundary conditions are as follows:

- Left and right boundary---adiabatic
- Top and bottom boundary-- convective heat transfer
Heat transfer coefficient: $h=3500 \text{ W}/(\text{m}^2\text{K})$;
Fluid temperature: $T_f=150^\circ\text{C}$.
- In the middle layer--- a constant source term

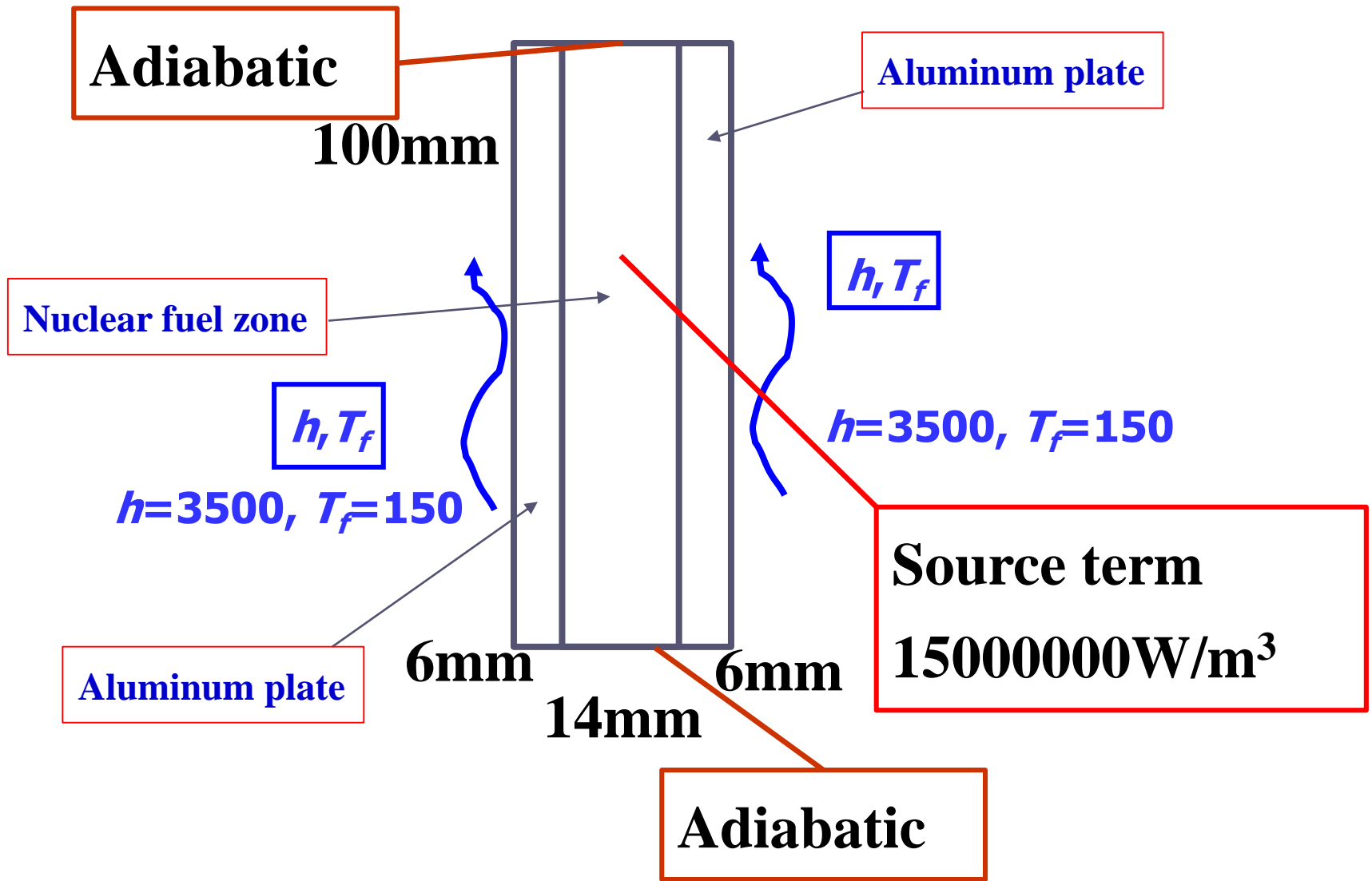


Fig.1 Computational domain

Find: temperature distribution in the domain.

Solution:
$$\text{div}(\Gamma_{\phi} \text{grad} \phi) + S_{\phi}^* = 0$$

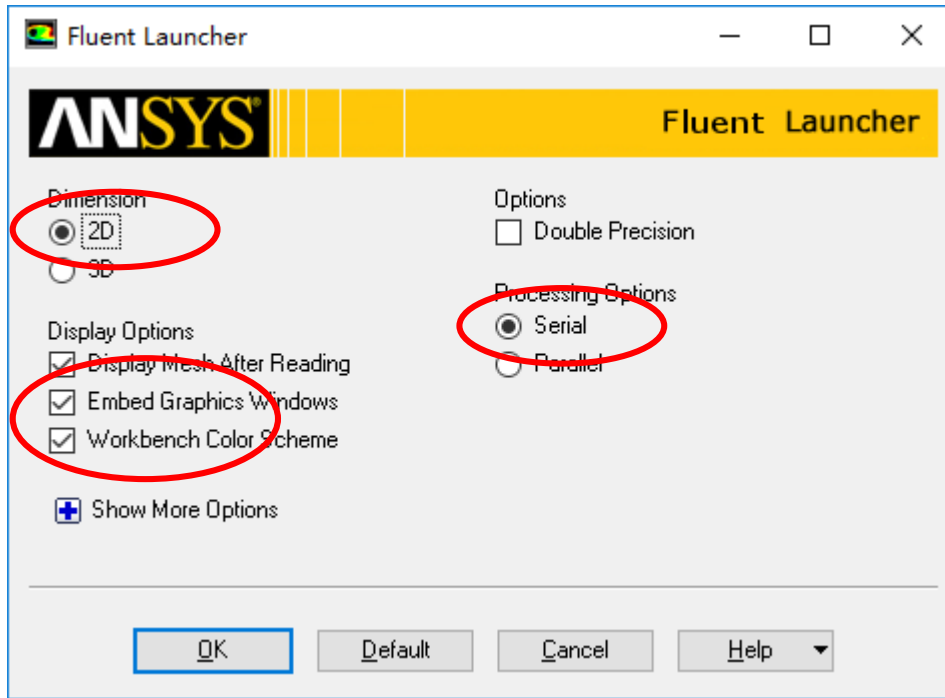
It is a heat conduction problem with given GAMA and source term. GAMA and source term are different for different layers.

Remark: construct the reasonable physical model and write down the right governing equation, BC and IC is the first and most important step before using Fluent.

Fluent is just a tool for solving above problem !

Background of NHT helps you better use the tool.

Start the Fluent software



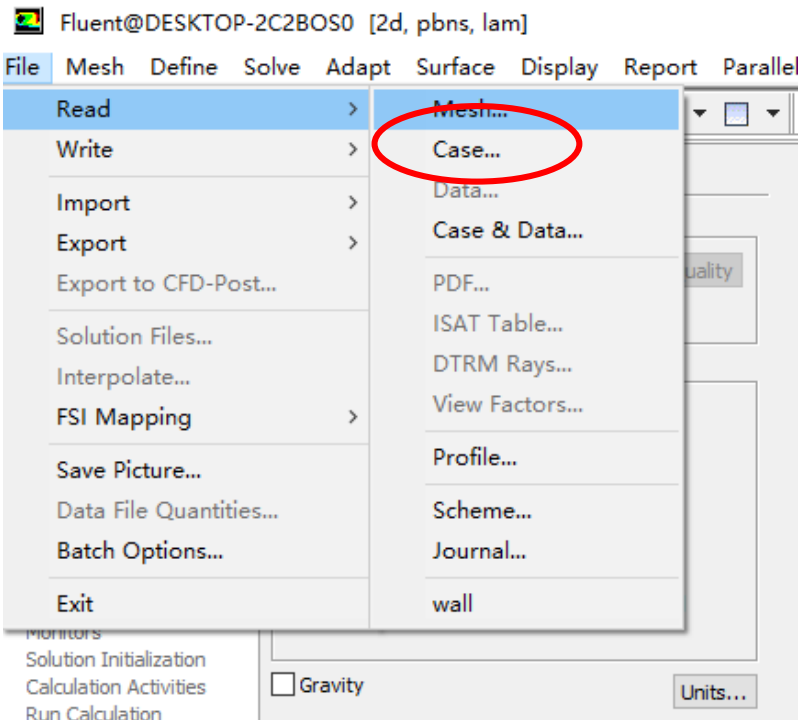
1. Choose **2-Dimension**
2. Choose **display options**
3. Choose **Serial processing option**

Note: Double precision or Single precision

For most cases the single precision version of Fluent is sufficient. For example, for heat transfer problem, **if the thermal conductivity between different components are high**, it is recommended to use Double Precision Version.

Step 1: **Read** and check the mesh

- The mesh is generated by pre-processing software such as **ICEM** and **GAMBIT**. The document is with suffix (后缀名) “**.msh**”
- This step is similar to the Grid subroutine (**UGRID, Setup1**) in our general teaching code.



Mesh → Read

```
> Reading "F:\Fluent\case1\wall.msh"...
Done.
  11457 nodes.
    200 mixed wall faces, zone 5.
    200 mixed wall faces, zone 6.
    56 mixed wall faces, zone 7.
    56 mixed wall faces, zone 8.
  22144 mixed interior faces, zone 10.
    5600 quadrilateral cells, zone 2.
    2800 quadrilateral cells, zone 3.
    2800 quadrilateral cells, zone 4.
```

Step 1: Read and **check** the mesh

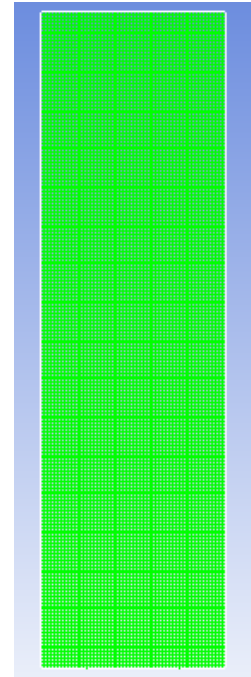
Mesh→Check

- Check the **quality and topological information** of the mesh

```

Mesh Check
  default-interior -> default-interior (10) and default-interior:017 (17)
Domain Extents:
  x-coordinate: min (m) = -1.400000e-02, max (m) = 1.400000e-02
  y-coordinate: min (m) = -5.000000e-02, max (m) = 5.000000e-02
Volume statistics:
  minimum volume (m3): 2.499982e-07
  maximum volume (m3): 2.500008e-07
  total volume (m3): 2.800000e-03
Face area statistics:
  minimum face area (m2): 4.999973e-04
  maximum face area (m2): 5.000010e-04
Checking mesh.....
Done.

```



- Sometimes the check will be failed if the quality is not good or there is a problem with the mesh.

```

Face area statistics:
WARNING: invalid or face with too small area exists.
minimum face area (m2): 0.000000e+00
maximum face area (m2): 5.081937e-03

```

```

WARNING: Mesh check failed.

```

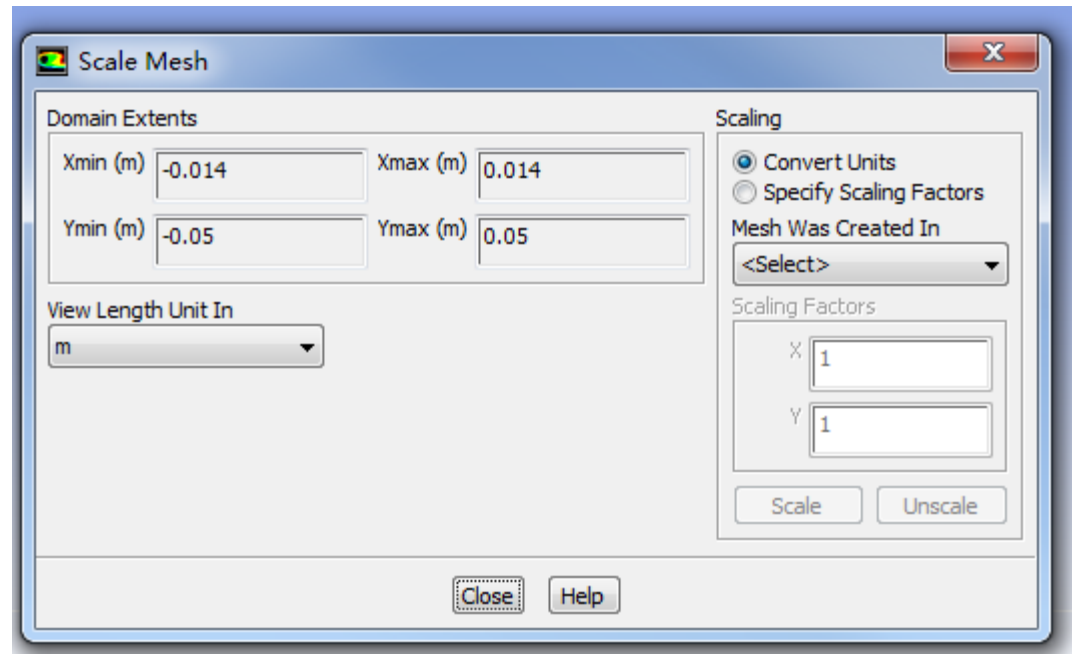
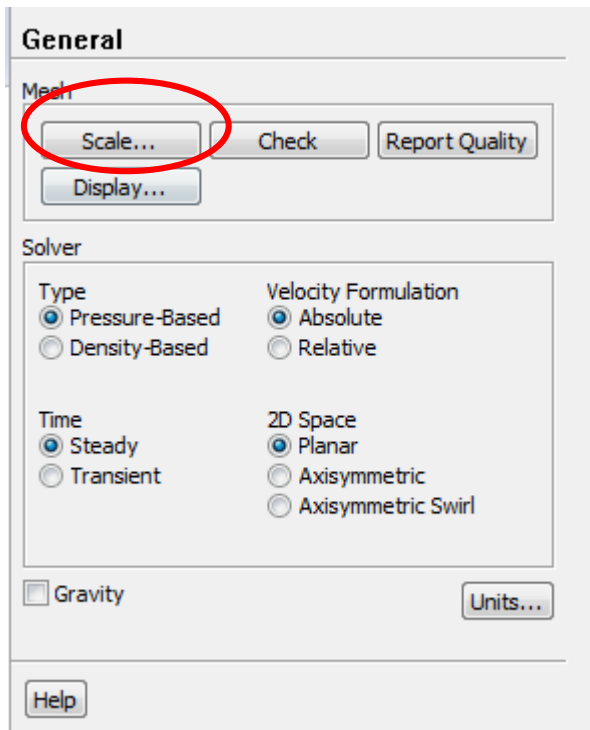
```

WARNING: The mesh contains high aspect ratio quadrilateral,
hexahedral, or polyhedral cells.

```

Step 2: Scale the domain size

General → Scale

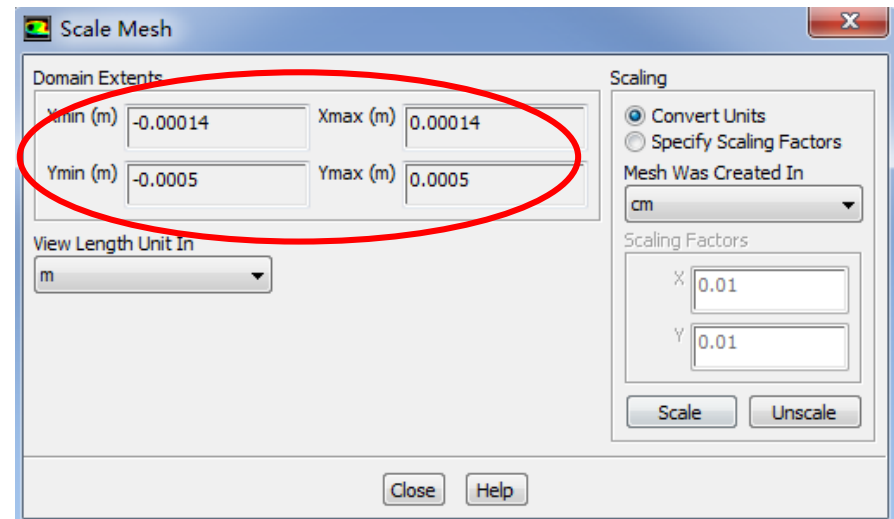
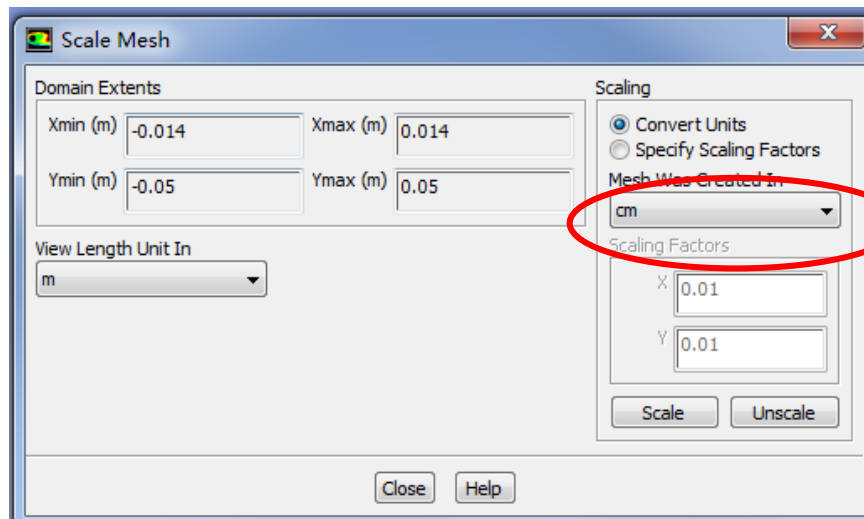


- **Fluent stores the mesh in units as “m”, SI unit. You can show it in different units such as cm, mm, in, or ft.**

- You also can scale the domain size use “Convert Units” or “Specify Scaling Factors” command.

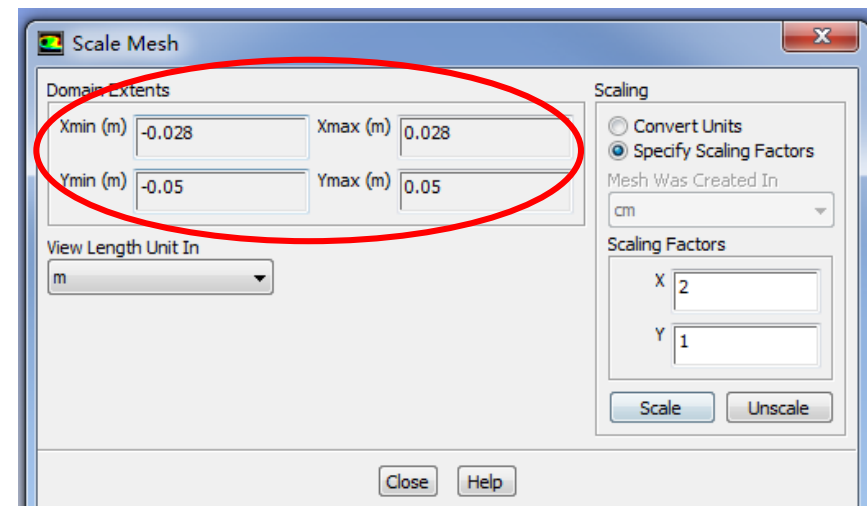
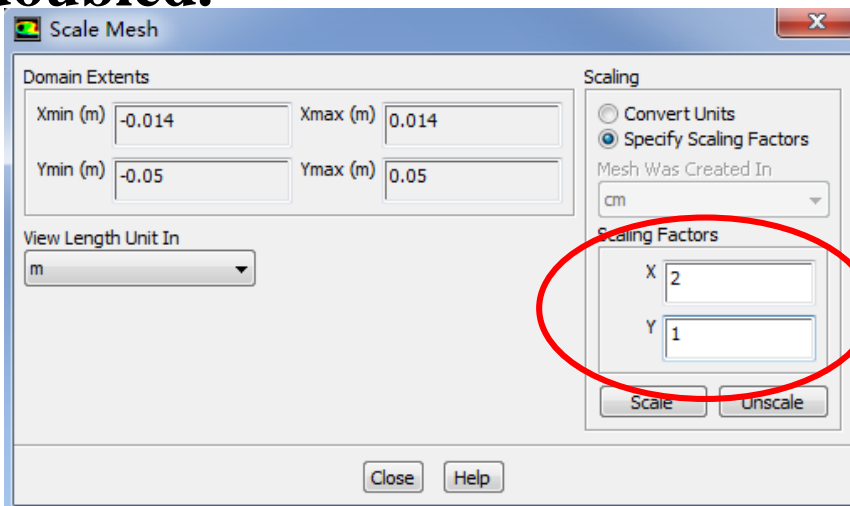
Remark: Fluent thought you create the mesh in units of m. However, if your mesh is created in a different unit, such as cm, you **must** use **Convert Units Command** to change the mesh into the right size. The values will be multiplied by the Scaling Factor.

ICEM: 1 cm -> Fluent: 1m -> Scale: cm, factor: 0.01



- “Specify Scaling factors”, by using this you can define your own scaling factor.

Remark: In “Convert Unit”, the scaling factor is fixed, You can also use “Specify Scaling Factors” to define your own scaling factor. For example, in the following the size of x direction is doubled.



- If you click the **Scale button** by accident or use the wrong Scaling factor, you can click the **Unscale button**. The old size will be recovered,

Step 3: Choose the physicochemical model

Based on the governing equations you are going to solve, select the related model in Fluent.

Remark: Understand the problem you are going to solve, and write down the right governing equations is the first and most important step for numerical simulation. Without background of “Fluid mechanics”, “Heat Transfer” and “numerical heat transfer”, it is hard to complete this step for fluid flow and heat transfer problem.

$$\text{div}(\Gamma_{\phi} \text{grad} \phi) + S_{\phi}^* = 0$$

This is the equation we will solve, the energy equation for heat conduction in solid, without radiation and convection.

Remark: The governing equation of energy solved by Fluent is as follows (Ansys Help: 5.2.1)

$$\frac{\partial}{\partial t}(\rho E) + \nabla \cdot (\bar{v} (\rho E + p)) = \nabla \cdot \left(k_{eff} \nabla T - \sum_j h_j \bar{J}_j + (\bar{\tau}_{eff} \cdot \bar{v}) \right) + S_h$$

$$E = h - \frac{p}{\rho} + \frac{v^2}{2}$$

Enthalpy

8.1 Format Improvement of General Governing Equation

$$\frac{\partial(\rho c_p T)}{\partial t} + \frac{\partial(\rho c_p u T)}{\partial x} + \frac{\partial(\rho c_p v T)}{\partial y} = \frac{\partial}{\partial x} \left(\lambda \frac{\partial T}{\partial x} \right) + \frac{\partial}{\partial y} \left(\lambda \frac{\partial T}{\partial y} \right) + S_T.$$

$$\frac{\partial(\rho T)}{\partial t} + \frac{\partial(\rho u T)}{\partial x} + \frac{\partial(\rho v T)}{\partial y} = \frac{\partial}{\partial x} \left(\frac{\lambda}{c_p} \frac{\partial T}{\partial x} \right) + \frac{\partial}{\partial y} \left(\frac{\lambda}{c_p} \frac{\partial T}{\partial y} \right) + \frac{S_T}{c_p}$$

$$-\frac{1}{c_p^2} \left[\rho c_p T \frac{\partial c_p}{\partial t} + \left(\rho c_p u T - \lambda \frac{\partial T}{\partial x} \right) \frac{\partial c_p}{\partial x} + \left(\rho c_p v T - \lambda \frac{\partial T}{\partial y} \right) \frac{\partial c_p}{\partial y} \right]$$

Improved format of the general G.E.

The frame work of the previous G.E. is retained (保留), but the diffusion coefficient is resumed to (恢复到) its original value by introducing a **nominal density** as follows:

$$\frac{\partial(\rho^* \phi)}{\partial t} + \text{div}(\rho^* \phi \vec{U}) = \text{div}(\Gamma_\phi \text{grad} \phi) + S_\phi^*$$

The new form of G.E. are:

Equation	ρ^*	ϕ	Γ_ϕ	S_ϕ^*
Continuity equation	ρ	1	0	0
Momentum eqn. (x direction)	ρ	u	μ	$\rho f_x - \frac{\partial p}{\partial x}$
Momentum eqn. (y direction)	ρ	v	μ	$\rho f_y - \frac{\partial p}{\partial y}$
Energy equation	ρc_p	T	λ	S_T

Fluent for solid phase:

V is rotational or translational motion of solids

h : enthalpy

$$\frac{\partial}{\partial t}(\rho h) + \nabla \cdot (\bar{v} \rho h) = \nabla \cdot (k \nabla T) + S_h$$

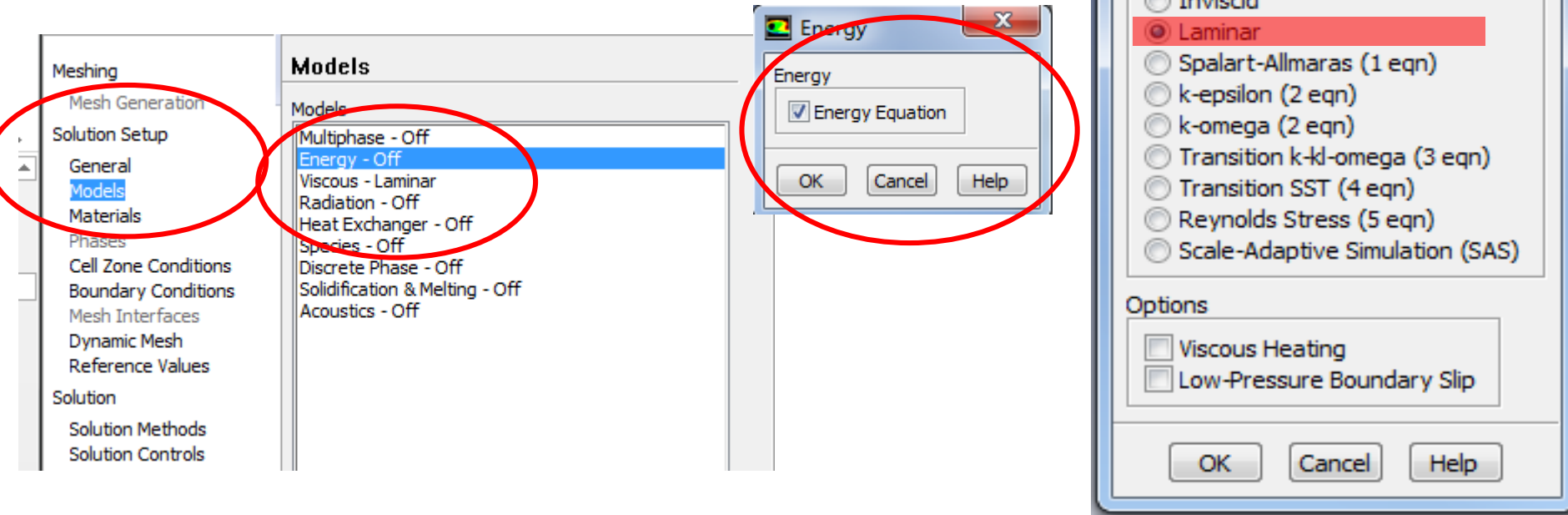
Our general Code:

$$\frac{\partial(\rho c_p T)}{\partial t} + \frac{\partial(\rho c_p u T)}{\partial x} + \frac{\partial(\rho c_p v T)}{\partial y} = \frac{\partial}{\partial x} \left(\lambda \frac{\partial T}{\partial x} \right) + \frac{\partial}{\partial y} \left(\lambda \frac{\partial T}{\partial y} \right) + S_T.$$

Such a treatment is much better than taking Γ / c_p as a nominal diffusion coefficient and S_T / c_p as a nominal source term.

To select the model, the command is as follows:

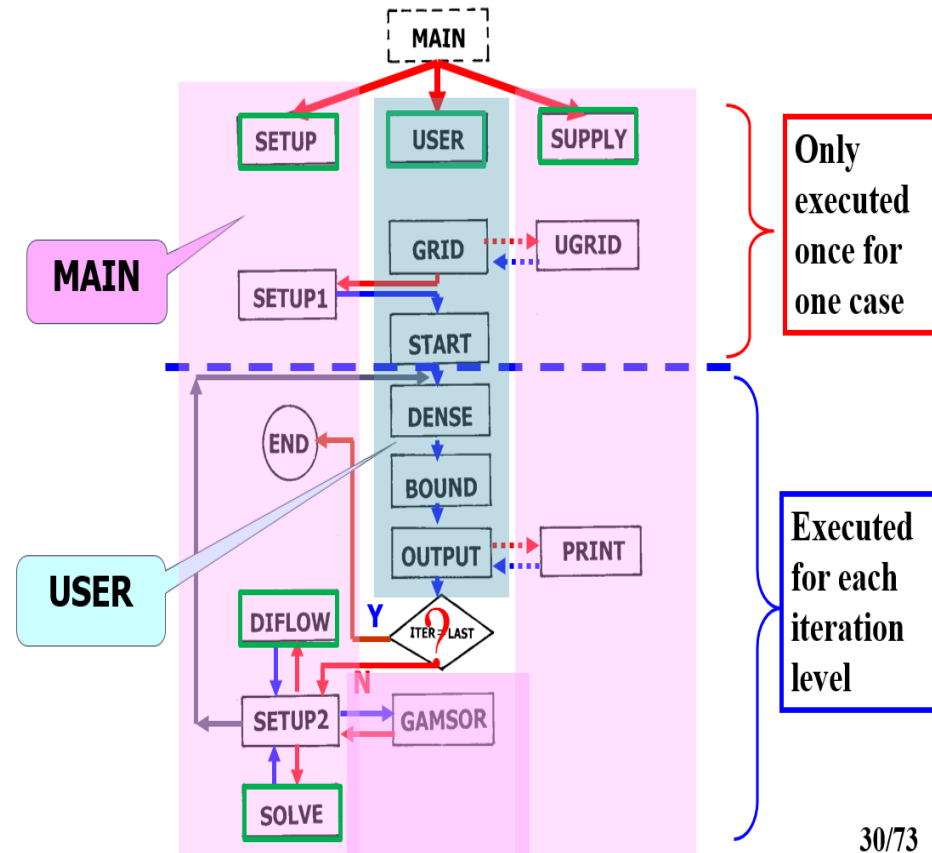
Solution Setup → Model



Remark: It is interesting to note that no matter what kind of problems you will solve, the model “Viscous-Laminar” is always activated. We can cancel it later.

Remark: In our general code,

In SETUP2, Visit NF from 1 to NFMAX in order; When some value of NF is visited and LSOLVE(NF)=.T. , then this variable is solved; Similarly in PRINT SUBROUTINE NF is visited form 1 to NFX4(=14) in order , as long as LPRINT(NF) = .T. , the variable is printed out.



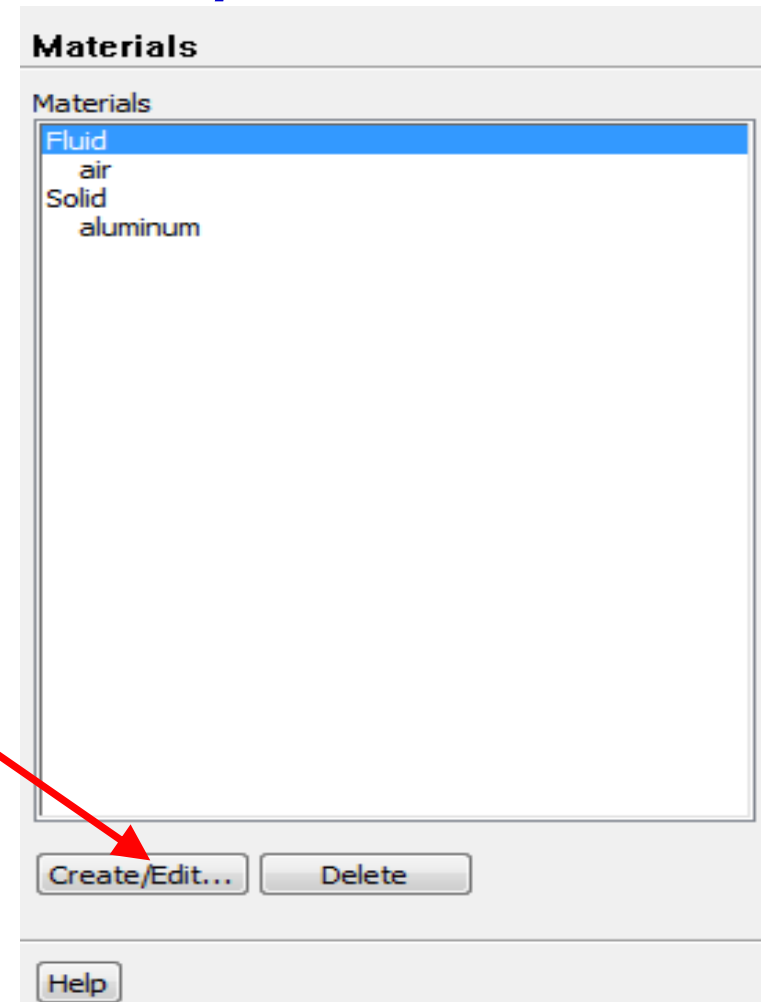
Step 4: Define the material properties

Define the properties required for modeling! For pure heat conduction problem studied here, ρ , C_p and λ should be defined.

Solution Setup → **Materials**

In Fluent, the default fluid is **air** and the default solid is **Al**.

Click the Create/Edit button to find or define new material!



Create/Edit Materials

Name: aluminum

Material Type: solid

Chemical Formula: al

Fluent Solid Materials: aluminum (al)

Mixture: none

Order Materials by:
 Name
 Chemical Formula

Fluent Database...
User-Defined Database...

Properties

Density (kg/m ³)	constant	Edit...
Cp (Specific Heat) (j/kg-k)	constant	Edit...
Thermal Conductivity (w/m-k)	constant	Edit...

871

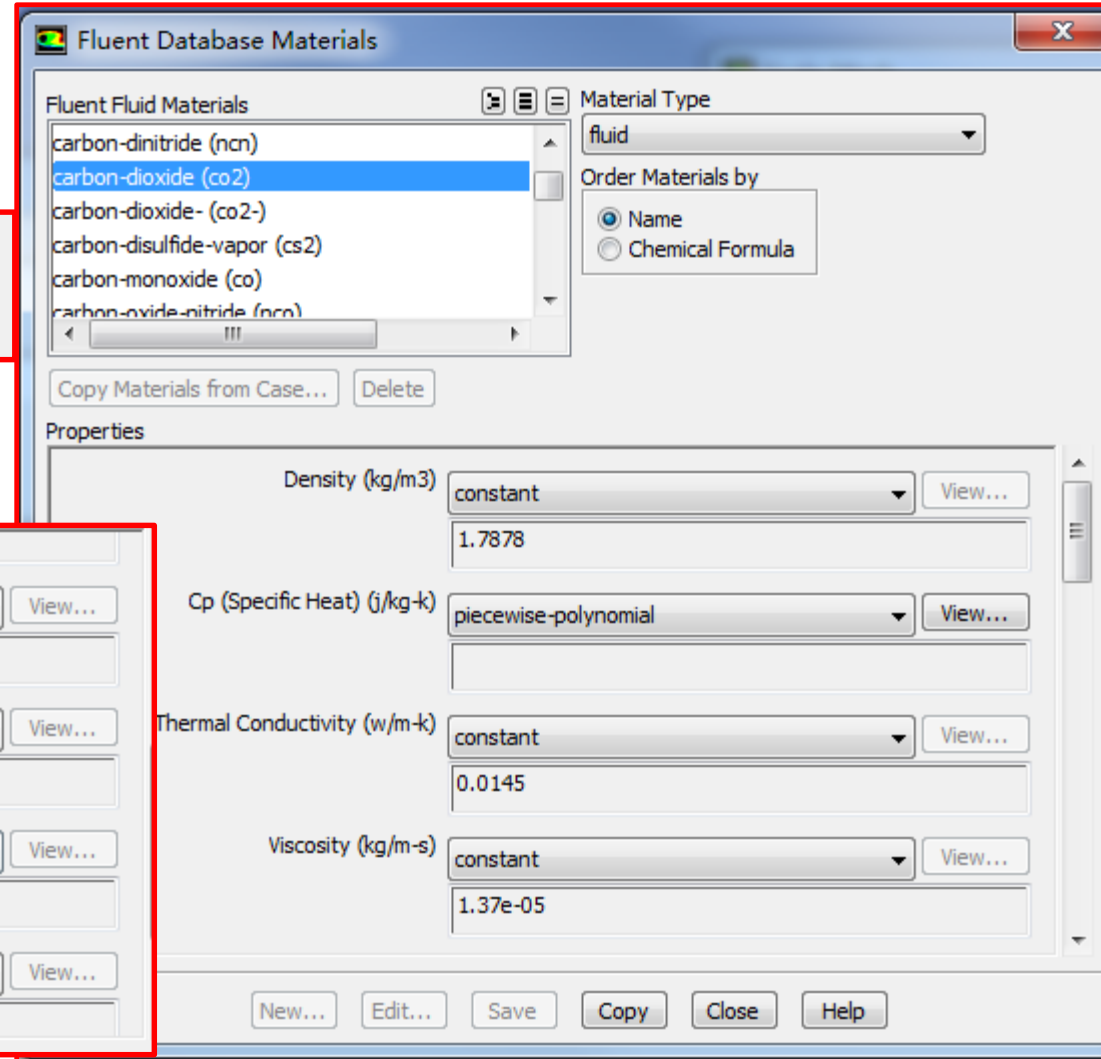
202.4

Change/Create Delete Close Help

**Fluent provide a lot of materials in its database. Usually,
You can find the material you need in the database.**

For example, CO₂

Acentric Factor	constant	View...
	0.33	



Fluent Database Materials

Fluent Fluid Materials

- carbon-dinitride (ncn)
- carbon-dioxide (co2)**
- carbon-dioxide- (co2-)
- carbon-disulfide-vapor (cs2)
- carbon-monoxide (co)
- carbon-oxide-nitride (nco)

Material Type: fluid

Order Materials by:

- Name
- Chemical Formula

Copy Materials from Case... Delete

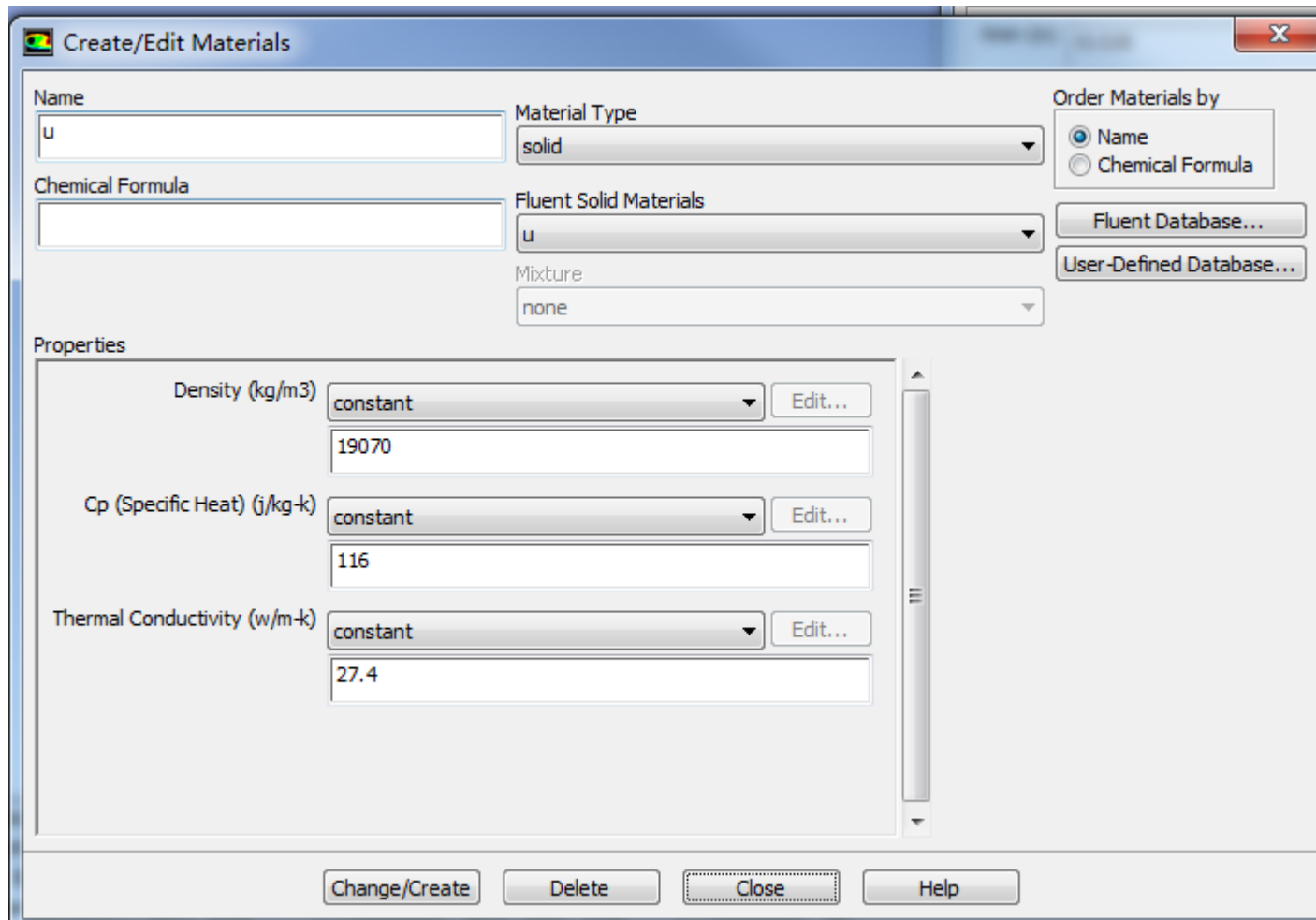
Properties

Density (kg/m3)	constant	View...
	1.7878	
Cp (Specific Heat) (j/kg-k)	piecewise-polynomial	View...
Thermal Conductivity (w/m-k)	constant	View...
	0.0145	
Viscosity (kg/m-s)	constant	View...
	1.37e-05	

New... Edit... Save Copy Close Help

	0.000551	
Molecular Weight (kg/kgmol)	constant	View...
	58.0498	
Critical Temperature (k)	constant	View...
	508.1	
Critical Pressure (pascal)	constant	View...
	4700000	
Critical Specific Volume (m3/kg)	constant	View...
	0.0036	

However, it will happen that the material you need is not in the database. You can input it manually.

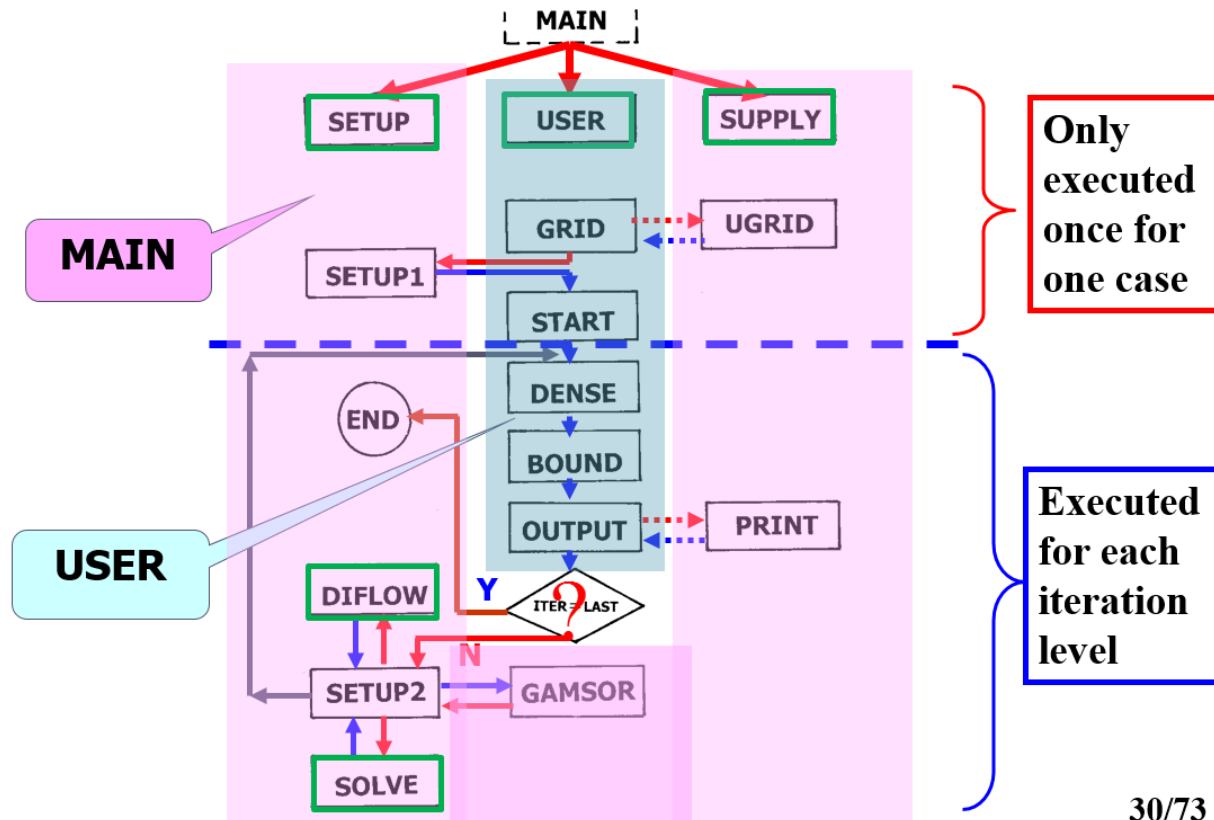


Our general Code:

12. GAMSOR

(1) Determine Γ_ϕ for different variables:

$$u, v - \eta ; T - \lambda$$



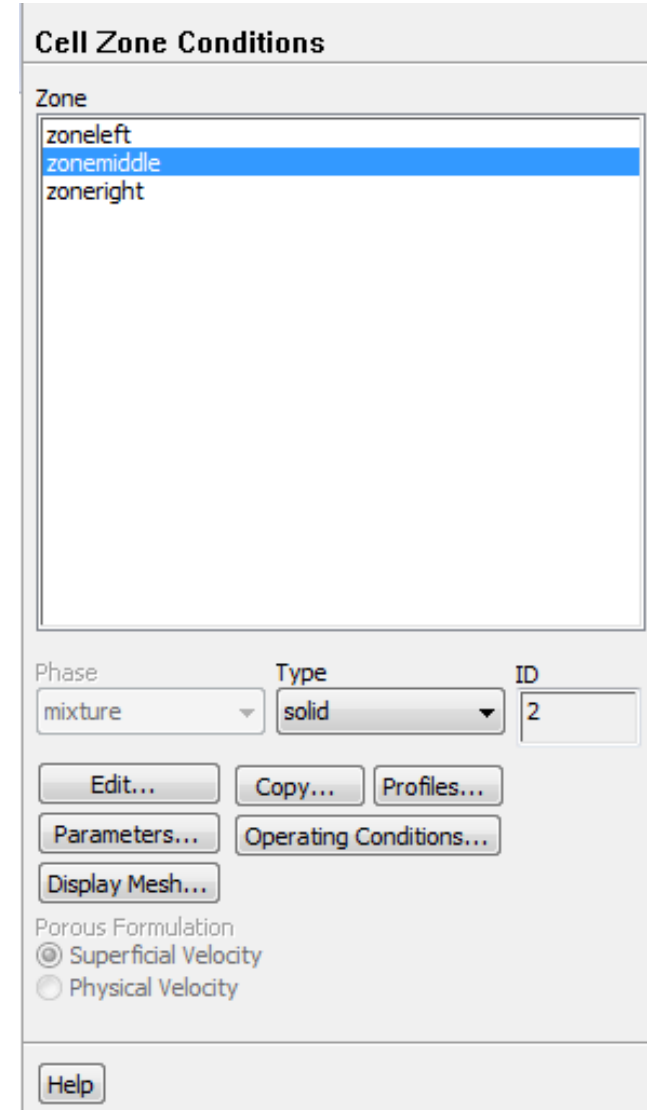
Step 5: Define zone condition

Solution Setup → Cell Zone Condition

Zone type:

- Fluid
- Solid

Porous media is treated as a type of fluid zone, in which parameters related to porous media should be given such as porosity, permeability (渗透率), etc. We will discuss it in Example 6.



Cell Zone Conditions

Zone

- zoneleft
- zonemiddle**
- zoneright

Phase: mixture Type: solid ID: 2

Edit... Copy... Profiles...

Parameters... Operating Conditions...

Display Mesh...

Porous Formulation

Superficial Velocity

Physical Velocity

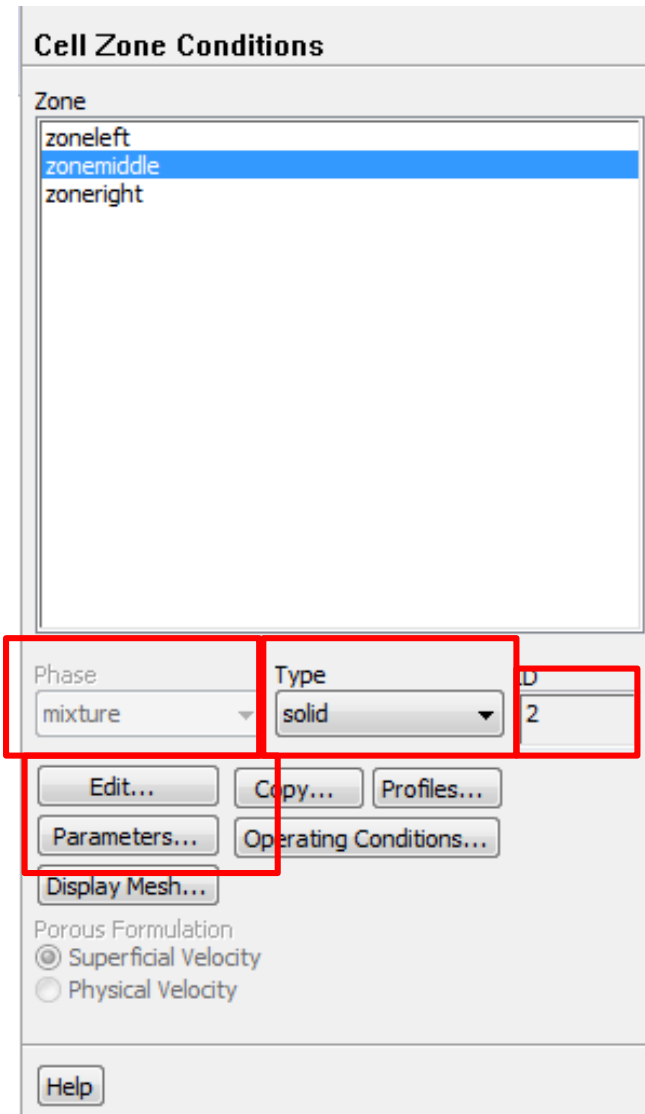
Help

Each zone has its ID.

Each zone should be assigned a type, either fluid or solid.

Phase is not activated here. It can be edited under other cases such as multiphase (多相流) flow model is activated.

Click Edit to define the zone condition of each zone.



Cell Zone Conditions

Zone

- zoneleft
- zonemiddle
- zoneright

Phase	Type	ID
mixture	solid	2

Edit... Copy... Profiles...
Parameters... Operating Conditions...
Display Mesh...

Porous Formulation
 Superficial Velocity
 Physical Velocity

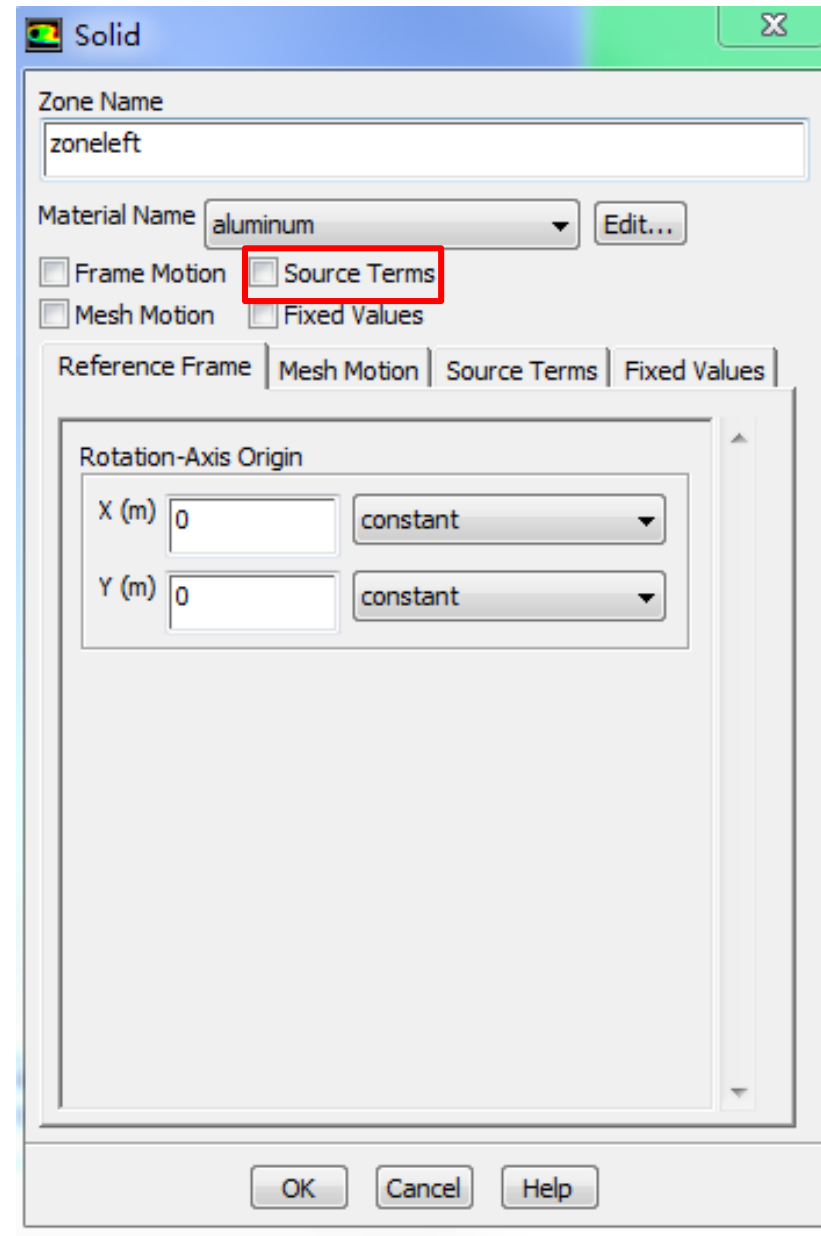
Help

A kind of material should be given to the zone, such as **Al** here for the two out layers.

Frame motion and Mesh motion is used if the solid or the frame is moving.

If T of the zone is fixed, you can select the Fixed value button.

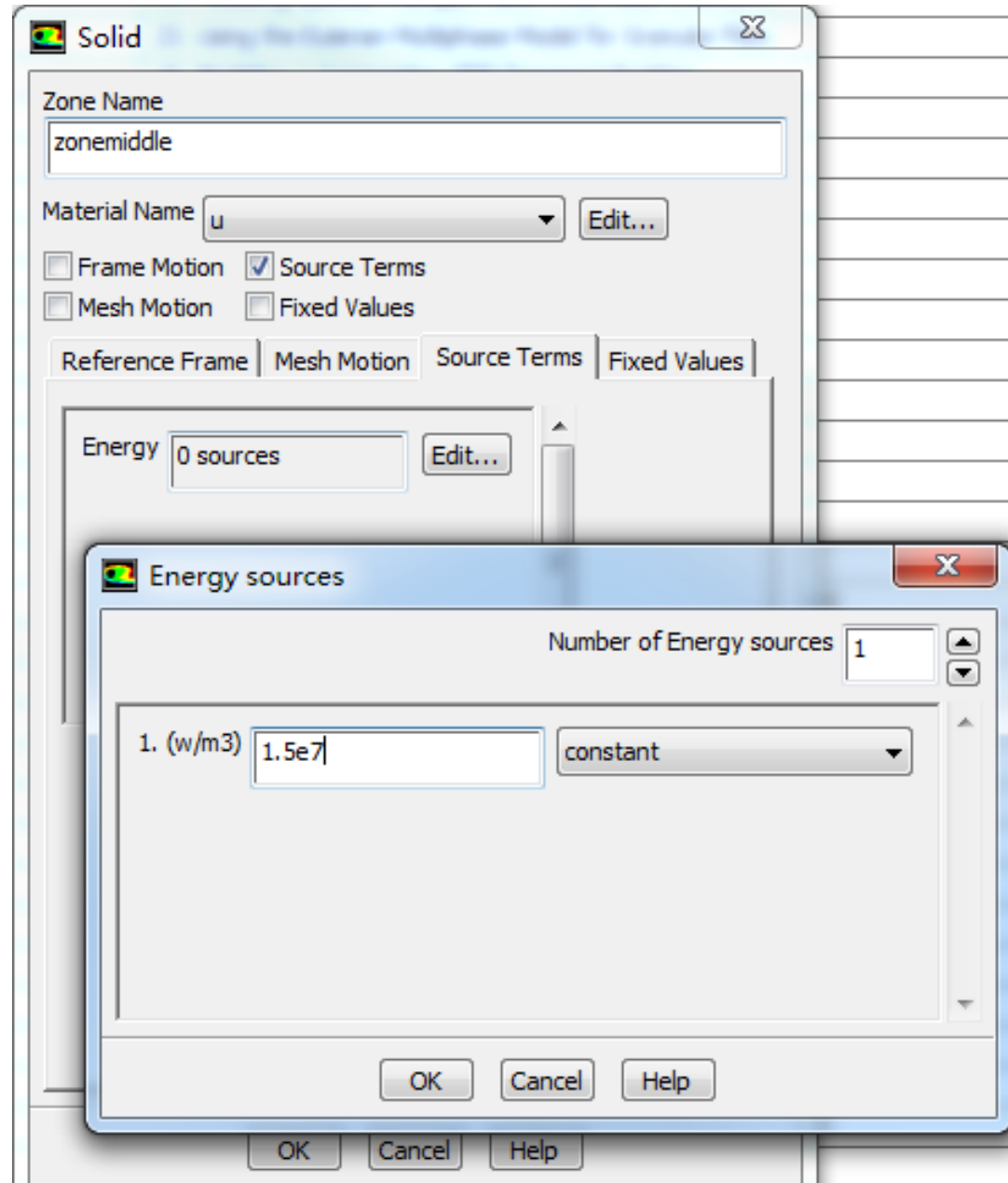
Source term should be selected if there is a source term in the zone.



For the middle zone, source term is activated, and the value of the source term is input in the **Energy Sources** interface.

Remark: Source term is given in GAMSOR, which is linearized and A_p and C_{on} is given.

$$S = S_C + S_P \phi_P, S_P \leq 0$$



Remark: In Fluent, if the source term is not a constant and is a function of the variable solved, local linearization of source term is also adopted.

$$S = S_C + S_P \phi_P, S_P \leq 0$$

- 1. Enhance the solution stability;**
- 2. Accelerate the convergence rate.**

For general source term that is not a constant, user defined function (UDF) is required in Fluent. Define_Source is adopted to specify custom source term for different transport equations.

Step 6: Define the boundary condition

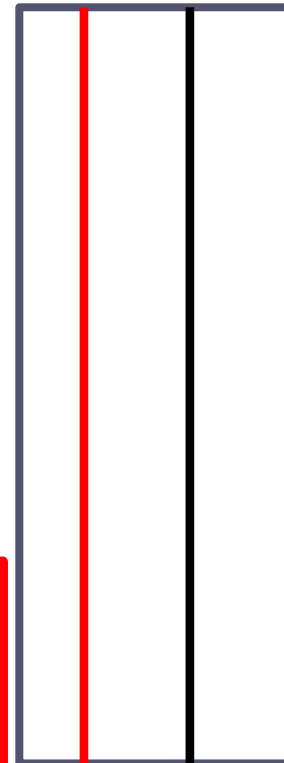
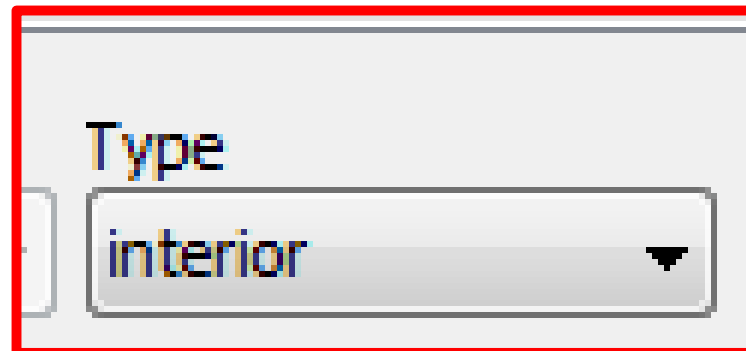
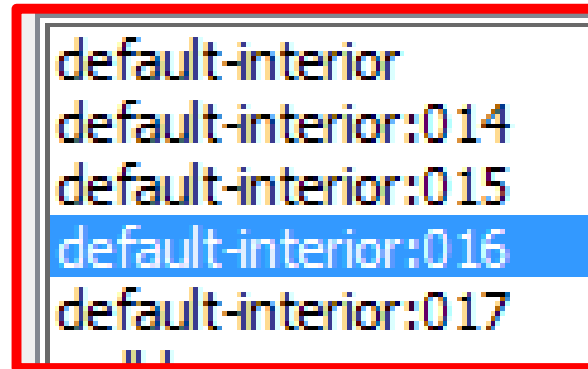
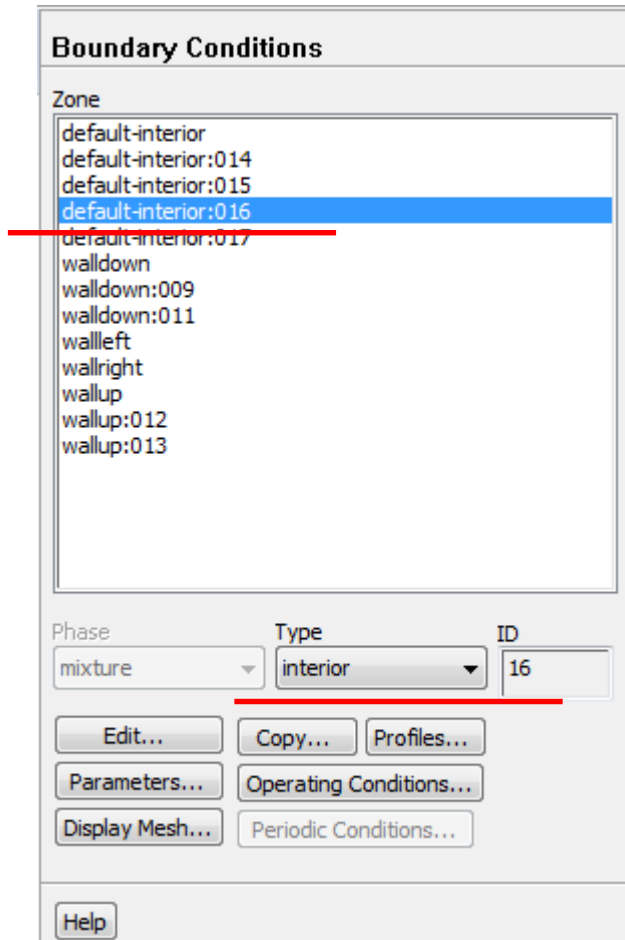
Boundary condition definition is one of the most important and difficult step during Fluent simulation. General boundary conditions in Fluent can be divided into two kinds:

1. BC at inlet and outlet: pressure, velocity, mass flow rate, outflow...

2. BC at wall: wall, periodic, symmetric...

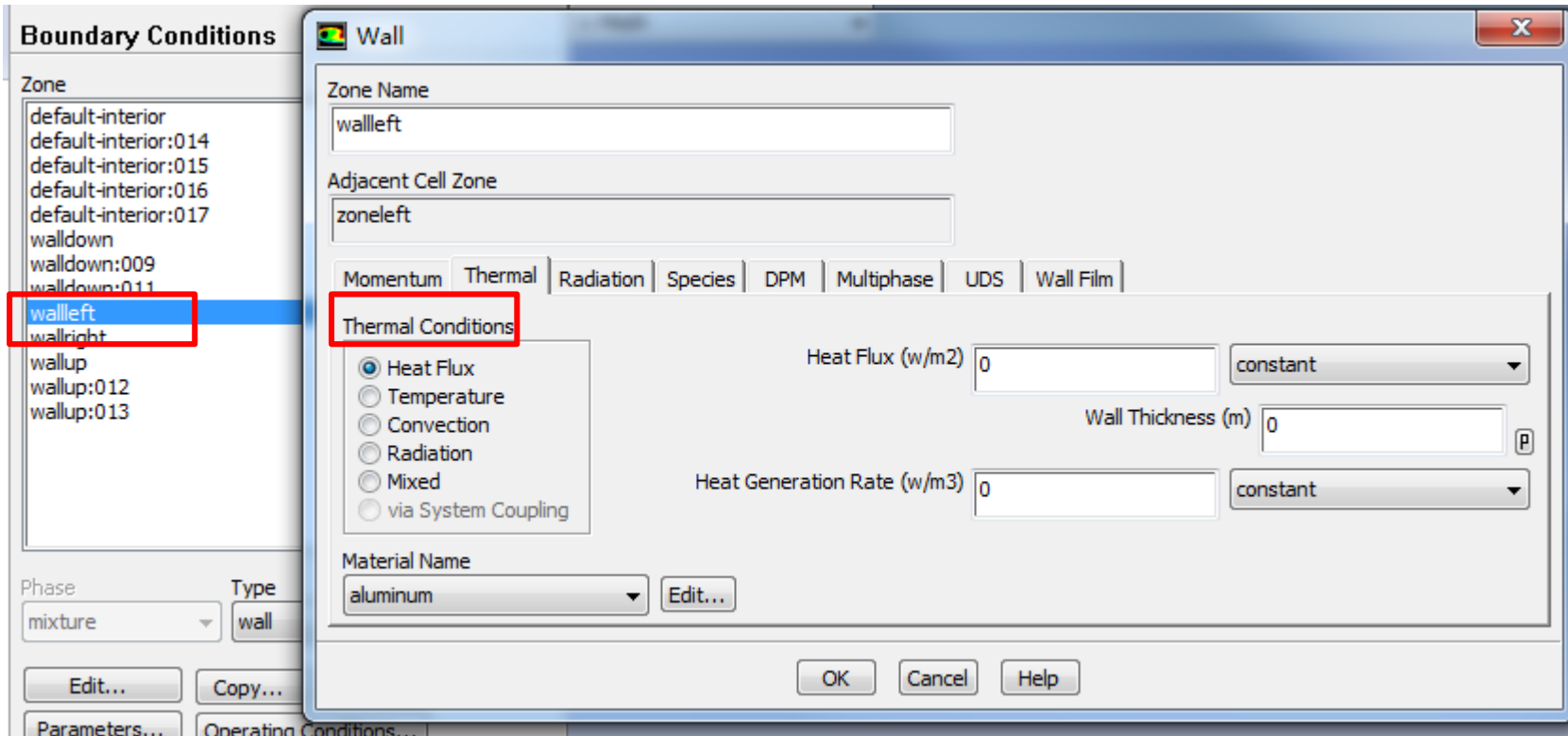
Remark: Interior cell zone and interior interface will also shown in the BC Window.

For example, **Default-interior:016** is also listed here. However, it is not a Boundary of the domain. Its type is **Interior** here.



Here, only the BCs related to the heat conduction problem studied here are introduced. Other types of BCs will be introduced in other examples.

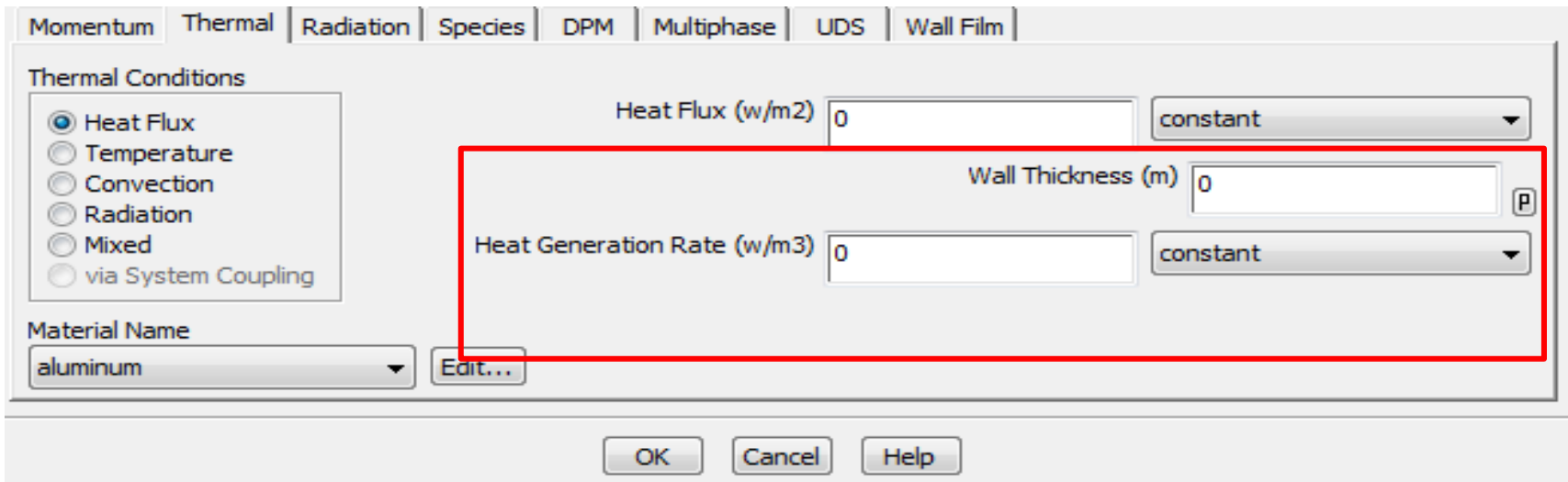
Solution Setup → Boundary conditions



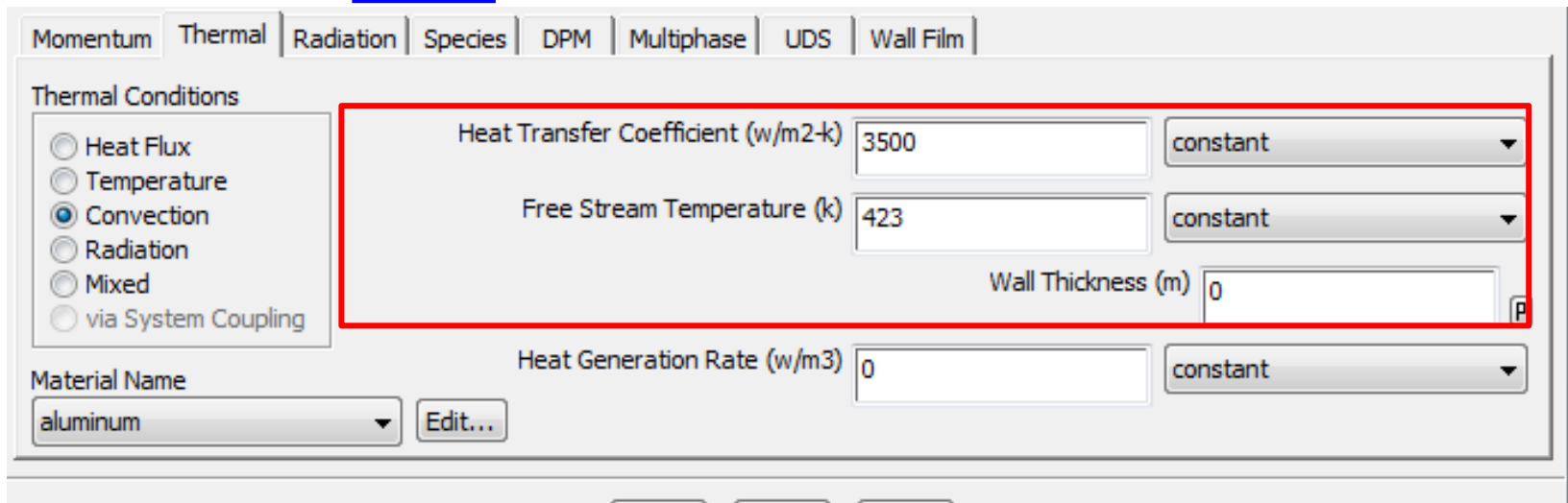
1. Heat Flux (2nd BC):
2. Temperature (1st BC)
3. Convection (3rd BC)

Input the related values

Here, if the wall thickness is greater than zero, and you describe it as a line in your model, a heat generation rate within the wall should also be given. You should also assign the wall a kind of material.



The screenshot shows the 'Thermal Conditions' dialog box in ANSYS Fluent. The 'Thermal Conditions' section has 'Heat Flux' selected. The 'Heat Flux (w/m2)' is set to 0 with a 'constant' dropdown. The 'Wall Thickness (m)' is set to 0. The 'Heat Generation Rate (w/m3)' is set to 0 with a 'constant' dropdown. The 'Material Name' is set to 'aluminum'. A red box highlights the 'Wall Thickness (m)' and 'Heat Generation Rate (w/m3)' fields. At the bottom are 'OK', 'Cancel', and 'Help' buttons.

h, T_f
 $h=3500, T_f=150$


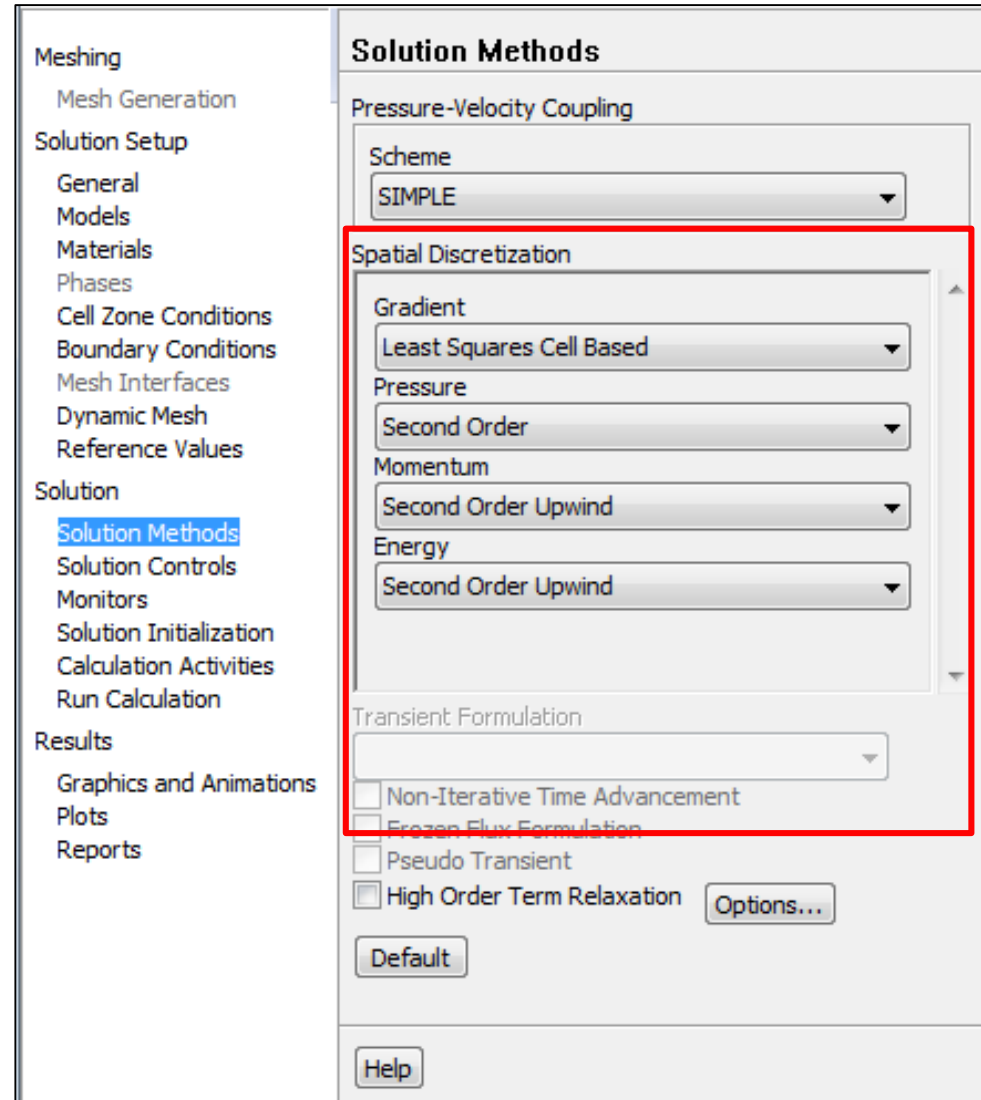
The screenshot shows the ANSYS Fluent software interface for setting thermal boundary conditions. The 'Thermal Conditions' panel is active, with 'Convection' selected. The 'Heat Transfer Coefficient (w/m2-k)' is set to 3500, 'Free Stream Temperature (k)' is set to 423, and 'Wall Thickness (m)' is set to 0. The 'Material Name' is set to 'aluminum'. The 'Heat Generation Rate (w/m3)' is set to 0. A red box highlights the convection parameters.

Remark: Other types of thermal BC in Fluent include radiation BC, mixed BC(combined radiation and convection BC), thermal conditions for two-sides walls, shell conduction.... For more details, you can refer to the Help File of Fluent.

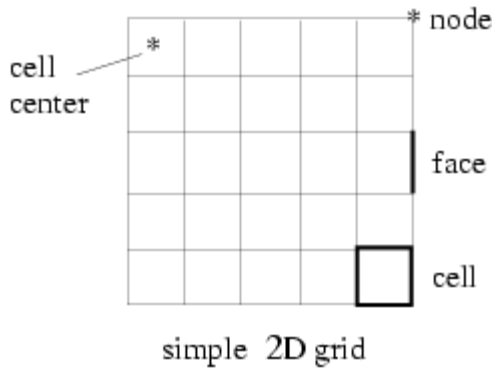
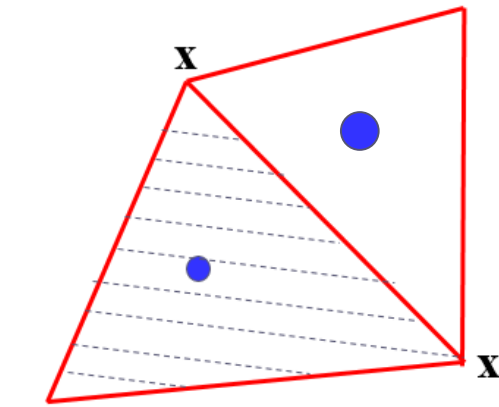
Step 7: Solution setup: algorithm and scheme




Remark: In Fluent, for the SIMPLE series algorithms, only **SIMPLE** and **SIMPLEC** are included.

Review: What is the difference between SIMPLE, SIMPLEC and SIMPLER?



Remark: Difference between the terminology in our NHT (Practice B) and Fluent software about the mesh information.



	Our NHT	Fluent
	Node/cell center	Cell center
	x	Node
	Interface	Interior face

Interface in Fluent is particularly used for the face between different materials.

Gradient calculation,
There are three schemes.

Gradient

Least Squares Cell Based

Green-Gauss Cell Based

Green-Gauss Node Based

Least Squares Cell Based

$\nabla \phi$

1. Green-Gauss Cell-Based (格林-高斯基于单元法)
2. Green-Gauss Node-Based (格林-高斯基于节点法)
3. Least-Squares Cell Based 基于单元体的最小二乘法
It is the default scheme for gradient calculation.

Green-Gauss Theory:

The averaged gradient over a control domain is:

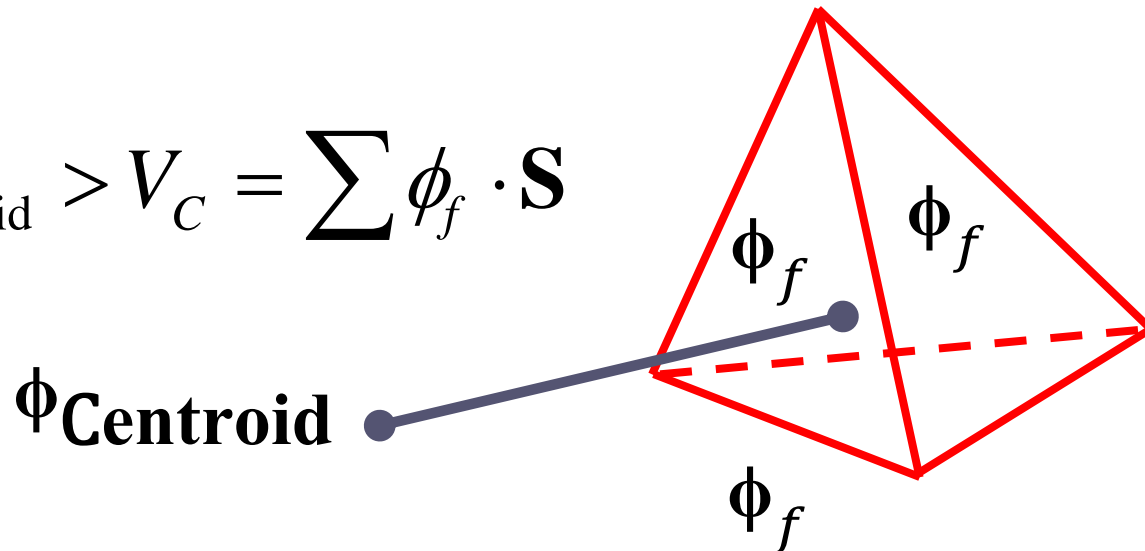
$$\langle \nabla \phi \rangle = \frac{1}{V_C} \int_{V_C} \nabla \phi dV$$

Using the Gauss integration theory (高斯定理), the volume integral (体积分) is transformed into a surface integral (面积分):

$$\langle \nabla \phi \rangle = \frac{1}{V_C} \int_{V_C} \nabla \phi dV = \frac{1}{V_C} \oint \phi \cdot \mathbf{n} dS$$

In the presence of discrete faces, the above equation can be written as:

$$\langle \nabla \phi_{\text{centroid}} \rangle V_C = \sum \phi_f \cdot \mathbf{S}$$



$$\nabla \phi_{\text{centroid}} V_C = \sum \phi_f \cdot \mathbf{n} S$$

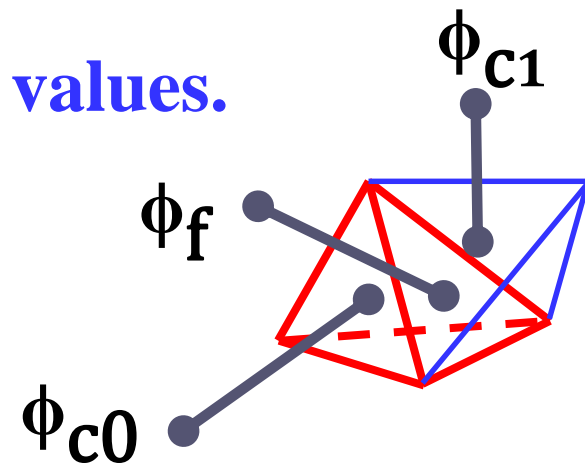
The problem of calculating gradient is transferred into the following equation:

How to determine ϕ_f at the face?

1. Green-Gauss Cell-Based (格林-高斯基于单元法)

Calculate ϕ_f using cell centroid values.

$$\phi_f = \frac{\phi_{C0} + \phi_{C1}}{2}$$



2. Green-Gauss Node-Based (格林-高斯基于节点法)

Calculate ϕ_f by the average of the node values. (代数平均值)

$$\phi_f = \frac{1}{N_f} \sum \phi_n$$

$$\phi_n = \sum_i^{N_{\text{cells}}(n)} \phi_{c_i} w_{c_i, n}$$

N_f : number of nodes on the face, ϕ_n : node value.

ϕ_n , is calculated by weighted average of the cell values surrounding the nodes ϕ_{c_i} .

Review: the node-based method is more accurate than the cell-based method.

3. Least-Squares Cell Based 基于单元体的最小二乘法

It is the default scheme for gradient calculation.

The basic idea is as follows. Consider two cell centroid C_0 and C_i , and their distance vector as δr . Then, the following equation

$$\phi_{C_i} = \phi_{C_0} + (\nabla \phi) \cdot (\mathbf{r}_{C_i} - \mathbf{r}_{C_0})$$

is exact only when the solution field is linear! **In other words, there is no second-order term for Taylor expansion of ϕ !**

For a cell centroid \mathbf{C}_0 with N neighboring nodes \mathbf{C}_i ,

$$\Phi_{C_i} = \phi_{C_i} - \left[\phi_{C_0} + (\nabla \phi) \cdot (\mathbf{r}_{C_i} - \mathbf{r}_{C_0}) \right]$$

True value

Calculated value

Making summation of all these Φ_{C_i} with a weighting factor w_i

$$\begin{aligned} \xi &= \sum_{i=1}^N w_i \Phi_{C_i} = \sum_{i=1}^N \left\{ w_i \left(\phi_{C_i} - \left[\phi_{C_0} + (\nabla \phi) \cdot (\mathbf{r}_{C_i} - \mathbf{r}_{C_0}) \right] \right)^2 \right\} \\ &= \sum_{i=1}^N \left\{ w_i \left(\phi_{C_i} - \phi_{C_0} - \left[\frac{\partial \phi}{\partial x} \Delta x_i + \frac{\partial \phi}{\partial y} \Delta y_i + \frac{\partial \phi}{\partial z} \Delta z_i \right] \right)^2 \right\} \end{aligned}$$

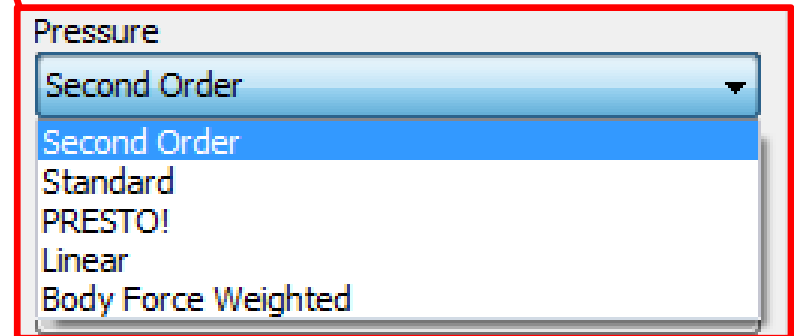
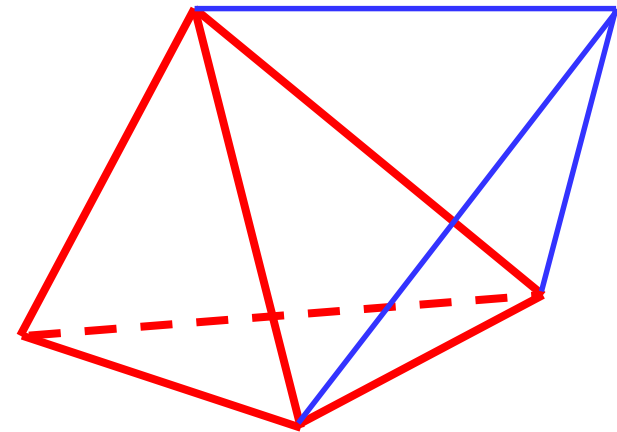
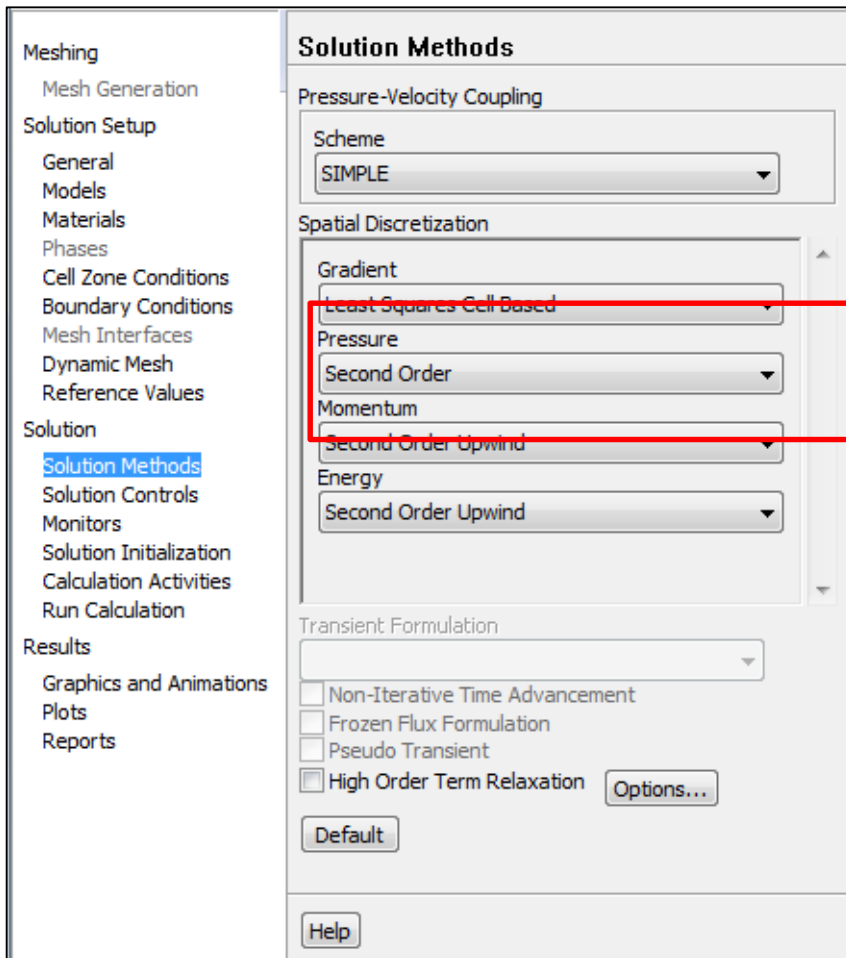
Therefore, to calculate the gradient $\nabla\phi$ it to find the one leading to the minimum ξ !

$$\xi = \sum_{i=1}^N \left\{ w_i \left(\phi_{Ci} - \phi_{C0} - \left[\frac{\partial\phi}{\partial x} \Delta x_i + \frac{\partial\phi}{\partial y} \Delta y_i + \frac{\partial\phi}{\partial z} \Delta z_i \right] \right)^2 \right\}$$

This is the idea of **Least-Squares method**.

Remark: On irregular (不规则) unstructured meshes, the accuracy of the least-squares gradient method is comparable to that of the node-based gradient. However, it is less expensive to compute the least-squares gradient than the node-based gradient.

Pressure calculation: to calculate the pressure value at the interface using centroid value.



1. Linear scheme

Computes the face pressure use the average of the pressure values in the adjacent cells.

$$P_f = \frac{P_{C0} + P_{C1}}{2}$$

2. Standard scheme

Interpolate the pressure using momentum equation coefficient.

$$P_f = \frac{\frac{P_{c0}}{a_{p,c0}} + \frac{P_{c1}}{a_{p,c1}}}{\frac{1}{a_{p,c0}} + \frac{1}{a_{p,c1}}}$$

3. Second Order

Calculate the pressure value using a central difference scheme

$$P_f \approx \frac{P_{C0} + \nabla P_{C0} \mathbf{r}_{C0} + P_{C1} + \nabla P_{C1} \mathbf{r}_{C1}}{2}$$

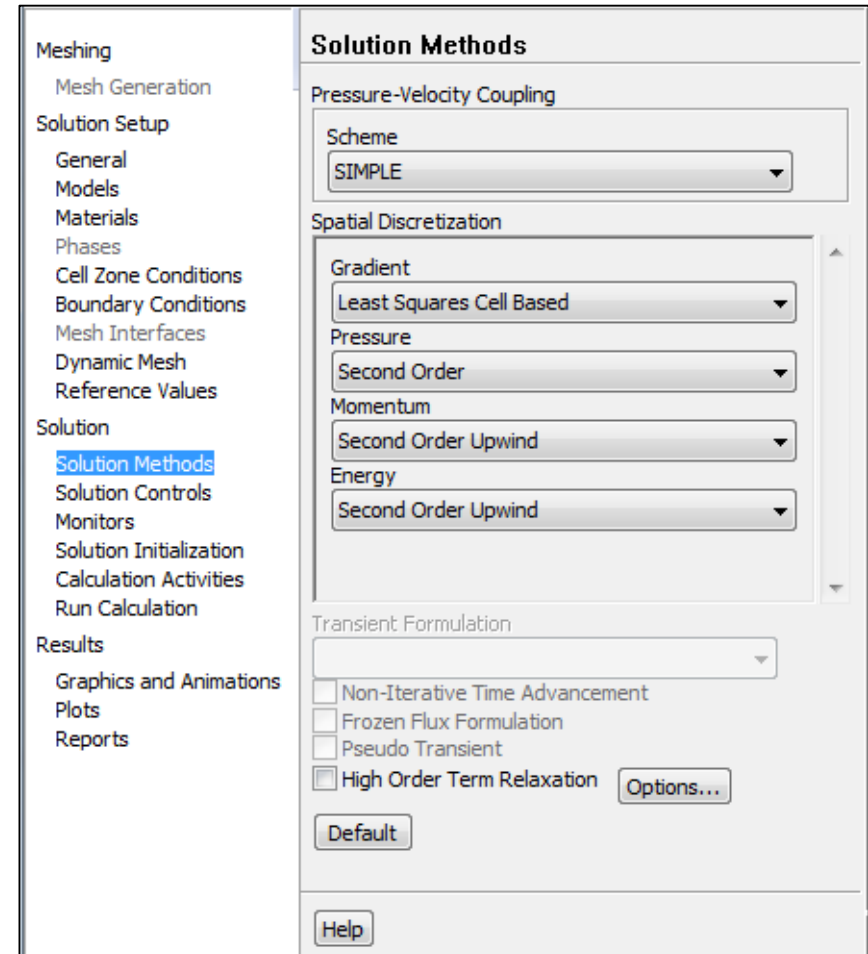
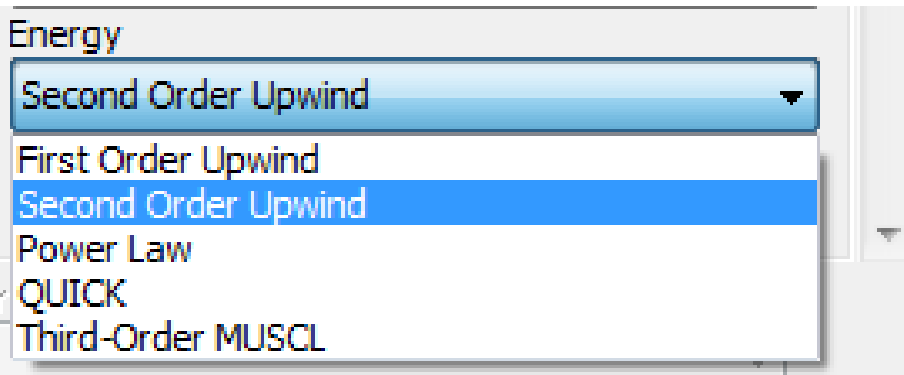
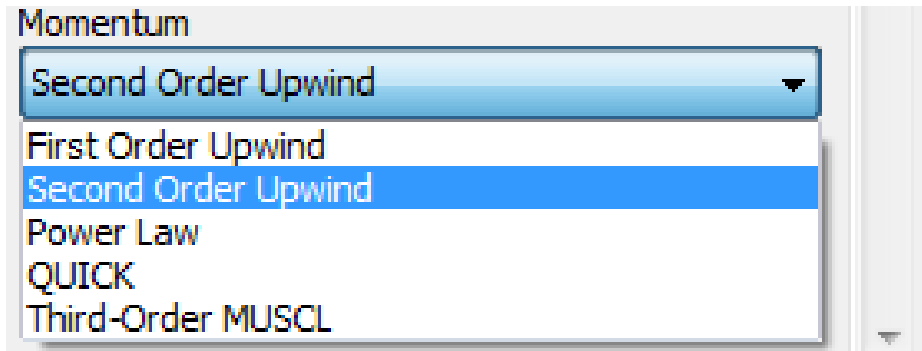
4. Body Force Weighted scheme

Calculate the pressure according to the body force. For multiphase flow such as VOF (Volume of Fluid, 体积函数法) or LS (Level Set, 水平集), it is recommended. Not for porous media!

5. PRESTO! (Pressure Staggering Option) scheme

For problem with high pressure gradient.

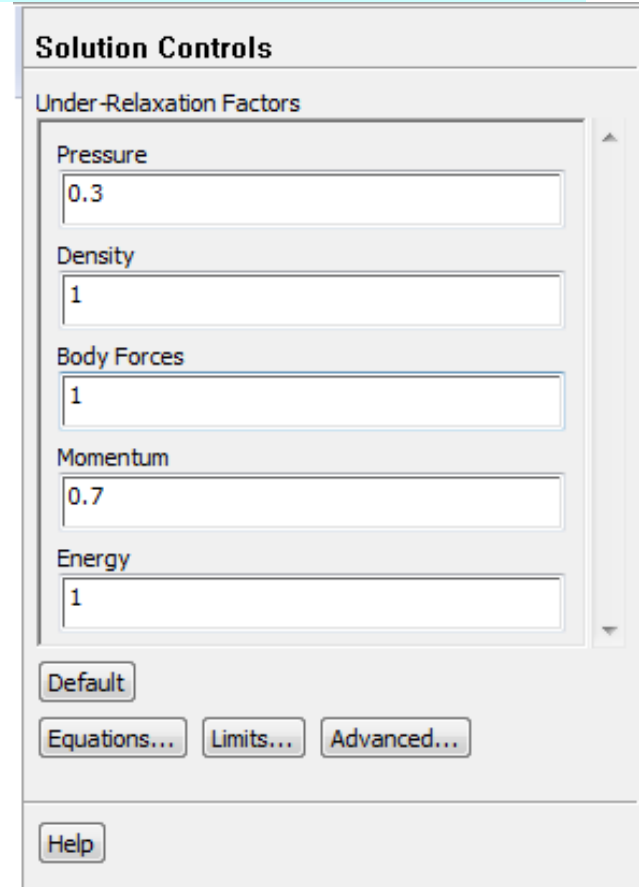
For convective term scheme, we are very familiar!



Step 7: Solution setup: relaxation

Under-relaxation is adopted to control the change rate of simulated variables in subsequent iterations.

The relaxation factor α for each variable has been optimized for the largest possible.



Solution Controls

Under-Relaxation Factors

Pressure	0.3
Density	1
Body Forces	1
Momentum	0.7
Energy	1

Default Equations... Limits... Advanced...

Help

In some cases, if your simulation is not converged, and you are sure there is no problem with other setting, you can try to reduce α !

Remark: In our teaching code, except the pressure correction equation, under-relaxation of other equations are implemented into the solution process.

$$\phi = \phi_P^0 + \alpha \left[\frac{\sum a_{nb} \phi_{nb} + b}{a_P} - \phi_P^0 \right]$$

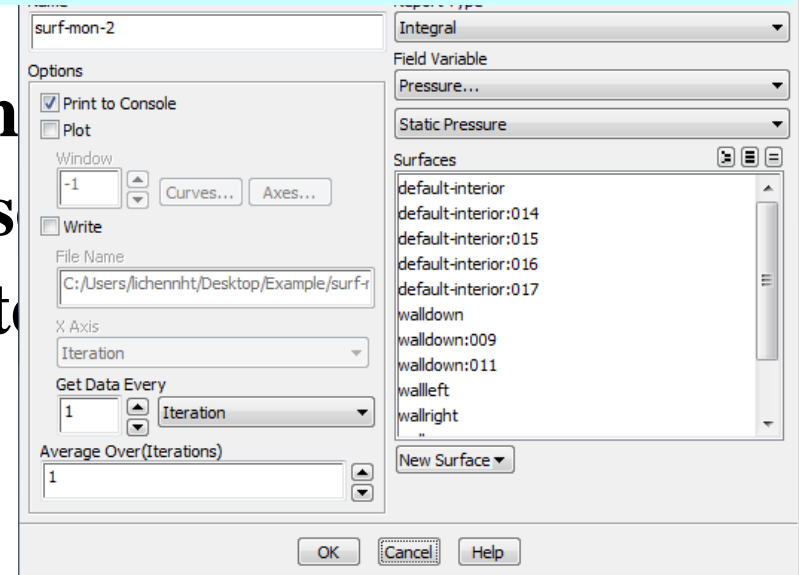
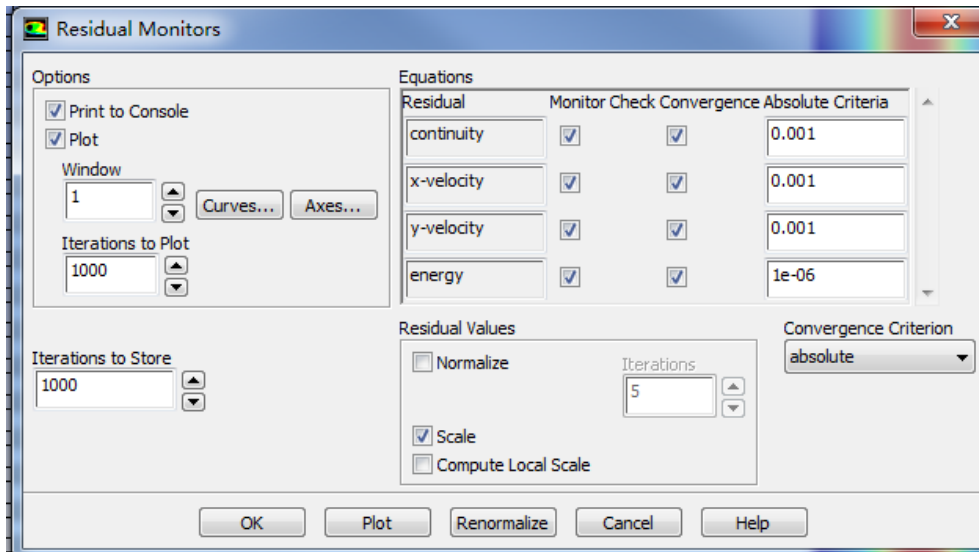
$$\left(\frac{a_P}{\alpha} \right) \phi_P = \sum a_{nb} \phi_{nb} + b + (1 - \alpha) \frac{a_P}{\alpha} \phi_P^0$$

New a_P, a_P'

New $b,$

$$a_P' \phi_P = \sum a_{nb} \phi_{nb} + b'$$

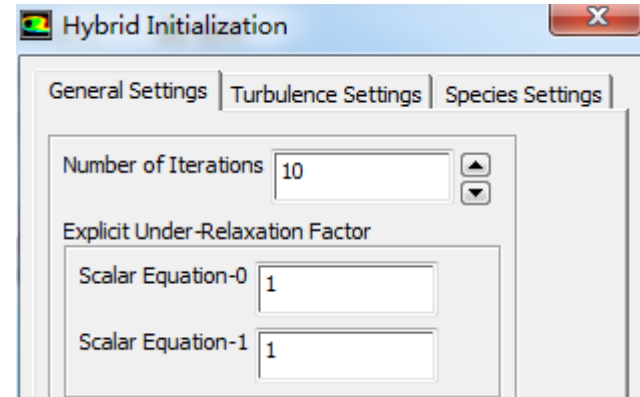
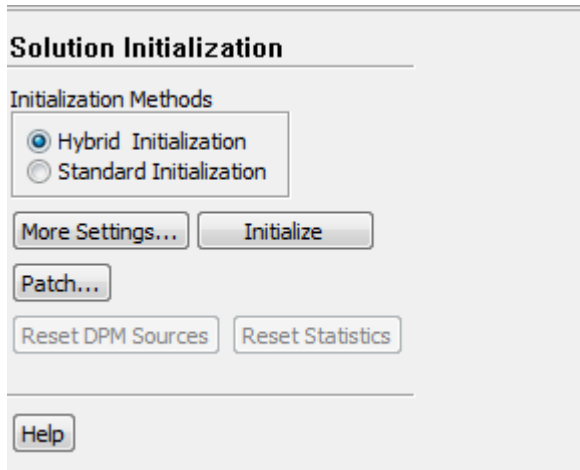
Step 7: Solution setup: monitors



The Residuals are the most important values to be monitored. You can double click it to set the related values.

You can also set other variables to monitor, such as temperature at a certain point.

Step 8: Initialization

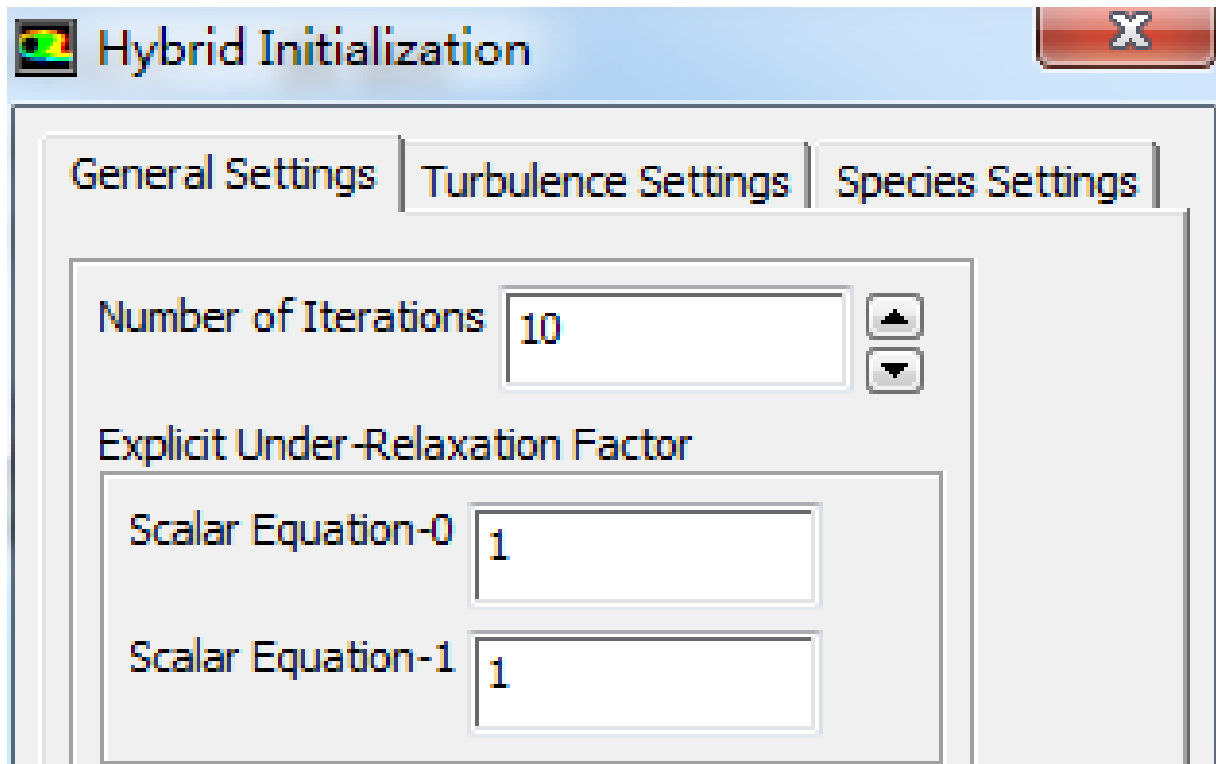


The default selection is Hybrid initialization (混合初始化).

The initial pressure and velocity field you give usually are not consistent, in other words, not meet the NS equation.

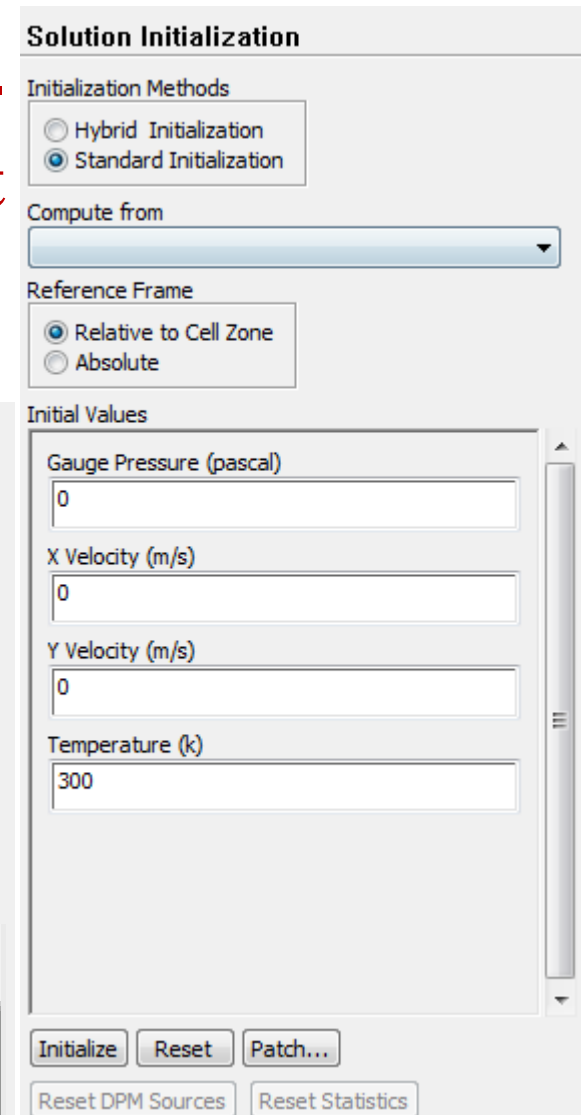
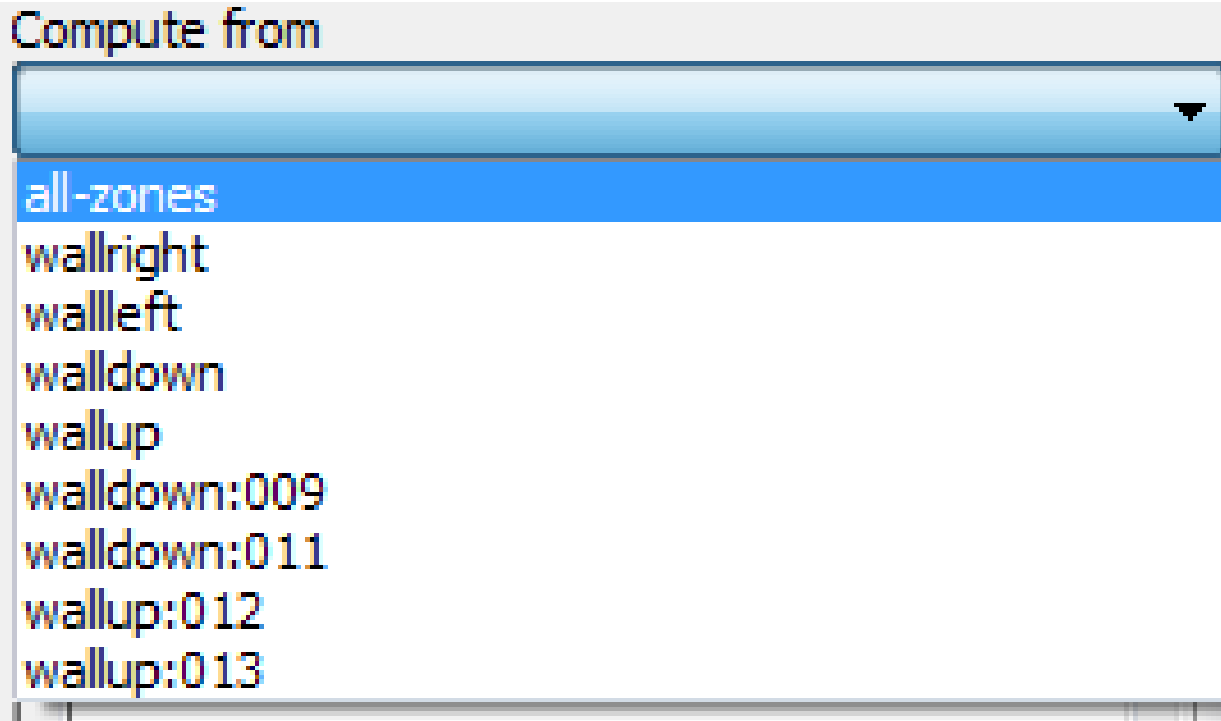
In SIMPLER algorithm, we solved an additional Poisson equation for pressure based on given velocity.

The Hybrid initialization method is similar that Poisson equation is solved to initialize the velocity and pressure equation. You can set the number of iterations to make sure the initial velocity and pressure are consistent.



Or you can simply chose Standard initialization method.

Click Compute from and the drop-down list will show and you can select an region.

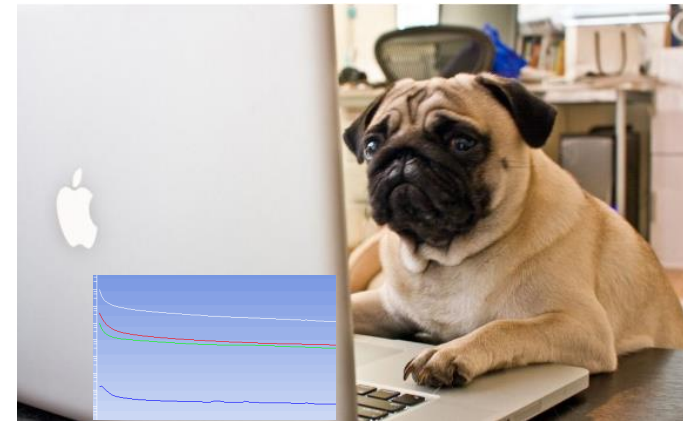


The eight steps for preparing a Fluent simulation have been completed!

1. Read mesh
2. scale domain
3. Choose model
4. define material
5. define zone condition
6. define boundary condition
7. Solution step
8. Initialization
9. Run the simulation.
10. Post-process

Step 9: Run the simulation

**What should you do in this step?
Just stare at the monitor to hope
that the residual curves are going
down for a steady problem.**



Diverged? Go back to Steps 1 to 8.

Step 10: Result post-processing

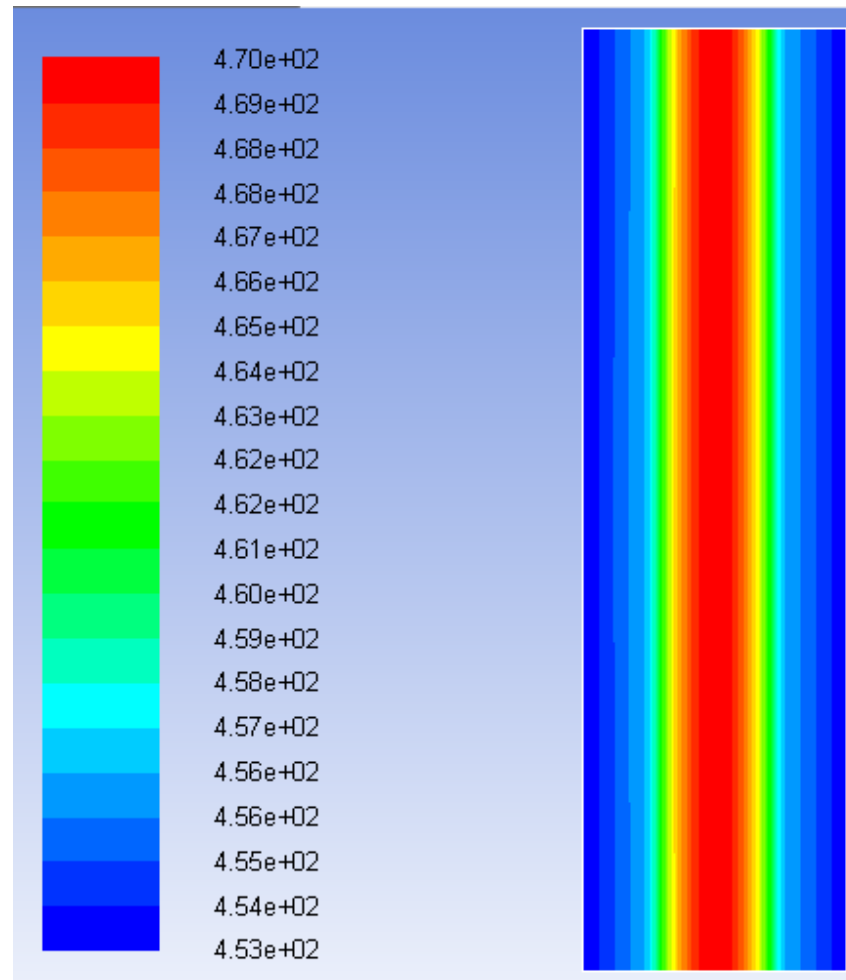


Fig.2 Temperature distribution

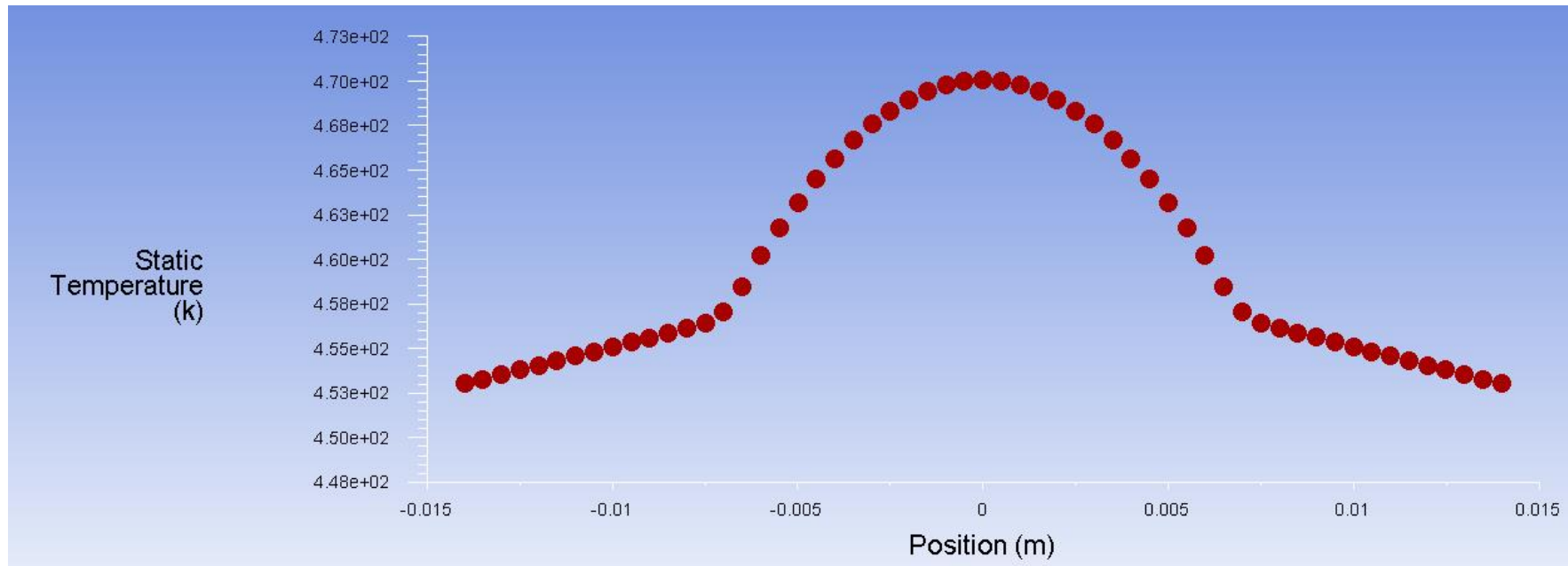


Fig.3 Temperature along x direction at half height

Review: The 10 steps for a Fluent simulation:

- 1. Read and check the mesh: mesh quality.**
- 2. Scale domain: make sure the domain size is right.**
- 3. Choose model: write down the right governing equation is very important.**
- 4. Define material: the solid and fluid related to your problem.**
- 5. Define zone condition: material of each zone and source term**
- 6. Define boundary condition: very important**
- 7. Solution step: algorithm and scheme. Have a background of NHT.**
- 8. Initialization: initial condition**
- 9. Run the simulation: monitor the residual curves and certain variable.**
- 10. Post-process: analyze the results.**

2: Operating the Fluent software to simulate the example and post-process the results. (运行软件)

Uranium: density: 19090 kg/m³; Cp: 116 J/(kg.K)

Thermal conductivity: 27.4 W/(m.K)

Thanks!

欢迎交流

陈黎，教授，博导

热流科学与工程教育部重点实验室，西安交通大学

邮箱：lichennht08@mail.xjtu.edu.cn

个人主页：<http://lichennht08.gr.xjtu.edu.cn/>

硕博士：每年招收3-4硕博士

博士后岗位：年薪18万，五险一金，子女入托上学，两年后考核副研岗



同舟共济渡彼岸!

**People in the same boat help each other to
cross to the other bank, where....**