

The Open Source CFD Toolbox

Programmer's Guide

Version 1.4 11th April 2007 P-2

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P-4

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P-6

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P-8

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P-7

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Contents

C	opyri	ight Notice	P-2
G	NU I	Free Documentation Licence	P-3
	1. A	APPLICABILITY AND DEFINITIONS	P-3
		VERBATIM COPYING	P-4
		COPYING IN QUANTITY	P-4
	4. N	MODIFICATIONS	P-5
	5. C	COMBINING DOCUMENTS	P-6
	6. C	COLLECTIONS OF DOCUMENTS	P-7
	7. A	AGGREGATION WITH INDEPENDENT WORKS	P-7
	8. T	TRANSLATION	P-7
	9. T	TERMINATION	P-7
	10.	FUTURE REVISIONS OF THIS LICENSE	P-8
Ti	ader	narks	P-9
\mathbf{C}	ontei	nts	P-11
1	Ten	asor mathematics	P-15
	1.1	Coordinate system	P-15
	1.2	Tensors	P-15
		1.2.1 Tensor notation	P-17
	1.3	Algebraic tensor operations	P-18
		1.3.1 The inner product	P-18
		1.3.2 The double inner product of two tensors	P-19
		1.3.3 The triple inner product of two third rank tensors	P-19
		1.3.4 The outer product	P-19
		1.3.5 The cross product of two vectors	P-20
		1.3.6 Other general tensor operations	P-20
		1.3.7 Geometric transformation and the identity tensor	P-21
		1.3.8 Useful tensor identities	P-21
		1.3.9 Operations exclusive to tensors of rank 2	P-22
		1.3.10 Operations exclusive to scalars	P-23
	1.4	OpenFOAM tensor classes	P-23
		1.4.1 Algebraic tensor operations in OpenFOAM	P-24
	1.5	Dimensional units	P-26

2 Discretisation procedures P-27 P-27 P-28 P-28 Laplacian P-28 P-28 P-29 P-29 P-30 P-31 P-32 P-33 P-38 P-38 First time derivative P-39 P-39 P-39 P-40 P-41 P-41 P-41 P-41 P-42 2.5.1 Treatment of temporal discretisation in OpenFOAM P-43 P-43 P-44 P-45 3 Examples of the use of OpenFOAM P-45 P-46

Exercise

Exercise

Contents
P-27

P-47 P-47

P-49 P-49

P-52

P-54 P-54

P-54 P-55

P-58

P-59

P-59

P-59

P-60

P-62

P-63

P-63

Open VFOAM-1.4

P-12

Contents			P-13
3.4	Decon	appression of a tank internally pressurised with water	P-63
	3.4.1	Problem specification	P-64
	3.4.2	Mesh Generation	P-65
	3.4.3	Preparing the Run	P-67
	3.4.4	Running the case	P-68
	3.4.5	Improving the solution by refining the mesh	P-69
3.5	Magne	etohydrodynamic flow of a liquid	P-69
	3.5.1	Problem specification	P-69
	3.5.2	Mesh generation	P-71
	3.5.3	Running the case	P-72
Index			P-75

P-14 Contents

Chapter 1

Tensor mathematics

This Chapter describes tensors and their algebraic operations and how they are represented in mathematical text in this book. It then explains how tensors and tensor algebra are programmed in OpenFOAM.

1.1 Coordinate system

OpenFOAM is primarily designed to solve problems in continuum mechanics, *i.e.* the branch of mechanics concerned with the stresses in solids, liquids and gases and the deformation or flow of these materials. OpenFOAM is therefore based in 3 dimensional space and time and deals with physical entities described by tensors. The coordinate system used by OpenFOAM is the right-handed rectangular Cartesian axes as shown in Figure 1.1. This system of axes is constructed by defining an origin O from which three lines are drawn at right angles to each other, termed the Ox, Oy, Oz axes. A right-handed set of axes is defined such that to an observer looking down the Oz axis (with O nearest them), the arc from a point on the Ox axis to a point on the Oy axis is in a clockwise sense.

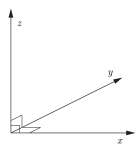


Figure 1.1: Right handed axes

1.2 Tensors

The term tensor describes an entity that belongs to a particular space and obeys certain mathematical rules. Briefly, tensors are represented by a set of *component values* relating

P-16 Tensor mathematics

to a set of unit base vectors; in OpenFOAM the unit base vectors \mathbf{i}_x , \mathbf{i}_y and \mathbf{i}_z are aligned with the right-handed rectangular Cartesian axes x, y and z respectively. The base vectors are therefore orthogonal, i.e. at right-angles to one another. Every tensor has the following attributes:

Dimension d of the particular space to which they belong, i.e. d = 3 in OpenFOAM;

Rank An integer r > 0, such that the number of component values $= d^r$.

While OpenFOAM 1.x is set to 3 dimensions, it offers tensors of ranks 0 to 3 as standard while being written in such a way to allow this basic set of ranks to be extended indefinitely. Tensors of rank 0 and 1, better known as scalars and vectors, should be familiar to readers; tensors of rank 2 and 3 may not be so familiar. For completeness all ranks of tensor offered as standard in OpenFOAM 1.x are reviewed below.

Rank 0 'scalar' Any property which can be represented by a single real number, denoted by characters in italics, e.q. mass m, volume V, pressure p and viscosity μ .

Rank 1 'vector' An entity which can be represented physically by both magnitude and direction. In component form, the vector $\mathbf{a}=(a_1,a_2,a_3)$ relates to a set of Cartesian axes x,y,z respectively. The *index notation* presents the same vector as $a_i, i=1,2,3$, although the list of indices i=1,2,3 will be omitted in this book, as it is intuitive since we are always dealing with 3 dimensions.

Rank 2 'tensor' or second rank tensor, T has 9 components which can be expressed in array notation as:

$$\mathbf{T} = T_{ij} = \begin{pmatrix} T_{11} & T_{12} & T_{13} \\ T_{21} & T_{22} & T_{23} \\ T_{31} & T_{32} & T_{33} \end{pmatrix}$$
 (1.1)

The components T_{ij} are now represented using 2 indices since r=2 and the list of indices i,j=1,2,3 is omitted as before. The components for which i=j are referred to as the diagonal components, and those for which $i\neq j$ are referred to as the off-diagonal components. The transpose of T is produced by exchanging components across the diagonal such that

$$\mathbf{T}^{\mathrm{T}} = T_{ji} = \begin{pmatrix} T_{11} & T_{21} & T_{31} \\ T_{12} & T_{22} & T_{32} \\ T_{13} & T_{23} & T_{33} \end{pmatrix}$$
(1.2)

Note: a rank 2 tensor is often colloquially termed 'tensor' since the occurrence of higher order tensors is fairly rare.

Symmetric rank 2 The term 'symmetric' refers to components being symmetric about the diagonal, i.e. $T_{ij} = T_{ji}$. In this case, there are only 6 independent components since $T_{12} = T_{21}$, $T_{13} = T_{31}$ and $T_{23} = T_{32}$. OpenFOAM distinguishes between symmetric and non-symmetric tensors to save memory by storing 6 components rather than 9 if the tensor is symmetric. Most tensors encountered in continuum mechanics are symmetric.

1.2 Tensors P-17

Rank 3 has 27 components and is represented in index notation as P_{ijk} which is too long to represent in array notation as in Equation 1.1.

Symmetric rank 3 Symmetry of a rank 3 tensor is defined in OpenFOAM to mean that $P_{ijk} = P_{ikj} = P_{jik} = P_{jki} = P_{kij} = P_{kji}$ and therefore has 10 independent components. More specifically, it is formed by the outer product of 3 identical vectors, where the outer product operation is described in Section 1.3.4.

1.2.1 Tensor notation

This is a book on computational continuum mechanics that deals with problems involving complex PDEs in 3 spatial dimensions and in time. It is vital from the beginning to adopt a notation for the equations which is compact yet unambiguous. To make the equations easy to follow, we must use a notation that encapsulates the idea of a tensor as an entity in the own right, rather than a list of scalar components. Additionally, any tensor operation should be perceived as an operation on the entire tensor entity rather than a series of operations on its components.

Consequently, in this book the *tensor notation* is preferred in which any tensor of rank 1 and above, *i.e.* all tensors other than scalars, are represented by letters in bold face, *e.g.* a. This actively promotes the concept of a tensor as a entity in its own right since it is denoted by a single symbol, and it is also extremely compact. The potential drawback is that the rank of a bold face symbol is not immediately apparent, although it is clearly not zero. However, in practice this presents no real problem since we are aware of the property each symbol represents and therefore intuitively know its rank, *e.g.* we know velocity **U** is a tensor of rank 1.

A further, more fundamental idea regarding the choice of notation is that the mathematical representation of a tensor should not change depending on our coordinate system, *i.e.* the vector ais the same vector irrespective of where we view it from. The tensor notation supports this concept as it implies nothing about the coordinate system. However, other notations, *e.g. a_i*, expose the individual components of the tensor which naturally implies the choice of coordinate system. The unsatisfactory consequence of this is that the tensor is then represented by a set of values which are not unique — they depend on the coordinate system.

That said, the index notation, introduced in Section 1.2, is adopted from time to time in this book mainly to expand tensor operations into the constituent components. When using the index notation, we adopt the *summation convention* which states that whenever the same letter subscript occurs twice in a term, the that subscript is to be given all values, *i.e.* 1, 2, 3, and the results added together, *e.q.*

$$a_i b_i = \sum_{i=1}^3 a_i b_i = a_1 b_1 + a_2 b_2 + a_3 b_3 \tag{1.3}$$

In the remainder of the book the symbol \sum is omitted since the repeated subscript indicates the summation.

OpenVFOAM-1.4

P-18 Tensor mathematics

1.3 Algebraic tensor operations

This section describes all the algebraic operations for tensors that are available in Open-FOAM. Let us first review the most simple tensor operations: addition, subtraction, and scalar multiplication and division. Addition and subtraction are both commutative and associative and are only valid between tensors of the same rank. The operations are performed by addition/subtraction of respective components of the tensors, *e.g.* the subtraction of two vectors **a** and **b** is

$$\mathbf{a} - \mathbf{b} = a_i - b_i = (a_1 - b_1, a_2 - b_2, a_3 - b_3) \tag{1.4}$$

Multiplication of any tensor \mathbf{a} by a scalar s is also commutative and associative and is performed by multiplying all the tensor components by the scalar. For example,

$$s\mathbf{a} = sa_i = (sa_1, sa_2, sa_3)$$
 (1.5)

Division between a tensor \mathbf{a} and a scalar is only relevant when the scalar is the second argument of the operation, *i.e.*

$$\mathbf{a}/s = a_i/s = (a_1/s, a_2/s, a_3/s) \tag{1.6}$$

Following these operations are a set of more complex products between tensors of rank 1 and above, described in the following Sections.

1.3.1 The inner product

The inner product operates on any two tensors of rank r_1 and r_2 such that the rank of the result $r=r_1+r_2-2$. Inner product operations with tensors up to rank 3 are described below:

• The inner product of two vectors \mathbf{a} and \mathbf{b} is commutative and produces a scalar $\mathbf{s} = \mathbf{a} \cdot \mathbf{b}$ where

$$s = a_i b_i = a_1 b_1 + a_2 b_2 + a_3 b_3 \tag{1.7}$$

ullet The inner product of a tensor T and vector a produces a vector $b = T \cdot a$, represented below as a column array for convenience

$$b_i = T_{ij}a_j = \begin{pmatrix} T_{11}a_1 + T_{12}a_2 + T_{13}a_3 \\ T_{21}a_1 + T_{22}a_2 + T_{23}a_3 \\ T_{31}a_1 + T_{32}a_2 + T_{33}a_3 \end{pmatrix}$$

$$(1.8)$$

It is non-commutative if **T** is non-symmetric such that $\mathbf{b} = \mathbf{a} \cdot \mathbf{T} = \mathbf{T}^{\mathrm{T}} \cdot \mathbf{a}$ is

$$b_i = a_j T_{ji} = \begin{pmatrix} a_1 T_{11} + a_2 T_{21} + a_3 T_{31} \\ a_1 T_{12} + a_2 T_{22} + a_3 T_{32} \\ a_1 T_{13} + a_2 T_{23} + a_3 T_{33} \end{pmatrix}$$

$$(1.9)$$

• The inner product of two tensors T and S produces a tensor $P = T \cdot S$ whose components are evaluated as:

$$P_{ij} = T_{ik}S_{kj} \tag{1.10}$$

It is non-commutative such that $\mathbf{T} \cdot \mathbf{S} = (\mathbf{S}^{\mathrm{T}} \cdot \mathbf{T}^{\mathrm{T}})^{\mathrm{T}}$

1.3 Algebraic tensor operations P-19

• The inner product of a vector \mathbf{a} and third rank tensor \mathbf{P} produces a second rank tensor $\mathbf{T} = \mathbf{a} \cdot \mathbf{P}$ whose components are

$$T_{ij} = a_k P_{kij} \tag{1.11}$$

Again this is non-commutative so that $T = P \cdot a$ is

$$T_{ij} = P_{ijk}a_k \tag{1.12}$$

• The inner product of a second rank tensor T and third rank tensor P produces a third rank tensor $Q = T \cdot P$ whose components are

$$Q_{ijk} = T_{il}P_{ljk} \tag{1.13}$$

Again this is non-commutative so that $\mathbf{Q} = \mathbf{P} \cdot \mathbf{T}$ is

$$Q_{ijk} = P_{ijl}T_{lk} \tag{1.14}$$

1.3.2 The double inner product of two tensors

The double inner product of two second-rank tensors T and S produces a scalar s = T : S which can be evaluated as the sum of the 9 products of the tensor components

$$s = T_{ij}S_{ij} = T_{11}S_{11} + T_{12}S_{12} + T_{13}S_{13} + T_{21}S_{21} + T_{22}S_{22} + T_{23}S_{23} + T_{31}S_{31} + T_{32}S_{32} + T_{33}S_{33}$$

$$(1.15)$$

The double inner product between a second rank tensor T and third rank tensor P produces a vector a = T : P with components

$$a_i = T_{ik}P_{iki} \tag{1.16}$$

This is non-commutative so that $\mathbf{a} = \mathbf{P} \cdot \mathbf{T}$ is

$$a_i = P_{ijk}T_{jk} \tag{1.17}$$

1.3.3 The triple inner product of two third rank tensors

The triple inner product of two third rank tensors **P** and **Q** produces a scalar $s = \mathbf{P} \cdot \mathbf{Q}$ which can be evaluated as the sum of the 27 products of the tensor components

$$s = P_{ijk}Q_{ijk} \tag{1.18}$$

1.3.4 The outer product

The outer product operates between vectors and tensors as follows:

• The outer product of two vectors \mathbf{a} and \mathbf{b} is non-commutative and produces a tensor $\mathbf{T} = \mathbf{a}\mathbf{b} = (\mathbf{b}\mathbf{a})^T$ whose components are evaluated as:

$$T_{ij} = a_i b_j = \begin{pmatrix} a_1 b_1 & a_1 b_2 & a_1 b_3 \\ a_2 b_1 & a_2 b_2 & a_2 b_3 \\ a_3 b_1 & a_3 b_2 & a_3 b_3 \end{pmatrix}$$
(1.19)

Open VFOAM-1.4

P-20 Tensor mathematics

ullet An outer product of a vector **a** and second rank tensor **T** produces a third rank tensor $\mathbf{P} = \mathbf{a}\mathbf{T}$ whose components are

$$P_{ijk} = a_i T_{jk} \tag{1.20}$$

This is non-commutative so that P = T a produces

$$P_{ijk} = T_{ij}a_k \tag{1.21}$$

1.3.5 The cross product of two vectors

The cross product operation is exclusive to vectors only. For two vectors \mathbf{a} with \mathbf{b} , it produces a vector $\mathbf{c} = \mathbf{a} \times \mathbf{b}$ whose components are

$$c_i = e_{ijk}a_ib_k = (a_2b_3 - a_3b_2, a_3b_1 - a_1b_3, a_1b_2 - a_2b_1)$$

$$(1.22)$$

where the *permutation symbol* is defined by

$$e_{ijk} = \begin{cases} 0 & \text{when any two indices are equal} \\ +1 & \text{when } i,j,k \text{ are an even permutation of } 1,2,3 \\ -1 & \text{when } i,j,k \text{ are an odd permutation of } 1,2,3 \end{cases}$$

$$(1.23)$$

in which the even permutations are 123, 231 and 312 and the odd permutations are 132, 213 and 321.

1.3.6 Other general tensor operations

Some less common tensor operations and terminology used by OpenFOAM are described below.

Square of a tensor is defined as the outer product of the tensor with itself, e.g. for a vector \mathbf{a} , the square $\mathbf{a}^2 = \mathbf{a}\mathbf{a}$.

*n*th power of a tensor is evaluated by *n* outer products of the tensor, *e.g.* for a vector \mathbf{a} , the 3rd power $\mathbf{a}^3 = \mathbf{a}\mathbf{a}\mathbf{a}$.

Magnitude squared of a tensor is the rth inner product of the tensor of rank r with itself, to produce a scalar. For example, for a second rank tensor \mathbf{T} , $|\mathbf{T}|^2 = \mathbf{T} \cdot \mathbf{T}$.

Magnitude is the square root of the magnitude squared, e.g. for a tensor \mathbf{T} , $|\mathbf{T}| = \sqrt{\mathbf{T} \cdot \mathbf{T}}$ Vectors of unit magnitude are referred to as unit vectors.

Component maximum is the component of the tensor with greatest value, inclusive of sign, *i.e.* not the largest magnitude.

Component minimum is the component of the tensor with smallest value.

Component average is the mean of all components of a tensor.

Scale As the name suggests, the scale function is a tool for scaling the components of one tensor by the components of another tensor of the same rank. It is evaluated as the product of corresponding components of 2 tensors, e.g., scaling vector \mathbf{a} by vector \mathbf{b} would produce vector \mathbf{c} whose components are

$$c_i = \text{scale}(\mathbf{a}, \mathbf{b}) = (a_1 b_1, a_2 b_2, a_3 b_3)$$
 (1.24)

1.3 Algebraic tensor operations P-21

1.3.7 Geometric transformation and the identity tensor

A second rank tensor \mathbf{T} is strictly defined as a linear vector function, i.e. it is a function which associates an argument vector \mathbf{a} to another vector \mathbf{b} by the inner product $\mathbf{b} = \mathbf{T} \cdot \mathbf{a}$. The components of \mathbf{T} can be chosen to perform a specific geometric transformation of a tensor from the x, y, z coordinate system to a new coordinate system x^*, y^*, z^* ; \mathbf{T} is then referred to as the *transformation tensor*. While a scalar remains unchanged under a transformation, the vector \mathbf{a} is transformed to \mathbf{a}^* by

$$\mathbf{a}^* = \mathbf{T} \cdot \mathbf{a} \tag{1.25}$$

A second rank tensor S is transformed to S^* according to

$$\mathbf{S}^* = \mathbf{T} \cdot \mathbf{S} \cdot \mathbf{T}^{\mathrm{T}} \tag{1.26}$$

The $identity\ tensor\ \mathbf{I}$ is defined by the requirement that it transforms another tensor onto itself. For all vectors \mathbf{a}

$$\mathbf{a} = \mathbf{I} \cdot \mathbf{a} \tag{1.27}$$

and therefore

$$\mathbf{I} = \delta_{ij} = \begin{pmatrix} 1 & 0 & 0 \\ 0 & 1 & 0 \\ 0 & 0 & 1 \end{pmatrix} \tag{1.28}$$

where δ_{ij} is known as the Kronecker delta symbol.

1.3.8 Useful tensor identities

Several identities are listed below which can be verified by under the assumption that all the relevant derivatives exist and are continuous. The identities are expressed for scalar s and vector \mathbf{a} .

$$\nabla \cdot (\nabla \times \mathbf{a}) \equiv 0$$

$$\nabla \times (\nabla s) \equiv \mathbf{0}$$

$$\nabla \cdot (s\mathbf{a}) \equiv s \nabla \cdot \mathbf{a} + \mathbf{a} \cdot \nabla s$$

$$\nabla \times (s\mathbf{a}) \equiv s \nabla \times \mathbf{a} + \nabla s \times \mathbf{a}$$

$$\nabla (\mathbf{a} \cdot \mathbf{b}) \equiv \mathbf{a} \times (\nabla \times \mathbf{b}) + \mathbf{b} \times (\nabla \times \mathbf{a}) + (\mathbf{a} \cdot \nabla) \mathbf{b} + (\mathbf{b} \cdot \nabla) \mathbf{a}$$

$$\nabla \cdot (\mathbf{a} \times \mathbf{b}) \equiv \mathbf{b} \cdot (\nabla \times \mathbf{a}) - \mathbf{a} \cdot (\nabla \times \mathbf{b})$$

$$\nabla \times (\mathbf{a} \times \mathbf{b}) \equiv \mathbf{a}(\nabla \cdot \mathbf{b}) - \mathbf{b}(\nabla \cdot \mathbf{a}) + (\mathbf{b} \cdot \nabla) \mathbf{a} - (\mathbf{a} \cdot \nabla) \mathbf{b}$$

$$\nabla \times (\nabla \times \mathbf{a}) \equiv \nabla(\nabla \cdot \mathbf{a}) - \nabla^2 \mathbf{a}$$

$$(\nabla \times \mathbf{a}) \times \mathbf{a} \equiv \mathbf{a} \cdot (\nabla \mathbf{a}) - \nabla(\mathbf{a} \cdot \mathbf{a})$$

It is sometimes useful to know the $e-\delta$ identity to help to manipulate equations in index notation:

$$e_{ijk}e_{irs} = \delta_{ir}\delta_{ks} - \delta_{is}\delta_{kr} \tag{1.30}$$

Open VFOAM-1.4

P-22 Tensor mathematics

1.3.9 Operations exclusive to tensors of rank 2

There are several operations that manipulate the components of tensors of rank 2 that are listed below:

Transpose of a tensor $T = T_{ij}$ is $T^T = T_{ii}$ as described in Equation 1.2.

Symmetric and skew (antisymmetric) tensors As discussed in section 1.2, a tensor is said to be symmetric if its components are symmetric about the diagonal, i.e. $\mathbf{T} = \mathbf{T}^{\mathrm{T}}$. A skew or antisymmetric tensor has $\mathbf{T} = -\mathbf{T}^{\mathrm{T}}$ which intuitively implies that $T_{11} = T_{22} = T_{33} = 0$. Every second order tensor can be decomposed into symmetric and skew parts by

$$\mathbf{T} = \underbrace{\frac{1}{2}(\mathbf{T} + \mathbf{T}^{\mathrm{T}})}_{summetric} + \underbrace{\frac{1}{2}(\mathbf{T} - \mathbf{T}^{\mathrm{T}})}_{skew} = \operatorname{symm} \mathbf{T} + \operatorname{skew} \mathbf{T}$$
(1.31)

Trace The trace of a tensor T is a scalar, evaluated by summing the diagonal components

$$\operatorname{tr} \mathbf{T} = T_{11} + T_{22} + T_{33} \tag{1.32}$$

 ${\bf Diagonal}$ returns a vector whose components are the diagonal components of the second rank tensor ${\bf T}$

$$\operatorname{diag} \mathbf{T} = (T_{11}, T_{22}, T_{33}) \tag{1.33}$$

Deviatoric and hydrostatic tensors Every second rank tensor \mathbf{T} can be decomposed into a deviatoric component, for which $\operatorname{tr} \mathbf{T} = 0$ and a hydrostatic component of the form $\mathbf{T} = s\mathbf{I}$ where s is a scalar. Every second rank tensor can be decomposed into deviatoric and hydrostatic parts as follows:

$$\mathbf{T} = \underbrace{\mathbf{T} - \frac{1}{3} (\operatorname{tr} \mathbf{T}) \mathbf{I}}_{deviatoric} + \underbrace{\frac{1}{3} (\operatorname{tr} \mathbf{T}) \mathbf{I}}_{hydrostatic} = \operatorname{dev} \mathbf{T} + \operatorname{hyd} \mathbf{T}$$
(1.34)

Determinant The determinant of a second rank tensor is evaluated by

$$\det \mathbf{T} = \begin{vmatrix} T_{11} & T_{12} & T_{13} \\ T_{21} & T_{22} & T_{23} \\ T_{31} & T_{32} & T_{33} \end{vmatrix} = T_{11}(T_{22}T_{33} - T_{23}T_{32}) - T_{12}(T_{21}T_{33} - T_{23}T_{31}) + T_{13}(T_{21}T_{32} - T_{22}T_{31})$$

$$= \frac{1}{6}e_{ijk}e_{pqr}T_{ip}T_{jq}T_{kr}$$
(1.35)

Cofactors The *minors* of a tensor are evaluated for each component by deleting the row and column in which the component is situated and evaluating the resulting entries as a 2×2 determinant. For example, the minor of T_{12} is

$$\begin{vmatrix} \overline{T_{11}} & \overline{T_{12}} & \overline{T_{13}} \\ T_{21} & T_{22} & T_{23} \\ T_{31} & T_{32} & T_{33} \end{vmatrix} = \begin{vmatrix} T_{21} & T_{23} \\ T_{31} & T_{33} \end{vmatrix} = T_{21}T_{33} - T_{23}T_{31}$$

$$(1.36)$$

1.4 OpenFOAM tensor classes P-23

The cofactors are *signed minors* where each minor is component is given a sign based on the rule

+ve if
$$i + j$$
 is even
-ve if $i + j$ is odd (1.37)

The cofactors of T can be evaluated as

$$\operatorname{cof} \mathbf{T} = \frac{1}{2} e_{jkr} e_{ist} T_{sk} T_{tr} \tag{1.38}$$

Inverse The inverse of a tensor can be evaluated as

$$\operatorname{inv} \mathbf{T} = \frac{\operatorname{cof} \mathbf{T}^{\mathrm{T}}}{\det \mathbf{T}} \tag{1.39}$$

Hodge dual of a tensor is a vector whose components are

$$*\mathbf{T} = (T_{23}, -T_{13}, T_{12}) \tag{1.40}$$

1.3.10 Operations exclusive to scalars

OpenFOAM supports most of the well known functions that operate on scalars, e.g. square root, exponential, logarithm, sine, cosine etc..., a list of which can be found in Table 1.2. There are 3 additional functions defined within OpenFOAM that are described below:

Sign of a scalar s is

$$sgn(s) = \begin{cases} 1 & \text{if } s \ge 0, \\ -1 & \text{if } s < 0. \end{cases}$$
 (1.41)

Positive of a scalar s is

$$pos(s) = \begin{cases} 1 & \text{if } s \ge 0, \\ 0 & \text{if } s < 0. \end{cases}$$
 (1.42)

Limit of a scalar s by the scalar n

$$limit(s, n) = \begin{cases} s & \text{if } s < n, \\ 0 & \text{if } s \ge n. \end{cases}$$
(1.43)

1.4 OpenFOAM tensor classes

OpenFOAM contains a C++ class library primitive that contains the classes for the tensor mathematics described so far. The basic tensor classes that are available as standard in OpenFOAM are listed in Table 1.1. The Table also lists the functions that allow the user to access individual components of a tensor, known as access functions.

We can declare the tensor

$$\mathbf{T} = \begin{pmatrix} 1 & 2 & 3 \\ 4 & 5 & 6 \\ 7 & 8 & 9 \end{pmatrix} \tag{1.44}$$

in OpenFOAM by the line:

OpenVFOAM-1.4

P-24 Tensor mathematics

Rank	Common name	Basic class	Access functions
0	Scalar	scalar	
1	Vector	vector	x(), y(), z()
2	Tensor	tensor	xx(), xy(), xz()

Table 1.1: Basic tensor classes in OpenFOAM

tensor T(1, 2, 3, 4, 5, 6, 7, 8, 9);

We can then access the component T_{13} , or T_{xz} using the xz() access function. For instance the code

outputs to the screen:

Txz = 3

1.4.1 Algebraic tensor operations in OpenFOAM

The algebraic operations described in Section 1.3 are all available to the OpenFOAM tensor classes using syntax which closely mimics the notation used in written mathematics. Some functions are represented solely by descriptive functions, e.g. symm(), but others can also be executed using symbolic operators, e.g. *. All functions are listed in Table 1.2.

Operation	Comment	Mathematical	Description
		Description	in OpenFOAM
Addition		a + b	a + b
Subtraction		a - b	a - b
Scalar multiplication		$s\mathbf{a}$	s * a
Scalar division		\mathbf{a}/s	a / s
Outer product	$rank \mathbf{a}, \mathbf{b} \ge 1$	ab	a * b
Inner product	$rank \mathbf{a}, \mathbf{b} \ge 1$	a•b	a & b
Double inner product	$rank \mathbf{a}, \mathbf{b} >= 2$	a:b	a && b
Cross product	$rank \mathbf{a}, \mathbf{b} = 1$	$\mathbf{a} \times \mathbf{b}$	a ^ b
Square		\mathbf{a}^2	sqr(a)
Magnitude squared		$ \mathbf{a} ^2$	magSqr(a)
Magnitude		a	mag(a)
Power	n = 0, 1,, 4	\mathbf{a}^n	pow(a,n)
Component average	i = 1,, N	$\overline{a_i}$	cmptAv(a)
Component maximum	i = 1,, N	$\max(a_i)$	max(a)
Component minimum	i = 1,, N	$\min(a_i)$	min(a)
Scale		$scale(\mathbf{a}, \mathbf{b})$	scale(a,b)
Geometric transformation	transforms \mathbf{a} u	sing tensor T	transform(T,a)

Operations exclusive to tensors of rank 2

Continued on next page

1.4 OpenFOAM tensor classes P-25

Continued from previous page			
Operation	Comment	Mathematical	Description
		Description	in OpenFOAM
Transpose		\mathbf{T}^{T}	T.T()
Diagonal		$\operatorname{diag} \mathbf{T}$	diag(T)
Trace		$\operatorname{tr} \mathbf{T}$	tr(T)
Deviatoric component		$\operatorname{dev} \mathbf{T}$	dev(T)
Symmetric component		$\operatorname{symm} \mathbf{T}$	symm(T)
Skew-symmetric component		skew T	skew(T)
Determinant		$\det \mathbf{T}$	det(T)
Cofactors		$\operatorname{cof}\mathbf{T}$	cof(T)
Inverse		$\operatorname{inv} \mathbf{T}$	inv(T)
Hodge dual		*T	*T

Operations exclusive to scalars

Sign (boolean)		sgn(s)	sign(s)
Positive (boolean)		s >= 0	pos(s)
Negative (boolean)		$s \le 0$	neg(s)
Limit	n scalar	limit(s, n)	limit(s,n)
Square root		\sqrt{s}	sqrt(s)
Exponential		$\exp s$	exp(s)
Natural logarithm		$\ln s$	log(s)
Base 10 logarithm		$\log_{10} s$	log10(s)
Sine		$\sin s$	sin(s)
Cosine		$\cos s$	cos(s)
Tangent		$\tan s$	tan(s)
Arc sine		a sin s	asin(s)
Arc cosine		$a\cos s$	acos(s)
Arc tangent		a tan s	atan(s)
Hyperbolic sine		$\sinh s$	sinh(s)
Hyperbolic cosine		$\cosh s$	cosh(s)
Hyperbolic tangent		$\tanh s$	tanh(s)
Hyperbolic arc sine		a s inh s	asinh(s)
Hyperbolic arc cosine		$a\cosh s$	acosh(s)
Hyperbolic arc tangent		$\operatorname{atanh} s$	atanh(s)
Error function		$\operatorname{erf} s$	erf(s)
Complement error function		$\operatorname{erfc} s$	erfc(s)
Logarithm gamma function		$\ln \Gamma s$	lgamma(s)
Type 1 Bessel function of order	0	$J_0 s$	j0(s)
Type 1 Bessel function of order	1	$J_1 s$	j1(s)
Type 2 Bessel function of order	0	$Y_0 s$	y0(s)
Type 2 Bessel function of order	1	$Y_1 s$	y1(s)
1 (1 1 1	1 1 11 .	1	

a, b are tensors of arbitrary rank unless otherwise stated

Table 1.2: Algebraic tensor operations in OpenFOAM

OpenVFOAM-1.4

P-26 Tensor mathematics

1.5 Dimensional units

In continuum mechanics, properties are represented in some chosen units, e.g. mass in kilograms (kg), volume in cubic metres (m³), pressure in Pascals (kg m s⁻²). Algebraic operations must be performed on these properties using consistent units of measurement; in particular, addition, subtraction and equality are only physically meaningful for properties of the same dimensional units. As a safeguard against implementing a meaningless operation, OpenFOAM encourages the user to attach dimensional units to any tensor and will then perform dimension checking of any tensor operation.

Units are defined using the dimensionSet class, e.g.

dimensionSet pressureDims(1, -1, -2, 0, 0, 0, 0);

No.	Property	Unit	Symbol
1	Mass	kilogram	k
2	Length	metre	m
3	Time	second	S
4	Temperature	Kelvin	K
5	Quantity	moles	mol
6	Current	ampere	A
7	Luminous intensity	candela	cd

Table 1.3: S.I. base units of measurement

where each of the values corresponds to the power of each of the S.I. base units of measurement listed in Table 1.3. The line of code declares pressureDims to be the dimensionSet for pressure kg m s $^{-2}$ since the first entry in the pressureDims array, 1, corresponds to k^1 , the second entry, -1, corresponds to m^{-1} etc.. A tensor with units is defined using the dimensioned<Type> template class, the <Type> being scalar, vector, tensor, etc.. The dimensioned<Type> stores a variable name of class word,the value <Type> and a dimensionSet

```
dimensionedTensor sigma
  (
     "sigma",
     dimensionSet(1, -1, -2, 0, 0, 0, 0),
     tensor(1e6,0,0,0,1e6,0,0,0,1e6),
  );
```

creates a tensor with correct dimensions of pressure, or stress

$$\sigma = \begin{pmatrix} 10^6 & 0 & 0\\ 0 & 10^6 & 0\\ 0 & 0 & 10^6 \end{pmatrix} \tag{1.45}$$

s is a scalar, N is the number of tensor components

Chapter 2

Discretisation procedures

So far we have dealt with algebra of tensors at a point. The PDEs we wish to solve involve derivatives of tensors with respect to time and space. We therefore need to extend our description to a tensor field, i.e. a tensor that varies across time and spatial domains. In this Chapter we will first present a mathematical description of all the differential operators we may encounter. We will then show how a tensor field is constructed in OpenFOAM and how the derivatives of these fields are discretised into a set of algebraic equations.

2.1 Differential operators

Before defining the spatial derivatives we first introduce the nabla vector operator ∇ , represented in index notation as ∂_i :

$$\nabla \equiv \partial_i \equiv \frac{\partial}{\partial x_i} \equiv \left(\frac{\partial}{\partial x_1}, \frac{\partial}{\partial x_2}, \frac{\partial}{\partial x_3}\right) \tag{2.1}$$

The nabla operator is a useful notation that obeys the following rules:

- it operates on the tensors to its right and the conventional rules of a derivative of a product, e.g. ∂_iab = (∂_ia) b + a (∂_ib);
- otherwise the nabla operator behaves like any other vector in an algebraic operation.

2.1.1 Gradient

If a scalar field s is defined and continuously differentiable then the gradient of s, ∇s is a vector field

$$\nabla s = \partial_i s = \left(\frac{\partial s}{\partial x_1}, \frac{\partial s}{\partial x_2}, \frac{\partial s}{\partial x_3}\right) \tag{2.2}$$

The gradient can operate on any tensor field to produce a tensor field that is one rank higher. For example, the gradient of a vector field \mathbf{a} is a second rank tensor field

$$\nabla \mathbf{a} = \partial_i a_j = \begin{pmatrix} \partial a_1 / \partial x_1 & \partial a_2 / \partial x_1 & \partial a_3 / \partial x_1 \\ \partial a_1 / \partial x_2 & \partial a_2 / \partial x_2 & \partial a_3 / \partial x_2 \\ \partial a_1 / \partial x_3 & \partial a_2 / \partial x_3 & \partial a_3 / \partial x_3 \end{pmatrix}$$

$$(2.3)$$

P-28 Discretisation procedures

2.1.2 Divergence

If a vector field \mathbf{a} is defined and continuously differentiable then the divergence of \mathbf{a} is a scalar field

$$\nabla \cdot \mathbf{a} = \partial_i a_i = \frac{\partial a_1}{\partial x_1} + \frac{\partial a_2}{\partial x_2} + \frac{\partial a_3}{\partial x_3} \tag{2.4}$$

The divergence can operate on any tensor field of rank 1 and above to produce a tensor that is one rank lower. For example the divergence of a second rank tensor field T is a vector field (expanding the vector as a column array for convenience)

$$\nabla \cdot \mathbf{T} = \partial_i T_{ij} = \begin{pmatrix} \partial T_{11}/\partial x_1 + \partial T_{12}/\partial x_1 + \partial T_{13}/\partial x_1 \\ \partial T_{21}/\partial x_2 + \partial T_{22}/\partial x_2 + \partial T_{23}/\partial x_2 \\ \partial T_{31}/\partial x_3 + \partial T_{32}/\partial x_3 + \partial T_{33}/\partial x_3 \end{pmatrix}$$
(2.5)

2.1.3 Curl

If a vector field \mathbf{a} is defined and continuously differentiable then the curl of \mathbf{a} , $\nabla \times \mathbf{a}$ is a vector field

$$\nabla \times \mathbf{a} = e_{ijk} \partial_j a_k = \left(\frac{\partial a_3}{\partial x_2} - \frac{\partial a_2}{\partial x_3}, \frac{\partial a_1}{\partial x_3} - \frac{\partial a_3}{\partial x_1}, \frac{\partial a_2}{\partial x_1} - \frac{\partial a_1}{\partial x_2} \right)$$
(2.6)

The curl is related to the gradient by

$$\nabla \times \mathbf{a} = 2 \,(* \,\mathrm{skew} \,\nabla \mathbf{a}) \tag{2.7}$$

2.1.4 Laplacian

The Laplacian is an operation that can be defined mathematically by a combination of the divergence and gradient operators by $\nabla^2 \equiv \nabla \cdot \nabla$. However, the Laplacian should be considered as a single operation that transforms a tensor field into another tensor field of the same rank, rather than a combination of two operations, one which raises the rank by 1 and one which reduces the rank by 1.

In fact, the Laplacian is best defined as a $scalar\ operator$, just as we defined nabla as a vector operator, by

$$\nabla^2 \equiv \partial^2 \equiv \frac{\partial^2}{\partial x_1^2} + \frac{\partial^2}{\partial x_2^2} + \frac{\partial^2}{\partial x_3^2} \tag{2.8}$$

For example, the Laplacian of a scalar field s is the scalar field

$$\nabla^2 s = \partial^2 s = \frac{\partial^2 s}{\partial x_1^2} + \frac{\partial^2 s}{\partial x_2^2} + \frac{\partial^2 s}{\partial x_3^2} \tag{2.9}$$

2.1.5 Temporal derivative

There is more than one definition of temporal, or time, derivative of a tensor. To describe the temporal derivatives we must first recall that the tensor relates to a property of a volume of material that may be moving. If we track an infinitesimally small volume of material, or

2.2 Overview of discretisation P-29

particle, as it moves and observe the change in the tensorial property ϕ in time, we have the total, or material time derivative denoted by

$$\frac{D\phi}{Dt} = \lim_{\Delta t \to 0} \frac{\Delta\phi}{\Delta t} \tag{2.10}$$

However in continuum mechanics, particularly fluid mechanics, we often observe the change of a ϕ in time at a fixed point in space as different particles move across that point. This change at a point in space is termed the *spatial* time derivative which is denoted by $\partial/\partial t$ and is related to the material derivative by:

$$\frac{D\phi}{Dt} = \frac{\partial\phi}{\partial t} + \mathbf{U} \cdot \nabla\phi \tag{2.11}$$

where **U** is the velocity field of property ϕ . The second term on the right is known as the convective rate of change of ϕ .

2.2 Overview of discretisation

The term discretisation means approximation of a problem into discrete quantities. The FV method and others, such as the finite element and finite difference methods, all discretise the problem as follows:

Spatial discretisation Defining the solution domain by a set of points that fill and bound a region of space when connected;

Temporal discretisation (For transient problems) dividing the time domain into into a finite number of time intervals, or steps;

Equation discretisation Generating a system of algebraic equations in terms of discrete quantities defined at specific locations in the domain, from the PDEs that characterise the problem.

2.2.1 OpenFOAM lists and fields

OpenFOAM frequently needs to store sets of data and perform functions, such as mathematical operations, on the data. OpenFOAM therefore provides an array template class List<Type>, making it possible to create a list of any object of class Type that inherits the functions of the Type. For example a List of vector is List<vector>.

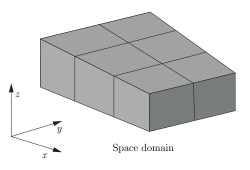
Lists of the tensor classes are defined as standard in OpenFOAM by the template class Field<Type>. For better code legibility, all instances of Field<Type>, e.g. Field<vector>, are renamed using typedef declarations as scalarField, vectorField, tensorField, symmTensorField, tensorThirdField and symmTensorThirdField. Algebraic operations can be performed between Fields subject to obvious restrictions such as the fields having the same number of elements. OpenFOAM also supports operations between a field and single tensor, e.g. all values of a Field U can be multiplied by the scalar 2 with the operation U = 2.0 * U.

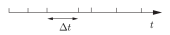
OpenVFOAM-1.4

P-30 Discretisation procedures

2.3 Discretisation of the solution domain

Discretisation of the solution domain is shown in Figure 2.1. The space domain is discretised into computational mesh on which the PDEs are subsequently discretised. Discretisation of time, if required, is simple: it is broken into a set of time steps Δt that may change during a numerical simulation, perhaps depending on some condition calculated during the simulation.





Time domain

Figure 2.1: Discretisation of the solution domain

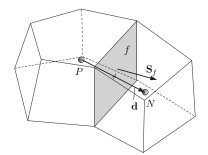


Figure 2.2: Parameters in finite volume discretisation

On a more detailed level, discretisation of space requires the subdivision of the domain into a number of cells, or control volumes. The cells are contiguous, *i.e.* they do not overlap one another and completely fill the domain. A typical cell is shown in Figure 2.2. Dependent variables and other properties are principally stored at the cell centroid P although they

2.3 Discretisation of the solution domain P-31 P-32 Discretisation procedures

may be stored on faces or vertices. The cell is bounded by a set of flat faces, given the generic label f. In OpenFOAM there is no limitation on the number of faces bounding each cell, nor any restriction on the alignment of each face. This kind of mesh is often referred to as "arbitrarily unstructured" to differentiate it from meshes in which the cell faces have a prescribed alignment, typically with the coordinate axes. Codes with arbitrarily unstructured meshes offer greater freedom in mesh generation and manipulation in particular when the geometry of the domain is complex or changes over time.

Whilst most properties are defined at the cell centroids, some are defined at cell faces. There are two types of cell face.

Internal faces Those faces that connect two cells (and it can never be more than two). For each internal face, OpenFOAM designates one adjoining cell to be the face owner and the other to be the neighbour;

Boundary faces Those belonging to one cell since they coincide with the boundary of the domain. These faces simply have an owner cell.

2.3.1 Defining a mesh in OpenFOAM

There are different levels of mesh description in OpenFOAM, beginning with the most basic mesh class, named polyMesh since it is based on polyhedra. A polyMesh is constructed using the minimum information required to define the mesh geometry described below and presented in Figure 2.3:

Points A list of cell vertex point coordinate vectors, *i.e.* a vectorField, that is renamed pointField using a typedef declaration;

Faces A list of cell faces List<face>, or faceList, where the face class is defined by a list of vertex numbers, corresponding to the pointField:

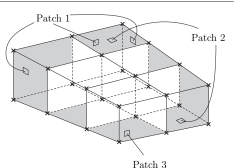
Cells a list of cells List<cell>, or cellList, where the cell class is defined by a list of face numbers, corresponding to the faceList described previously.

Boundary a polyBoundaryMesh decomposed into a list of patches, polyPatchList representing different regions of the boundary. The boundary is subdivided in this manner to allow different boundary conditions to be specified on different patches during a solution. All the faces of any polyPatch are stored as a single block of the faceList, so that its faces can be easily accessed using the slice class which stores references to the first and last face of the block. Each polyPatch is then constructed from

- a slice;
- a word to assign it a name.

FV discretisation uses specific data that is derived from the mesh geometry stored in polyMesh. OpenFOAM therefore extends the polyMesh class to fvMesh which stores the additional data needed for FV discretisation. fvMesh is constructed from polyMesh and stores the data in Table 2.1 which can be updated during runtime in cases where the mesh moves, is refined etc..

OpenVFOAM-1.4



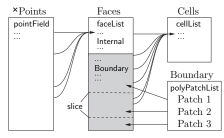


Figure 2.3: Schematic of the basic mesh description used in OpenFOAM

2.3.2 Defining a geometric Field in OpenFOAM

So far we can define a field, i.e. a list of tensors, and a mesh. These can be combined to define a tensor field relating to discrete points in our domain, specified in OpenFOAM by the template class <code>geometricField<Type></code>. The Field values are separated into those defined within the internal region of the domain, e.g. at the cell centres, and those defined on the domain boundary, e.g. on the boundary faces. The <code>geometricField<Type></code> stores the following information:

Internal field This is simply a Field<Type>, described in Section 2.2.1;

BoundaryField This is a GeometricBoundaryField, in which a Field is defined for the faces of each patch and a Field is defined for the patches of the boundary. This is then a field of fields, stored within an object of the FieldField<Type> class. A reference to the fvBoundaryMesh is also stored [**].

Mesh A reference to an fvMesh, with some additional detail as to the whether the field is defined at cell centres, faces, etc..

Dimensions A dimensionSet, described in Section 1.5.

Old values Discretisation of time derivatives requires field data from previous time steps.

2.4 Equation discretisation	P-33	P-34	

Class	Description	Symbol	Access function
volScalarField	Cell volumes	V	V()
surfaceVectorField	Face area vectors	\mathbf{S}_f	Sf()
surfaceScalarField	Face area magnitudes	$ \mathbf{S}_f $	magSf()
volVectorField	Cell centres	C	C()
surfaceVectorField	Face centres	\mathbf{C}_f	Cf()
surfaceScalarField	Face motion fluxes **	ϕ_a	phi()

Table 2.1: fvMesh stored data.

The geometricField<Type> will store references to stored fields from the previous, or old, time step and its previous, or old-old, time step where necessary.

Previous iteration values The iterative solution procedures can use under-relaxation which requires access to data from the previous iteration. Again, if required, geometricField<Type> stores a reference to the data from the previous iteration.

As discussed in Section 2.3, we principally define a property at the cell centres but quite often it is stored at the cell faces and on occasion it is defined on cell vertices. The geometricField<Type> is renamed using typedef declarations to indicate where the field variable is defined as follows:

volField<Type> A field defined at cell centres;

surfaceField<Type> A field defined on cell faces;

pointField<Type> A field defined on cell vertices.

These typedef field classes of geometricField<Type>are illustrated in Figure 2.4. A geometricField<Type> inherits all the tensor algebra of Field<Type> and has all operations subjected to dimension checking using the dimensionSet. It can also be subjected to the FV discretisation procedures described in the following Section. The class structure used to build geometricField<Type> is shown in Figure 2.51.

Equation discretisation

Equation discretisation converts the PDEs into a set of algebraic equations that are commonly expressed in matrix form as:

$$[A][x] = [b] \tag{2.12}$$

where [A] is a square matrix, [x] is the column vector of dependent variable and [b] is the source vector. The description of [x] and [b] as 'vectors' comes from matrix terminology rather than being a precise description of what they truly are: a list of values defined at locations in the geometry, i.e. a geometricField<Type>, or more specifically a volField<Type> when using FV discretisation.

OpenVFOAM-1.4

Discretisation procedures

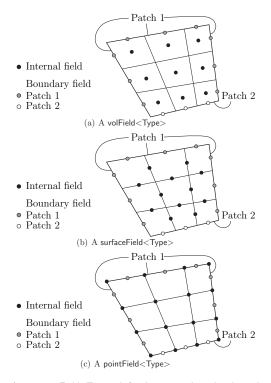


Figure 2.4: Types of geometricField<Type> defined on a mesh with 2 boundary patches (in 2 dimensions for simplicity)

¹The diagram is not an exact description of the class hierarchy, rather a representation of the general structure leading from some primitive classes to geometric<Type>Field.

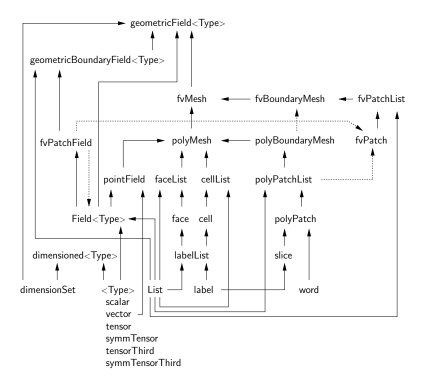


Figure 2.5: Basic class structure leading to geometricField<Type>

OpenVFOAM-1.4

P-36 Discretisation procedures

[A] is a list of coefficients of a set of algebraic equations, and cannot be described as a geometricField<Type>. It is therefore given a class of its own: fvMatrix. fvMatrix<Type> is created through discretisation of a geometric<Type>Field and therefore inherits the <Type>. It supports many of the standard algebraic matrix operations of addition +, subtraction - and multiplication *.

Each term in a PDE is represented individually in OpenFOAM code using the classes of static functions finiteVolumeMethod and finiteVolumeCalculus, abbreviated by a typedef to fvm and fvc respectively. fvm and fvc contain static functions, representing differential operators, e.g. ∇^2 , $\nabla \cdot$ and $\partial/\partial t$, that discretise geometricField<Type>s. The purpose of defining these functions within two classes, fvm and fvc, rather than one, is to distinguish:

- functions of fvm that calculate implicit derivatives of and return an fvMatrix<Type>
- some functions of fvc that calculate explicit derivatives and other explicit calculations, returning a geometricField<Type>.

Figure 2.6 shows a geometric Field<Type> defined on a mesh with 2 boundary patches and illustrates the explicit operations merely transform one field to another and drawn in 2D for simplicity.

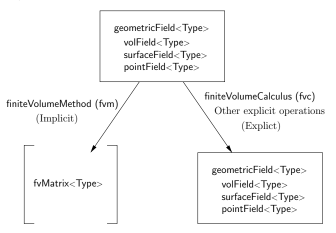


Figure 2.6: A geometricField<Type> and its operators

Table 2.2 lists the main functions that are available in fvm and fvc to discretise terms that may be found in a PDE. FV discretisation of each term is formulated by first integrating the term over a cell volume V. Most spatial derivative terms are then converted to integrals over the cell surface S bounding the volume using Gauss's theorem

$$\int_{V} \nabla \star \phi \ dV = \int_{S} d\mathbf{S} \star \phi \tag{2.13}$$

where **S** is the surface area vector, ϕ can represent any tensor field and the star notation \star is used to represent any tensor product, *i.e.* inner, outer and cross and the respective

Term description	Implicit /	Text	fvm::/fvc:: functions
	Explicit	expression	
Laplacian	Imp/Exp	$\nabla^2 \phi$	laplacian(phi)
		$\nabla \cdot \Gamma \nabla \phi$	laplacian(Gamma, phi)
Time derivative	${\rm Imp}/{\rm Exp}$	$\frac{\partial \phi}{\partial t}$	ddt(phi)
		$\frac{\partial \rho \phi}{\partial t}$	ddt(rho,phi)
Second time derivative	Imp/Exp	$\frac{\partial}{\partial t} \left(\rho \frac{\partial \phi}{\partial t} \right)$	d2dt2(rho, phi)
Convection	Imp/Exp	$\nabla \cdot (\psi)$	$\mathtt{div}(\mathtt{psi},\mathtt{scheme})^*$
		$\nabla \cdot (\psi \phi)$	$\mathtt{div}(\mathtt{psi},\ \mathtt{phi},\ \mathtt{word})^*$
			div(psi, phi)
Divergence	Exp	$\nabla \cdot \chi$	div(chi)
Gradient	Exp	$\nabla \chi$	grad(chi)
		$\nabla \phi$	gGrad(phi)
			lsGrad(phi)
			<pre>snGrad(phi)</pre>
			${\tt snGradCorrection(phi)}$
Grad-grad squared	Exp	$ \nabla\nabla\phi ^2$	sqrGradGrad(phi)
Curl	Exp	$\nabla \times \phi$	curl(phi)
Source	Imp	$\rho\phi$	Sp(rho,phi)
16 9 9	Imp/Exp†	1: :: 1:	SuSp(rho,phi)

†fvm::SuSp source is discretised implicit or explicit depending on the sign of rho. †An explicit source can be introduced simply as a vol<Type>Field, e.g. rho*phi. Function arguments can be of the following classes:

phi: vol<Type>Field

Gamma: scalar volScalarField, surfaceScalarField, volTensorField, surfaceTensorField,

rho: scalar, volScalarField psi: surfaceScalarField.

chi: surface<Type>Field, vol<Type>Field.

Table 2.2: Discretisation of PDE terms in OpenFOAM

Open VFOAM-1.4

P-38 Discretisation procedures

derivatives: divergence $\nabla \cdot \phi$, gradient $\nabla \phi$ and $\nabla \times \phi$. Volume and surface integrals are then linearised using appropriate schemes which are described for each term in the following Sections. Some terms are always discretised using one scheme, a selection of schemes is offered in OpenFOAM for the discretisation of other terms. The choice of scheme is either made by a direct specification within the code or it can be read from an input file at job run-time and stored within an fvSchemes class object.

2.4.1 The Laplacian term

The Laplacian term is integrated over a control volume and linearised as follows:

$$\int_{V} \nabla \cdot (\Gamma \nabla \phi) \ dV = \int_{S} d\mathbf{S} \cdot (\Gamma \nabla \phi) = \sum_{f} \Gamma_{f} \mathbf{S}_{f} \cdot (\nabla \phi)_{f}$$
(2.14)

The face gradient discretisation is implicit when the length vector \mathbf{d} between the centre of the cell of interest P and the centre of a neighbouring cell N is orthogonal to the face plane, *i.e.* parallel to \mathbf{S}_f :

$$\mathbf{S}_{f} \bullet (\nabla \phi)_{f} = |S_{f}| \frac{\phi_{N} - \phi_{P}}{|\mathbf{d}|} \tag{2.15}$$

In the case of non-orthogonal meshes, an additional explicit term is introduced [?] which is evaluated by interpolating cell centre gradients, themselves calculated by central differencing cell centre values.

2.4.2 The convection term

OpenVFOAM-1.4

The convection term is integrated over a control volume and linearised as follows:

$$\int_{V} \nabla \cdot (\rho \mathbf{U} \phi) \ dV = \int_{S} d\mathbf{S} \cdot (\rho \mathbf{U} \phi) = \sum_{f} \mathbf{S}_{f} \cdot (\rho \mathbf{U})_{f} \phi_{f} = \sum_{f} F \phi_{f}$$
(2.16)

The face field ϕ_f can be evaluated using a variety of schemes:

Central differencing (CD) is second-order accurate but unbounded

$$\phi_f = f_x \phi_P + (1 - f_x) \phi_N \tag{2.17}$$

where $f_x \equiv \overline{fN}/\overline{PN}$ where \overline{fN} is the distance between f and cell centre N and \overline{PN} is the distance between cell centres P and N.

Upwind differencing (UD) determines ϕ_f from the direction of flow and is bounded at the expense of accuracy

$$\phi_f = \begin{cases} \phi_P & \text{for } F \ge 0\\ \phi_N & \text{for } F < 0 \end{cases}$$
 (2.18)

Blended differencing (BD) schemes combine UD and CD in an attempt to preserve boundedness with reasonable accuracy,

$$\phi_f = (1 - \gamma) \left(\phi_f\right)_{UD} + \gamma \left(\phi_f\right)_{CD} \tag{2.19}$$

OpenFOAM has several implementations of the Gamma differencing scheme to select the blending coefficient γ [?] but it offers other well-known schemes such as van Leer, SUPERBEE, MINMOD etc.

2.4.3 First time derivative

The first time derivative $\partial/\partial t$ is integrated over a control volume as follows:

$$\frac{\partial}{\partial t} \int_{V} \rho \phi \ dV$$
 (2.20)

The term is discretised by simple differencing in time using:

new values $\phi^n \equiv \phi(t + \Delta t)$ at the time step we are solving for;

old values $\phi^o \equiv \phi(t)$ that were stored from the previous time step;

old-old values $\phi^{oo} \equiv \phi(t - \Delta t)$ stored from a time step previous to the last.

One of two discretisation schemes can be declared using the timeScheme keyword in the appropriate input file, described in detail in section 4.4 of the User Guide.

Euler implicit scheme, timeScheme EulerImplicit, that is first order accurate in time:

$$\frac{\partial}{\partial t} \int_{V} \rho \phi \ dV = \frac{(\rho_{P} \phi_{P} V)^{n} - (\rho_{P} \phi_{P} V)^{o}}{\Delta t}$$
(2.21)

Backward differencing scheme, timeScheme BackwardDifferencing, that is second order accurate in time by storing the old-old values and therefore with a larger overhead in data storage than EulerImplicit:

$$\frac{\partial}{\partial t} \int_{V} \rho \phi \ dV = \frac{3 \left(\rho_{P} \phi_{P} V\right)^{n} - 4 \left(\rho_{P} \phi_{P} V\right)^{o} + \left(\rho_{P} \phi_{P} V\right)^{oo}}{2 \Delta t} \tag{2.22}$$

2.4.4 Second time derivative

The second time derivative is integrated over a control volume and linearised as follows:

$$\frac{\partial}{\partial t} \int_{V} \rho \frac{\partial \phi}{\partial t} dV = \frac{(\rho_{P} \phi_{P} V)^{n} - 2(\rho_{P} \phi_{P} V)^{o} + (\rho_{P} \phi_{P} V)^{oo}}{\Delta t^{2}}$$
(2.23)

It is first order accurate in time.

2.4.5 Divergence

The divergence term described in this Section is strictly an explicit term that is distinguished from the convection term of Section 2.4.2, *i.e.* in that it is not the divergence of the product of a velocity and dependent variable. The term is integrated over a control volume and linearised as follows:

$$\int_{V} \nabla \cdot \phi \ dV = \int_{S} d\mathbf{S} \cdot \phi = \sum_{f} \mathbf{S}_{f} \cdot \phi_{f}$$
(2.24)

The fvc::div function can take as its argument either a surface<Type>Field, in which case ϕ_f is specified directly, or a vol<Type>Field which is interpolated to the face by central differencing as described in Section 2.4.10:

OpenVFOAM-1.4

P-40 Discretisation procedures

2.4.6 Gradient

The gradient term is an explicit term that can be evaluated in a variety of ways. The scheme can be evaluated either by selecting the particular grad function relevant to the discretisation scheme, e.g. fvc::gGrad, fvc::lsGrad etc., or by using the fvc::grad function combined with the appropriate timeScheme keyword in an input file

Gauss integration is invoked using the fvc::grad function with timeScheme Gauss or directly using the fvc::gGrad function. The discretisation is performed using the standard method of applying Gauss's theorem to the volume integral:

$$\int_{V} \nabla \phi \ dV = \int_{S} d\mathbf{S} \ \phi = \sum_{f} \mathbf{S}_{f} \phi_{f} \tag{2.25}$$

As with the fvc::div function, the Gaussian integration fvc::grad function can take either a surfaceField<Type> or a volField<Type> as an argument.

Least squares method is based on the following idea:

- a value at point P can be extrapolated to neighbouring point N using the gradient at P;
- 2. the extrapolated value at N can be compared to the actual value at N, the difference being the error:
- if we now minimise the sum of the square of weighted errors at all neighbours of P with the respect to the gradient, then the gradient should be a good approximation.

Least squares is invoked using the fvc::grad function with timeScheme leastSquares or directly using the fvc::lsGrad function. The discretisation is performed as by first calculating the tensor G at every point P by summing over neighbours N:

$$\mathbf{G} = \sum_{N} w_{N}^{2} \mathbf{d} \mathbf{d} \tag{2.26}$$

where **d** is the vector from P to N and the weighting function $w_N = 1/|\mathbf{d}|$. The gradient is then evaluated as:

$$(\nabla \phi)_P = \sum_N w_N^2 \mathbf{G}^{-1} \cdot \mathbf{d} (\phi_N - \phi_P)$$
(2.27)

Surface normal gradient The gradient normal to a surface $\mathbf{n}_f \cdot (\nabla \phi)_f$ can be evaluated at cell faces using the scheme

$$(\nabla \phi)_f = \frac{\phi_N - \phi_P}{|\mathbf{d}|} \tag{2.28}$$

This gradient is called by the function fvc::snGrad and returns a surfaceField<Type>. The scheme is directly analogous to that evaluated for the Laplacian discretisation scheme in Section 2.4.1, and in the same manner, a correction can be introduced to improve the accuracy of this face gradient in the case of non-orthogonal meshes. This correction is called using the function fvc::snGradCorrection [Check**].

2.4.7 Grad-grad squared

The grad-grad squared term is evaluated by: taking the gradient of the field; taking the gradient of the resulting gradient field; and then calculating the magnitude squared of the result. The mathematical expression for grad-grad squared of ϕ is $|\nabla (\nabla \phi)|^2$.

2.4.8 Curl

The curl is evaluated from the gradient term described in Section 2.4.6. First, the gradient is discretised and then the curl is evaluated using the relationship from Equation 2.7, repeated here for convenience

$$\nabla \times \phi = 2 * (\text{skew } \nabla \phi)$$

2.4.9 Source terms

Source terms can be specified in 3 ways

Explicit Every explicit term is a volField<Type>. Hence, an explicit source term can be incorporated into an equation simply as a field of values. For example if we wished to solve Poisson's equation $\nabla^2 \phi = f$, we would define phi and f as volScalarField and then do

Implicit An implicit source term is integrated over a control volume and linearised by

$$\int_{V} \rho \phi \ dV = \rho_P V_P \phi_P \tag{2.29}$$

Implicit/Explicit The implicit source term changes the coefficient of the diagonal of the matrix. Depending on the sign of the coefficient and matrix terms, this will either increase or decrease diagonal dominance of the matrix. Decreasing the diagonal dominance could cause instability during iterative solution of the matrix equation. Therefore OpenFOAM provides a mixed source discretisation procedure that is implicit when the coefficients that are greater than zero, and explicit for the coefficients less than zero. In mathematical terms the matrix coefficient for node P is $V_P \max(\rho_P, 0)$ and the source term is $V_P \phi_P \min(\rho_P, 0)$.

2.4.10 Other explicit discretisation schemes

There are some other discretisation procedures that convert volField < Type > s into surface < Type > Fields and visa versa.

Surface integral fvc::surfaceIntegrate performs a summation of surface
-Type>Field face values bounding each cell and dividing by the cell volume, i.e.
 $(\sum_f \phi_f)/V_P$. It returns a volField
-Type>.

Surface sum fvc::surfaceSum performs a summation of surface
-Type>Field face values bounding each cell, i.e.
 $\sum_f \phi_f$ returning a volField
-Type>.

OpenVFOAM-1.4

P-42 Discretisation procedures

Average fvc::average produces an area weighted average of surface<Type>Field face values, i.e. $(\sum_f S_f \phi_f)/\sum_f S_f$, and returns a volField<Type>.

Reconstruct

Face interpolate The geometric<Type>Field function faceInterpolate() interpolates volField<Type> cell centre values to cell faces using central differencing, returning a surface<Type>Field.

2.5 Temporal discretisation

Although we have described the discretisation of temporal derivatives in Sections 2.4.3 and 2.4.4, we need to consider how to treat the spatial derivatives in a transient problem. If we denote all the spatial terms as $\mathcal{A}\phi$ where \mathcal{A} is any spatial operator, e.g. Laplacian, then we can express a transient PDE in integral form as

$$\int_{t}^{t+\Delta t} \left[\frac{\partial}{\partial t} \int_{V} \rho \phi \ dV + \int_{V} \mathcal{A} \phi \ dV \right] \ dt = 0$$
(2.30)

Using the Euler implicit method of Equation 2.21, the first term can be expressed as

$$\int_{t}^{t+\Delta t} \left[\frac{\partial}{\partial t} \int_{V} \rho \phi \ dV \right] \ dt = \int_{t}^{t+\Delta t} \frac{(\rho_{P} \phi_{P} V)^{n} - (\rho_{P} \phi_{P} V)^{o}}{\Delta t} \ dt$$

$$= \frac{(\rho_{P} \phi_{P} V)^{n} - (\rho_{P} \phi_{P} V)^{o}}{\Delta t} \Delta t$$
(2.31)

The second term can be expressed as

$$\int_{t}^{t+\Delta t} \left[\int_{V} \mathcal{A}\phi \ dV \right] \ dt = \int_{t}^{t+\Delta t} \mathcal{A}^{*}\phi \ dt \tag{2.32}$$

where \mathcal{A}^* represents the spatial discretisation of \mathcal{A} . The time integral can be discretised in three ways:

Euler implicit uses implicit discretisation of the spatial terms, thereby taking current values ϕ^n .

$$\int_{t}^{t+\Delta t} A^* \phi \ dt = A^* \phi^n \Delta t \tag{2.33}$$

It is first order accurate in time, guarantees boundedness and is unconditionally stable.

Explicit uses explicit discretisation of the spatial terms, thereby taking old values ϕ^o .

$$\int_{t}^{t+\Delta t} \mathcal{A}^{*} \phi \ dt = \mathcal{A}^{*} \phi^{o} \Delta t \tag{2.34}$$

It is first order accurate in time and is unstable if the Courant number Co is greater than 1. The Courant number is defined as

$$Co = \frac{\mathbf{U}_f \cdot \mathbf{d}}{|\mathbf{d}|^2 \Delta t} \tag{2.35}$$

where \mathbf{U}_f is a characteristic velocity, e.g. velocity of a wave front, velocity of flow.

2.6 Boundary Conditions P-43

Crank Nicholson uses the trapezoid rule to discretise the spatial terms, thereby taking a mean of current values ϕ^n and old values ϕ^o .

$$\int_{t}^{t+\Delta t} \mathcal{A}^{*} \phi \ dt = \mathcal{A}^{*} \left(\frac{\phi^{n} + \phi^{o}}{2} \right) \Delta t \tag{2.36}$$

It is second order accurate in time, is unconditionally stable but does not guarantee boundedness.

2.5.1 Treatment of temporal discretisation in OpenFOAM

At present the treatment of the temporal discretisation is controlled by the implementation of the spatial derivatives in the PDE we wish to solve. For example, let us say we wish to solve a transient diffusion equation

$$\frac{\partial \phi}{\partial t} = \kappa \nabla^2 \phi \tag{2.37}$$

An Euler implicit implementation of this would read

```
solve(fvm::ddt(phi) == kappa*fvm::laplacian(phi))
```

where we use the fvm class to discretise the Laplacian term implicitly. An explicit implementation would read

```
solve(fvm::ddt(phi) == kappa*fvc::laplacian(phi))
```

where we now use the fvc class to discretise the Laplacian term explicitly. The Crank Nicholson scheme can be implemented by the mean of implicit and explicit terms:

```
solve
  (
  fvm::ddt(phi)
  ==
  kappa*0.5*(fvm::laplacian(phi) + fvc::laplacian(phi))
  )
```

2.6 Boundary Conditions

Boundary conditions are required to complete the problem we wish to solve. We therefore need to specify boundary conditions on all our boundary faces. Boundary conditions can be divided into 2 types:

Dirichlet prescribes the value of the dependent variable on the boundary and is therefore termed 'fixed value' in this guide;

Neumann prescribes the gradient of the variable normal to the boundary and is therefore termed 'fixed gradient' in this guide.

OpenVFOAM-1.4

P-44 Discretisation procedures

When we perform discretisation of terms that include the sum over faces \sum_f , we need to consider what happens when one of the faces is a boundary face.

Fixed value We specify a fixed value at the boundary ϕ_b

- We can simply substitute φ_b in cases where the discretisation requires the value on a boundary face φ_f, e.g. in the convection term in Equation 2.16.
- In terms where the face gradient $(\nabla \phi)_f$ is required, e.g. Laplacian, it is calculated using the boundary face value and cell centre value,

$$\mathbf{S}_{f} \cdot (\nabla \phi)_{f} = |S_{f}| \frac{\phi_{b} - \phi_{P}}{|\mathbf{d}|} \tag{2.38}$$

Fixed gradient The fixed gradient boundary condition g_b is a specification on inner product of the gradient and unit normal to the boundary, or

$$g_b = \left(\frac{\mathbf{S}}{|\mathbf{S}|} \bullet \nabla \phi\right)_f \tag{2.39}$$

• When discretisation requires the value on a boundary face ϕ_f we must interpolate the cell centre value to the boundary by

$$\phi_f = \phi_P + \mathbf{d} \cdot (\nabla \phi)_f$$

$$= \phi_P + |\mathbf{d}| q_h$$
(2.40)

• ϕ_b can be directly substituted in cases where the discretisation requires the face gradient to be evaluated,

$$\mathbf{S}_f \bullet (\nabla \phi)_f = |S_f| \, g_b \tag{2.41}$$

2.6.1 Physical boundary conditions

The specification of boundary conditions is usually an engineer's interpretation of the true behaviour. Real boundary conditions are generally defined by some physical attributes rather than the numerical description as described of the previous Section. In incompressible fluid flow there are the following physical boundaries

Inlet The velocity field at the inlet is supplied and, for consistency, the boundary condition on pressure is zero gradient.

Outlet The pressure field at the outlet is supplied and a zero gradient boundary condition on velocity is specified.

No-slip impermeable wall The velocity of the fluid is equal to that of the wall itself, *i.e.* a fixed value condition can be specified. The pressure is specified zero gradient since the flux through the wall is zero.

In a problem whose solution domain and boundary conditions are symmetric about a plane, we only need to model half the domain to one side of the symmetry plane. The boundary condition on the plane must be specified according to

Symmetry plane The symmetry plane condition specifies the component of the gradient normal to the plane should be zero. [Check**]

Chapter 3

Examples of the use of OpenFOAM

In this section we shall describe several test cases supplied with the OpenFOAM distribution. The intention is to provide example cases, including those in the tutorials in chapter 2 of the User Guide, for every standard solver. The examples are designed to introduce certain tools and features of OpenFOAM, e.g. within pre-/post-processing, numerical schemes, algorithms. They also provide a means for validation of solvers although that is not their principal function.

Each example contains a description of the problem: the geometry, initial and boundary conditions, a brief description of the equations being solved, models used, and physical properties required. The solution domain is selected which may be a portion of the original geometry, e.g. if we introduce symmetry planes. The method of meshing, usually blockMesh, is specified; of course the user can simply view the mesh since every example is distributed with the polyMesh directory containing the data files that describe the mesh.

The examples coexist with the tutorials in the *tutorials* subdirectory of the OpenFOAM installation. They are organised into a set of subdirectories by solver, *e.g.* all the <code>icoFoam</code> cases are stored within a subdirectory <code>icoFoam</code>. Before running a particular example, the user is urged to copy it into their user account. We recommend that the user stores all OpenFOAM cases in a directory we recommend that the tutorials are copied into a directory <code>\$FOAM_RUN</code>. If this directory structure has not yet been created in the user's account, it can be created with

```
mkdir -p $FOAM_RUN
```

The tutorials can then be copied into this directory with

cp -r \$FOAM_TUTORIALS/* \$FOAM_RUN

3.1 Flow around a cylinder

In this example we shall investigate potential flow around a cylinder using potentialFoam. This example introduces the following OpenFOAM features:

- non-orthogonal meshes;
- generating an analytical solution to a problem in OpenFOAM.

P-46

Examples of the use of OpenFOAM

3.1.1 Problem specification

The problem is defined as follows:

Solution domain The domain is 2 dimensional and consists of a square domain with a cylinder collocated with the centre of the square as shown in Figure 3.1.

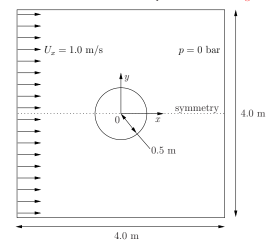


Figure 3.1: Geometry of flow round a cylinder

Governing equations

• Mass continuity for an incompressible fluid

$$\mathbf{V} \cdot \mathbf{U} = 0 \tag{3.1}$$

 Pressure equation for an incompressible, irrotational fluid assuming steady-state conditions

$$\nabla^2 p = 0 \tag{3.2}$$

$Boundary\ conditions$

- Inlet (left) with fixed velocity U = (1,0,0) m/s.
- Outlet (right) with a fixed pressure p = 0 Pa.
- No-slip wall (bottom);
- Symmetry plane (top).

Initial conditions U=0 m/s, p=0 Pa — required in OpenFOAM input files but not necessary for the solution since the problem is steady-state.

Solver name potentialFoam: a potential flow code, *i.e.* assumes the flow is incompressible, steady, irrotational, inviscid and it ignores gravity.

Case name cylinder case located in the \$FOAM_TUTORIALS/potentialFoam directory.

3.1.2 Note on potentialFoam

potential Foam is a useful solver to validate OpenFOAM since the assumptions of potential flow are such that an analytical solution exists for cases whose geometries are relatively simple. In this example of flow around a cylinder an analytical solution exists with which we can compare our numerical solution. potential Foam can also be run more like a utility to provide a (reasonably) conservative initial U field for a problem. When running certain cases, this can useful for avoiding instabilities due to the initial field being unstable. In short, potential Foam creates a conservative field from a non-conservative initial field supplied by the user.

3.1.3 Mesh generation

Mesh generation using blockMesh has been described in tutorials in the User Guide. In this case, the mesh consists of 10 blocks as shown in Figure 3.2. Remember that all meshes

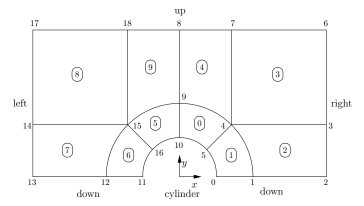


Figure 3.2: Blocks in cylinder geometry

are treated as 3 dimensional in OpenFOAM. If we wish to solve a 2 dimensional problem, we must describe a 3 dimensional mesh that is only one cell thick in the third direction that is not solved. In Figure 3.2 we show only the back plane of the geometry, along z=-0.5, in which the vertex numbers are numbered 0-18. The other 19 vertices in the front plane, z=+0.5, are numbered in the same order as the back plane, as shown in the mesh description file below:

OpenVFOAM-1.4

Examples of the use of OpenFOAM

```
13
14
                                  ""
15
            case
                                  ....
16
17
            instance
                                  dictionary
19
           object
                                  blockMeshDict:
21
22
                                    23
24
25
      convertToMeters 1:
27
      vertices
            (0.5 \ 0 \ -0.5)
29
           (1 0 -0.5)
(2 0 -0.5)
(2 0.707107 -0.5)
(0.707107 0.707107 -0.5)
32
            (0.353553 0.353553
(2 2 -0.5)
            (0.707107 2 -0.5)
            (0 2 -0.5)
(0 1 -0.5)
            (0 0.5 -0.5)
(-0.5 0 -0.5)
            (-1 0 -0.5)
(-2 0 -0.5)
            (-2 0 -0.5)

(-2 0.707107 -0.5)

(-0.707107 0.707107 -0.5)

(-0.353553 0.353553 -0.5)

(-2 2 -0.5)

(-0.707107 2 -0.5)
44
\frac{46}{47}
            (0.5 0 0.5)
(1 0 0.5)
49
            (1 0 0.5)
(2 0 0.5)
(2 0.707107 0.5)
(0.707107 0.707107 0.5)
50
51
52
53
           (0.707107 0.707107 0.5)
(0.353553 0.353553 0.5)
(2 2 0.5)
(0.707107 2 0.5)
(0 2 0.5)
(0 1 0.5)
(0 0.5 0.5)
54
55
56
57
58
59
60
61
            (-0.5 0 0.5)
(-1 0 0.5)
(-2 0 0.5)
            (-2 0.707107 0.5)
(-0.707107 0.707107 0.5)
63
             (-0.353553 0.353553 0.5)
64
65
             (-0.707107 2 0.5)
66
67
     );
68
69
70
     blocks
71
           hex (5 4 9 10 24 23 28 29) (10 10 1) simpleGrading (1 1 1)
72
           hex (0 1 4 5 19 20 23 24) (10 10 1) simpleGrading (1 1 1)
           hex (1 2 3 4 20 21 22 23) (20 10 1) simpleGrading (1 1 1)
73
           hex (4 3 6 7 23 22 25 26) (20 20 1) simpleGrading (1 1 1)
74
           hex (9 4 7 8 28 23 26 27) (10 20 1) simpleGrading (1 1 1)
75
           hex (15 16 10 9 34 35 29 28) (10 10 1) simpleGrading (1 1 1)
           hex (12 11 16 15 31 30 35 34) (10 10 1) simpleGrading (1 1 1)
           hex (13 12 15 14 32 31 34 33) (20 10 1) simpleGrading (1 1 1)
78
           hex (14 15 18 17 33 34 37 36) (20 20 1) simpleGrading (1 1 1)
79
           hex (15 9 8 18 34 28 27 37) (10 20 1) simpleGrading (1 1 1)
81
82
83
      edges
84
85
                 0 5 (0.469846 0.17101 -0.5)
           arc 5 10 (0.17101 0.469846 -0.5)
arc 1 4 (0.939693 0.34202 -0.5)
                 4 9 (0.34202 0.939693
88
           arc
           arc 19 24 (0.469846 0.17101 0.5)
arc 24 29 (0.17101 0.469846 0.5)
arc 20 23 (0.939693 0.34202 0.5)
```

OpenVFOAM-1.4

P-48

```
arc 23 28 (0.34202 0.939693 0.5)
          arc 11 16 (-0.469846 0.17101 -0.5)
arc 16 10 (-0.17101 0.469846 -0.5)
          arc 12 15
arc 15 9 (
                       -0.939693 0.34202 -0.5)
                         34202 0.939693
          arc 30 35
                       -0.469846 0.17101 0.5
                       -0.17101 0.469846 0.5)
          arc 31 34 (-0.939693 0.34202 0.5)
          arc 34 28 (-0.34202 0.939693 0.5)
100
101
103
     natches
104
          symmetryPlane down
105
106
               (0 1 20 19)
(1 2 21 20)
107
               (12 11 30 31)
(13 12 31 32)
109
110
111
          patch right
112
               (2 3 22 21)
114
               (3 6 25 22)
115
          symmetryPlane up
117
118
119
               (6 7 26 25)
(8 18 37 27)
121
               (18 17 36 37)
122
123
          patch left
124
125
               (14 13 32 33)
126
127
128
          symmetryPlane cylinder
129
130
               (10 5 24 29)
(5 0 19 24)
131
133
               (16 10 29 35
134
     );
\frac{136}{137}
     mergePatchPairs
139
141
```

3.1.4 Boundary conditions and initial fields

Using FoamX or editing case files by hand, set the boundary conditions in accordance with the problem description in Figure 3.1, i.e. the left boundary should be an Inlet, the right boundary should be an Outlet and the down and cylinder boundaries should be symmetryPlane. The top boundary conditions is chosen so that we can make the most genuine comparison with our analytical solution which uses the assumption that the domain is infinite in the y direction. The result is that the normal gradient of \mathbf{U} is small along a plane coinciding with our boundary. We therefore impose the condition that the normal component is zero, i.e. specify the boundary as a symmetryPlane, thereby ensuring that the comparison with the analytical is reasonable.

3.1.5 Running the case

No fluid properties need be specified in this problem since the flow is assumed to be incompressible and inviscid. In the *system* subdirectory, the *controlDict* specifies the control

Open√FOAM-1.4

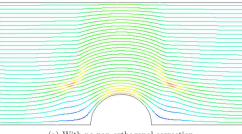
Examples of the use of OpenFOAM

parameters for the run. Note that since we assume steady flow, we only run for 1 time step:

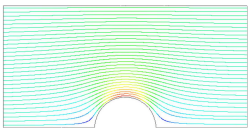
```
OpenFOAM: The Open Source CFD Toolbox
               F ield
              O peration
                             Version: 1.4
      //
               A nd
                             Web:
                                      http://www.openfoam.org
               M anipulation
   FoamFile
10
11
       version
       format
                     ascii;
       case
       instance
                     11.11
       class
                     dictionary
21
   application potentialFoam;
   startFrom
                 startTime;
   startTime
                 0;
                 endTime:
   stopAt
                 1;
   endTime
                 1;
   writeControl
                 timeStep;
   writeInterval
                 1;
   purgeWrite
   writeFormat
                 ascii:
   writePrecision 6:
   writeCompression uncompressed;
                 general;
   timeFormat
   timePrecision
                 6;
   runTimeModifiable ves:
```

potentialFoam executes an iterative loop around the pressure equation which it solves in order that explicit terms relating to non-orthogonal correction in the Laplacian term may be updated in successive iterations. The number of iterations around the pressure equation is controlled by the nNonOrthogonalCorrectors keyword in controlDict. In the first instance we can set nNonOrthogonalCorrectors to 0 so that no loops are performed, i.e. the pressure equation is solved once, and there is no non-orthogonal correction. The solution is shown in Figure 3.3(a) (at t=1, when the steady-state simulation is complete). We expect the solution to show smooth streamlines passing across the domain as in the analytical solution in Figure 3.3(c), yet there is clearly some error in the regions where there is high non-orthogonality in the mesh, e.g. at the join of blocks 0, 1 and 3. The case can be run a second time with some non-orthogonal correction by setting nNonOrthogonalCorrectors to 3. The solution shows smooth streamlines with no significant error due to non-orthogonality as shown in Figure 3.3(b).

P-50



(a) With no non-orthogonal correction



(b) With non-orthogonal correction

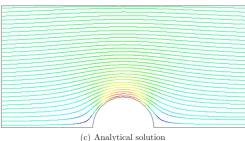


Figure 3.3: Streamlines of potential flow

OpenVFOAM-1.4

Examples of the use of OpenFOAM

3.1.6 Generating the analytical solution

P-52

Source code is included in the $FOAM_TUTORIALS/potentialFoam/analyticalCylinder$ directory to generate the analytical solution for the potential flow case. The velocity at any point at a distance d and angle θ from the cylinder centre is described analytically as

$$U_x = U_{\infty} \left[1 - \left(\frac{r}{d} \right)^2 \cos 2\theta \right]$$

$$U_y = U_{\infty} \left(\frac{r}{d} \right)^2 \sin 2\theta \quad (3.3)$$

where r is the cylinder radius and U_{∞} is the inlet flow velocity. Here, θ describes the angle subtended from the x-axis.

Let us examine some details of the source code in the <code>analyticalCylinder</code> directory. In <code>createFields.H</code>, the velocity field is read in using the <code>IOobject::NO_WRITE</code> option to ensure that the field data can never be overwritten during execution of <code>analyticalCylinder</code>. The inlet velocity and cylinder radius are taken from data read from the mesh and a field <code>UA</code> is set up to store the analytical solution:

```
Info<< "Reading field U\n" << endl;
yolVectorField U</pre>
           IOobject
                 runTime.timeName(),
                mesh,
IOobject::MUST_READ,
IOobject::NO_WRITE
11
12
     Info<< "Reading inlet velocity uInfX\n" << endl;</pre>
16
17
     dimensionedScalar uInfX
18
19
           "uInfx",
dimensionSet(0, 1, -1, 0, 0),
U.boundaryField()[3][0].x()
22
23
     Info << "U at inlet = " << uInfX.value() << " m/s" << endl;</pre>
      dimensionedScalar radius
            "radius"
          dimensionSet(0, 1, 0, 0, 0),
mag(U.mesh().boundary()[4].Cf()[0])
     Info << "Cylinder radius = " << radius.value() << " m" << endl;</pre>
     volVectorField UA
34
35
36
           IOobject
37
                 runTime.timeName(),
                mesh,
IOobject::NO_READ,
IOobject::AUTO_WRITE
41
42
43
44
     );
```

Thea main code analyticalCylinder. C performs the following tasks:

• increments the time step by runTime++;

• generates the analytical solution for field UA using tensor arithmetic;

```
• writes the solution to file by runTime.writeObjects().
                     F ield
                                           OpenFOAM: The Open Source CFD Toolbox
                     O peration
                                           Copyright (C) 1991-2007 OpenCFD Ltd.
                     A nd
                     M anipulation
     License
          This file is part of OpenFOAM.
          {\tt OpenFOAM} is free software; you can redistribute it and/or modify it under the terms of the GNU General Public License as published by the
          Free Software Foundation; either version 2 of the License, or (at your
13
          option) any later version.
14
          OpenFOAM is distributed in the hope that it will be useful, but WITHOUT ANY WARRANTY; without even the implied warranty of MERCHANTABILITY or FINNESS FOR A PARTICULAR PURPOSE. See the GNU General Public License
          for more details.
          You should have received a copy of the GNU General Public License along with OpenFOAM; if not, write to the Free Software Foundation, Inc., 51 Franklin St, Fifth Floor, Boston, MA 02110-1301 USA
     Application
25
          analyticalCylinder
      \begin{array}{c} {\tt Description} \\ {\tt Generates} \ \ {\tt an \ analytical \ solution \ for \ potential \ flow \ around \ a \ cylinder.} \end{array} 
28
          Can be compared with the solution from the potentialFlow/cylinder example.
33
     #include "fvCFD.H"
     int main(int argc, char *argv[])
39
          include "setRootCase.H"
          include "createTime.H' include "createMesh.H'
45
          include "createFields.H'
         Info << "\nEvaluating analytical solution" << endl;</pre>
          volVectorField centres = UA.mesh().C();
52
          volScalarField magCentres = mag(centres);
          volScalarField theta = acos((centres & vector(1,0,0))/magCentres);
          volVectorField cs2theta =
             cos(2*theta)*vector(1,0,0)
+ sin(2*theta)*vector(0,1,0);
          UA = uInfX*(dimensionedVector(vector(1,0,0))
61
             - pow((radius/magCentres),2)*cs2theta);
          runTime.write();
          Info<< "end" << endl;</pre>
          return(0);
68
```

The utility must be compiled with wmake as normal. It can then be run by typing

analyticalCylinder \$FOAM_RUN/potentialFoam cylinder

Open VFOAM-1.4

P-54

The analytical solution is plotted as streamlines as shown in Figure 3.3(c). Note that differences in the analytical and numerical solutions at the top plane are due to the fact that the analytical solution assumes an infinite boundary and the numerical solution specifies a zeroGradient boundary condition at that boundary.

3.1.7 Exercise

Investigate the accuracy of the numerical solution by implementing some measure of comparison between the numerical and analytical in analytical Cylinder.

3.2 Steady turbulent flow over a backward-facing step

In this example we shall investigate steady turbulent flow over a backward-facing step. The problem description is taken from one used by Pitz and Daily in an experimental investigation [**] against which the computed solution can be compared. This example introduces the following OpenFOAM features for the first time:

- generation of a mesh using blockMesh using full mesh grading capability;
- steady turbulent flow.

3.2.1 Problem specification

The problem is defined as follows:

Solution domain The domain is 2 dimensional, consisting of a short inlet, a backward-facing step and converging nozzle at outlet as shown in Figure 3.4.



Figure 3.4: Geometry of backward-facing step

Governing equations

• Mass continuity for incompressible flow

$$\nabla \cdot \mathbf{U} = 0 \tag{3.4}$$

Steady flow momentum equation

$$\nabla \cdot (\mathbf{U}\mathbf{U}) + \nabla \cdot \mathbf{R} = -\nabla p \tag{3.5}$$

where p is kinematic pressure and (in slightly over-simplistic terms) $\mathbf{R} = \nu_{eff} \nabla \mathbf{U}$ is the viscous stress term with an effective kinematic viscosity ν_{eff} , calculated from selected transport and turbulence models.

Initial conditions U = 0 m/s, p = 0 Pa — required in OpenFOAM input files but not necessary for the solution since the problem is steady-state.

Boundary conditions

- Inlet (left) with fixed velocity $\mathbf{U} = (10, 0, 0) \text{ m/s}$;
- Outlet (right) with fixed pressure p = 0 Pa;
- No-slip walls on other boundaries.

Transport properties

• Kinematic viscosity of air $\nu = \mu/\rho = 18.1 \times 10^{-6}/1.293 = 14.0 \ \mu m^2/s$

Turbulence model

- Standard $k \epsilon$:
- Coefficients: $C_{\mu} = 0.09$; $C_1 = 1.44$; $C_2 = 1.92$; $\alpha_k = 1$; $\alpha_{\epsilon} = 0.76923$.

Solver name simpleFoam: an implementation for steady incompressible flow.

Case name pitzDaily, located in the \$FOAM_TUTORIALS/simpleFoam directory.

The problem is solved using simpleFoam, so-called as it is an implementation for steady flow using the SIMPLE algorithm [**]. The solver has full access to all the turbulence models in the incompressibleTurbulenceModels library and the non-Newtonian models incompressibleTransportModels library of the standard OpenFOAM release.

3.2.2 Mesh generation

We expect that the flow in this problem is reasonably complex and an optimum solution will require grading of the mesh. In general, the regions of highest shear are particularly critical, requiring a finer mesh than in the regions of low shear. We can anticipate where high shear will occur by considering what the solution might be in advance of any calculation. At the inlet we have strong uniform flow in the x direction and, as it passes over the step, it generates shear on the fluid below, generating a vortex in the bottom half of the domain. The regions of high shear will therefore be close to the centreline of the domain and close to the walls.

The domain is subdivided into 12 blocks as shown in Figure 3.5.

The mesh is 3 dimensional, as always in OpenFOAM, so in Figure 3.5 we are viewing the back plane along z=-0.5. The full set of vertices and blocks are given in the mesh description file below:

OpenVFOAM-1.4

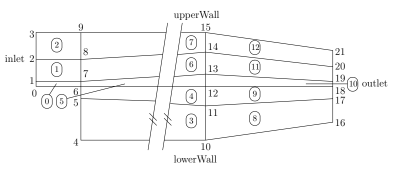


Figure 3.5: Blocks in backward-facing step

```
F ield
                                            OpenFOAM: The Open Source CFD Toolbox
                      0 peration
                                            Version: 1.4
                                                         http://www.openfoam.org
                      A nd
                      M anipulation
     FoamFile
                               2.0;
ascii:
          version
11
12
          root
          instance
local
           class
                               dictionary;
blockMeshDict;
20
21
22
23
24
25
     convertToMeters 0.001;
26
27
28
     vertices
            (-20.6 25.4 -0
(0 -25.4 -0.5)
           (290 16 6 -0.5)
52
```

OpenVFOAM-1.4

P-56

P-58

```
(0 -25.4 0.5)
(0 -5 0.5)
(0 0 0.5)
(0 3 0.5)
(0 12.7 0.5)
(0 25.4 0.5)
(206 -25.4 0.5)
(206 -8.5 0.5)
(206 0.5)
             (206 0 0.5)
(206 6.5 0.5)
             (206 17 0.5)
(206 25.4 0.5)
(290 -16.6 0.5)
(290 -6.3 0.5)
(290 0 0.5)
             (290 4.5 0.5)
(290 11 0.5)
(290 16.6 0.5)
       blocks
            hex (0 6 7 1 22 28 29 23) (18 7 1) simpleGrading (0.5 1.8 1)
            hex (1 7 8 2 23 29 30 24) (18 10 1) simpleGrading (0.5 4 1)
             hex (2 8 9 3 24 30 31 25) (18 13 1) simpleGrading (0.5 0.25 1)
 79
            hex (4 10 11 5 26 32 33 27) (180 18 1) simpleGrading (4 1 1)
 80
            hex (5 11 12 6 27 33 34 28) (180 9 1) edgeGrading (4 4 4 4 0.5 1 1 0.5 1 1 1 1) hex (6 12 13 7 28 34 35 29) (180 7 1) edgeGrading (4 4 4 4 1.8 1 1 1.8 1 1 1 1)
 82
            hex (7 13 14 8 29 35 36 30) (180 10 1) edgeGrading (4 4 4 4 4 1 1 4 1 1 1 1)
 83
             hex (8 14 15 9 30 36 37 31) (180 13 1) simpleGrading (4 0.25 1)
84
            hex (10 16 17 11 32 38 39 33) (25 18 1) simpleGrading (2.5 1 1)
 85
            hex (11 17 18 12 33 39 40 34) (25 9 1) simpleGrading (2.5 1 1)
            hex (12 18 19 13 34 40 41 35) (25 7 1) simpleGrading (2.5 1 1)
             hex (13 19 20 14 35 41 42 36) (25 10 1) simpleGrading (2.5 1 1)
            hex (14 20 21 15 36 42 43 37) (25 13 1) simpleGrading (2.5 0.25 1)
90
93
94
       patches
97
             patch inlet
                   (0 22 23 1)
(1 23 24 2)
101
                   (2 24 25 3)
102
             patch outlet
104
105
106
                   (10 17 39 30)
(17 18 40 39)
(18 19 41 40)
(19 20 42 41)
(20 21 43 42)
108
109
111
112
             wall upperWall
113
                   (3 25 31 9)
(9 31 37 15)
114
115
                   (15 37 43 21)
116
117
             wall lowerWall
118
119
                   (0 6 28 22)
(6 5 27 28)
(5 4 26 27)
120
121
122
                   (4 10 32 26)
(10 16 38 32)
124
125
126
             empty frontAndBack
127
                   (22 28 29 23)
(23 29 30 24)
(24 30 31 25)
(26 32 33 27)
120
130
131
```

Open VFOAM-1.4

```
(27 33 34 28)
(28 34 35 29)
(29 35 36 30)
(30 36 37 31)
(32 38 39 33)
(33 39 40 34)
132
133
134
135
136
137
138
                         (33 39 40 34)
(34 40 41 35)
(35 41 42 36)
(36 42 43 37)
(0 1 7 6)
(1 2 8 7)
(2 3 9 8)
(4 5 11 10)
(5 6 12 11)
(6 7 13 12)
(7 8 14 12)
140
141
143
144
145
146
147
                          (8 9 15 14)
(10 11 17 16
(11 12 18 17
\frac{149}{150}
                          (12 13 19 18)
(13 14 20 19)
151
152
153
                          (14 15 21 20)
154
155
         );
156
         mergePatchPairs
157
158
159
```

A major feature of this problem is the use of the full mesh grading capability of blockMesh that is described in section 6.3.1 of the User Guide. The user can see that blocks 4,5 and 6 use the full list of 12 expansion ratios. The expansion ratios correspond to each edge of the block, the first 4 to the edges aligned in the local x_1 direction, the second 4 to the edges in the local x_2 direction and the last 4 to the edges in the local x_3 direction. In blocks 4, 5, and 6, the ratios are equal for all edges in the local x_1 and x_3 directions but not for the edges in the x_2 direction that corresponds in all blocks to the global y. If we consider the ratios used in relation to the block definition in section 6.3.1 of the User Guide, we realize that different gradings have been prescribed along the left and right edges in blocks 4,5 and 6 in Figure 3.5. The purpose of this differential grading is to generate a fine mesh close to the most critical region of flow, the corner of the step, and allow it to expand into the rest of the domain.

The mesh can be generated using blockMesh from the command line or from within FoamX and viewed as described in previous examples.

3.2.3 Boundary conditions and initial fields

The case files can be viewed, or edited from within FoamX or by hand. In this case, we are required to set the initial and boundary fields for velocity \mathbf{U} , pressure p, turbulent kinetic energy k and dissipation rate ε . The boundary conditions can be specified by setting the physical patch types in FoamX: the upper and lower walls are set to Wall, the left patch to Inlet and the right patch to Outlet. These physical boundary conditions require us to specify a fixedValue at the inlet on \mathbf{U} , k and ε . \mathbf{U} is given in the problem specification, but the values of k and ϵ must be chosen by the user in a similar manner to that described in section 2.1.8.1 of the User Guide. We assume that the inlet turbulence is isotropic and estimate the fluctuations to be 5% of \mathbf{U} at the inlet. We have

$$U'_x = U'_y = U'_z = \frac{5}{100} 10 = 0.5 \text{ m/s}$$
 (3.6)

$$k = \frac{3}{2}(0.5)^2 = 0.375 \text{ m}^2/\text{s}^2$$
 (3.7)

If we estimate the turbulent length scale l to be 10% of the width of the inlet then

$$\varepsilon = \frac{C_{\mu}^{0.75}k^{1.5}}{l} = \frac{0.09^{0.75}0.375^{1.5}}{0.1 \times 25.4 \times 10^{-3}} = 14.855 \,\mathrm{m}^2/\mathrm{s}^3 \tag{3.8}$$

At the outlet we need only specify the pressure p = 0Pa.

3.2.4 Case control

The choices of fvSchemes are as follows: the timeScheme should be SteadyState; the gradScheme and laplacianScheme should be set as default to Gauss; and, the divScheme should be set to UD to ensure boundedness.

Special attention should be paid to the settings of $\mathit{fvTolerances}$. Although the top level simpleFoam code contains only equations for p and \mathbf{U} , the turbulent model solves equations for k, ε and \mathbf{R} , and tolerance settings are required for all 5 equations. A solverTolerance of 10^{-5} and solverRelativeTolerance of 0.1 are acceptable for all variables with the exception of p when 10^{-6} and 0.01 are recommended. Under-relaxation of the solution is required since the problem is steady. A relaxationFactor of 0.7 is acceptable for $\mathbf{U}, k, \varepsilon$ and \mathbf{R} but 0.3 is required for p to avoid numerical instability.

Finally, in *controlDict*, the time step deltaT should be set to 1 since in steady state cases such as this is effectively an iteration counter. With benefit of hindsight we know that the solution requires 1000 iterations reach reasonable convergence, hence endTime is set to 1000. Ensure that the writeFrequency is sufficiently high, *e.g.* 50, that you will not fill the hard disk with data during run time.

3.2.5 Running the case and post-processing

Run the case and post-process the results. After a few iterations, e.g. 50, a vortex develops beneath the corner of the step that is the height of the step but narrow in the x-direction as shown by the vector plot of velocities is shown Figure 3.6(a). Over several iterations the vortex stretches in the x-direction from the step to the outlet until at 1000 iterations the system reaches a steady-state in which the vortex is fully developed as shown in Figure 3.6(b-c).

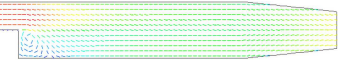
3.3 Supersonic flow over a forward-facing step

In this example we shall investigate supersonic flow over a forward-facing step. The problem description involves a flow of Mach 3 at an inlet to a rectangular geometry with a step near the inlet region that generates shock waves.

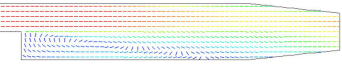
This example introduces the following OpenFOAM features for the first time:

supersonic flow;

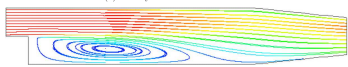
OpenVFOAM-1.4



(a) Velocity vectors after 50 iterations



(b) Velocity vectors at 1000 iterations



(c) Streamlines at 1000 iterations

Figure 3.6: Development of a vortex in the backward-facing step.

3.3.1 Problem specification

The problem is defined as follows:

P-60

Solution domain The domain is 2 dimensional and consists of a short inlet section followed by a forward-facing step of 20% the height of the section as shown in Figure 3.7

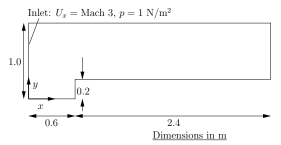


Figure 3.7: Geometry of the forward step geometry

Governing equations

Mass continuity

$$\frac{\partial \rho}{\partial t} + \nabla \cdot (\rho \mathbf{U}) = 0 \tag{3.9}$$

$$p = \rho RT \tag{3.10}$$

• Momentum equation for Newtonian fluid

$$\frac{\partial \rho \mathbf{U}}{\partial t} + \nabla \cdot (\rho \mathbf{U} \mathbf{U}) - \nabla \cdot \mu \nabla \mathbf{U} = -\nabla p \tag{3.11}$$

• Energy equation for fluid (ignoring some viscous terms), $e = C_v T$, with Fourier's Law $\mathbf{q} = -k \nabla T$

$$\frac{\partial \rho e}{\partial t} + \nabla \cdot (\rho \mathbf{U} e) - \nabla \cdot \left(\frac{k}{C_v}\right) \nabla e = p \nabla \cdot \mathbf{U}$$
(3.12)

Initial conditions U = 0 m/s, p = 1 Pa, T = 1 K.

Boundary conditions

- Inlet (left) with fixed Value for velocity $U=3~\mathrm{m/s}=\mathrm{Mach}\ 3,$ pressure $p=1~\mathrm{Pa}$ and temperature $T=1~\mathrm{K};$
- Outlet (right) with zeroGradient on U, p and T;
- No-slip adiabatic wall (bottom);
- Symmetry plane (top).

Transport properties

• Dynamic viscosity of air $\mu = 18.1 \mu Pa s$

Thermodynamic properties

- Specific heat at constant volume $C_v = 1.78571 \text{ J/kg K}$
- Gas constant R = 0.714286 J/kg K
- Conductivity $k = 32.3 \, \mu \text{W/m K}$

Case name forwardStep case located in the \$FOAM_TUTORIALS/sonicFoam directory.

 ${\bf Solver\ name\ sonic}$ sonic Foam: an implementation for compressible trans-sonic/supersonic laminar gas flow.

The case is designed such that the speed of sound of the gas $c = \sqrt{\gamma RT} = 1$ m/s, the consequence being that the velocities are directly equivalent to the Mach number, e.g. the inlet velocity of 3 m/s is equivalent to Mach 3. This speed of sound calculation can be verified using the relationship for a perfect gas, $C_p - Cv = R$, i.e. the ratio of specific heats

$$\gamma = C_p/C_v = \frac{R}{C_v} + 1 \tag{3.13}$$

OpenVFOAM-1.4

3.3.2 Mesh generation

P-62

The mesh used in this case is relatively simple, specified with uniform rectangular cells of length $0.06~\mathrm{m}$ in the x direction and $0.05~\mathrm{m}$ in the y direction. The geometry can simply be divided into 3 blocks, one below the top of the step, and two above the step, one either side of the step front. The full set of vertices and blocks are given in the mesh description file below:

```
OpenFOAM: The Open Source CFD Toolbox
                     O peration
                                          Version:
                     A nd
                                                      http://www.openfoam.org
                    M anipulation
     FoamFile
          version
          format
          instance
          class
                             dictionary;
blockMeshDict;
21
22
     convertToMeters 1;
26
27
     vertices
          (0 0 -0.05)
(0.6 0 -0.05)
(0 0.2 -0.05)
(0.6 0.2 -0.05)
          (0.6 0.2 0.05)
44
45
46
47
48
          hex (0 1 3 2 8 9 11 10) (25 10 1) simpleGrading (1 1 1)
         hex (2 3 6 5 10 11 14 13) (25 40 1) simpleGrading (1 1 1)
          hex (3 4 7 6 11 12 15 14) (100 40 1) simpleGrading (1 1 1)
51
52
53
54
     edges
55
56
57
58
     patches
59
          patch inlet
               (0 8 10 2)
(2 10 13 5)
          patch outlet
               (4 7 15 12)
```

3.3.3 Running the case

The case approaches a steady-state at some time after 5 s. The results for pressure at 10 s are shown in Figure 3.8. The results clearly show discontinuities in pressure, *i.e.* shock waves, emanating from ahead of the base of the step.

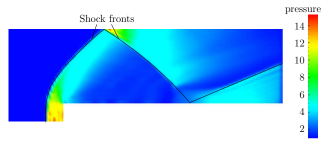


Figure 3.8: Shock fronts in the forward step problem

3.3.4 Exercise

The user can examine the effect on the solution of increasing the inlet velocity.

3.4 Decompression of a tank internally pressurised with water

In this example we shall investigate a problem of rapid opening of a pipe valve close to a pressurised liquid-filled tank. The prominent feature of the result in such cases is the propagation of pressure waves which must therefore be modelled as a compressible liquid.

This tutorial introduces the following OpenFOAM features for the first time:

• Mesh refinement

OpenVFOAM-1.4

Pressure waves in liquids

P-64

3.4.1 Problem specification

Solution domain The domain is 2 dimensional and consists of a tank with a small outflow pipe as shown in Figure 3.9

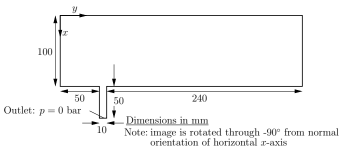


Figure 3.9: Geometry of a tank with outflow pipe

Governing equations This problem requires a model for compressibility ψ in the fluid in order to be able to resolve waves propagating at a finite speed. A barotropic relationship is used to relate density ρ and pressure p are related to ψ .

Mass continuity

$$\frac{\partial \rho}{\partial t} + \nabla \cdot (\rho \mathbf{U}) = 0 \tag{3.14}$$

• The barotropic relationship

$$\frac{\partial \rho}{\partial p} = \frac{\rho}{K} = \psi \tag{3.15}$$

where K is the bulk modulus

• Equation 3.15 is linearised as

$$\rho \approx \rho_0 + \psi \left(p - p_0 \right) \tag{3.16}$$

where ρ_0 and p_0 are the reference density and pressure respectively such that $\rho(p_0) = \rho_0$.

• Momentum equation for Newtonian fluid

$$\frac{\partial \rho \mathbf{U}}{\partial t} + \nabla \cdot (\rho \mathbf{U} \mathbf{U}) - \nabla \cdot \mu \nabla \mathbf{U} = -\nabla p \tag{3.17}$$

 $Boundary\ conditions$ Using FoamX the following physical boundary conditions can be set:

- outerWall is specified the wall condition;
- axis is specified as the symmetryPlane;
- nozzle is specified as a pressureOutlet where p = 0 bar.
- front and back boundaries are specified as empty.

Initial conditions U = 0 m/s, p = 100 bar.

Transport properties

• Dynamic viscosity of water $\mu = 1.0 \text{ mPa s}$

Thermodynamic properties

- Density of water $\rho = 1000 \text{ kg/m}^3$
- Reference pressure $p_0 = 1$ bar
- Compressibility of water $\psi = 4.54 \times 10^{-7} \text{ s}^2/\text{m}^2$

Solver name sonicLiquidFoam: a compressible sonic laminar liquid flow code.

 $\label{located} \textit{Case name decompressionTank} \ \text{case located in the $FOAM_TUTORIALS/sonicLiquidFoam directory.}$

3.4.2 Mesh Generation

The full geometry is modelled in this case; the set of vertices and blocks are given in the mesh description file below:

```
F ield
                                             OpenFOAM: The Open Source CFD Toolbox
                      O peration
                                            Version: 1.4
                                                          http://www.openfoam.org
                      A nd
                      M anipulation
      FoamFile
           version
                                2.0;
           instance
                                dictionary;
blockMeshDict;
           class
20
21
24
25
     convertToMeters 0.1;
     vertices
           (0 \ 0 \ -0.1)
           (1 0 -0.1)
(1 0 -0.1)
(0 0.5 -0.1)
(1 0.5 -0.1)
(1.5 0.5 -0.1)
           (0 0.6 -0.1)
(1 0.6 -0.1)
```

OpenVFOAM-1.4

```
(0 3 -0.1)
(1 3 -0.1)
(0 0 0.1)
(1 0 0.1)
(0 0.5 0.1)
(1 0.5 0.1)
(1.5 0.5 0.1)
(1 0.6 0.1)
(1 0.6 0.1)
(1 0.5 0.6 0.1)
37
38
39
40
41
42
43
            (0 3 0.1)
(1 3 0.1)
48
49
50
51
52
      );
      blocks
           hex (0 1 3 2 10 11 13 12) (30 20 1) simpleGrading (1 1 1)
54
           hex (2 3 6 5 12 13 16 15) (30 5 1) simpleGrading (1 1 1)
           hex (3 4 7 6 13 14 17 16) (25 5 1) simpleGrading (1 1 1)
55
           hex (5 6 9 8 15 16 19 18) (30 95 1) simpleGrading (1 1 1)
56
57
58
59
      edges
60
61
62
63
      patches
64
65
            wall outerWall
                 (0 1 11 10)
(1 3 13 11)
(3 4 14 13)
(7 6 16 17)
(6 9 19 16)
                 (9 8 18 19)
            symmetryPlane axis
75
76
77
78
79
80
                 (0 10 12 2)
(2 12 15 5)
(5 15 18 8)
           patch nozzle
                 (4 7 17 14)
82
            empty back
84
                  (0 2 3 1)
            empty front
                  (10 11 13 12)
                 (12 13 16 15)
(13 14 17 16)
                  (15 16 19 18)
      );
99
100
      mergePatchPairs
\frac{101}{102}
```

In order to improve the numerical accuracy, we shall use the reference level of 1 bar for the pressure field. Note that both the internal field level and the boundary conditions are offset by the reference level.

P-66

3.4.3 Preparing the Run

Before we commence the setup of the calculation, we need to consider the characteristic velocity of the phenomenon we are trying to capture. In the case under consideration, the fluid velocity will be very small, but the pressure wave will propagate with the speed of sound in water. The speed of sound is calculated as:

$$c = \sqrt{\frac{1}{\psi}} = \sqrt{\frac{1}{4.54 \times 10^{-7}}} = 1483.2 \text{m/s}.$$
 (3.18)

For the mesh described above, the characteristic mesh size is approximately 2 mm (note the scaling factor of 0.1 in the *blockMeshDict* file). Using

$$Co = \frac{U \,\Delta t}{\Delta x} \tag{3.19}$$

a reasonable time step is around $\Delta t = 5 \times 10^{-7} \mathrm{s}$, giving the Co number of 0.35, based on the speed of sound. Also, note that the reported Co number by the code (associated with the convective velocity) will be two orders of magnitude smaller. As we are interested in the pressure wave propagation, we shall set the simulation time to 0.25 ms. For reference, the *controlDict* file is quoted below.

```
OpenFOAM: The Open Source CFD Toolbox
                  F ield
                 O peration
                                   Version:
                                             1.4
                                              http://www.openfoam.org
                  M anipulation
    FoamFile
10
         version
        instance
        class
                         dictionary;
20
21
23
    application sonicLiquidFoam;
27
    startFrom
                     startTime;
29
    startTime
    stopAt
                     endTime;
    endTime
                    0.0001;
                     5e-07;
    writeControl
                    timeStep;
    writeInterval
                    20:
    purgeWrite
                    0;
    writeFormat
                    ascii;
    writePrecision 6;
    writeCompression compressed;
```

OpenVFOAM-1.4



3.4.4 Running the case

P-68

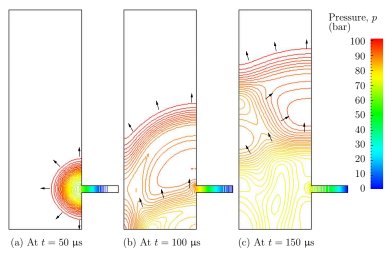


Figure 3.10: Propagation of pressure waves

The user can run the case and view results in dxFoam. The liquid flows out through the nozzle causing a wave to move along the nozzle. As it reaches the inlet to the tank, some of the wave is transmitted into the tank and some of it is reflected. While a wave is reflected up and down the inlet pipe, the waves transmitted into the tank expand and propagate through the tank. In Figure 3.10, the pressures are shown as contours so that the wave fronts are more clearly defined than if plotted as a normal isoline plot.

If the simulation is run for a long enough time for the reflected wave to return to the pipe, we can see that negative absolute pressure is detected. The modelling permits this and has some physical basis since liquids can support tension, *i.e.* negative pressures. In reality, however, impurities or dissolved gases in liquids act as sites for cavitation, or vapourisation/boiling, of the liquid due to the low pressure. Therefore in practical situations, we generally do not observe pressures falling below the vapourisation pressure of the liquid; not at least for longer than it takes for the cavitation process to occur.

Figure 3.11: Propagation of pressure waves with refined mesh

3.4.5 Improving the solution by refining the mesh

Looking at the evolution of the resulting pressure field in time, we can clearly see the propagation of the pressure wave into the tank and numerous reflections from the inside walls. It is also obvious that the pressure wave is smeared over a number of cells. We shall now refine the mesh and reduce the time step to obtain a sharper front resolution. Simply edit the *blockMeshDict* and increase the number of cells by a factor of 4 in the x and y directions, *i.e.* block 0 becomes (120 80 1) from (30 20 1) and so on. Run blockMesh on this file. In addition, in order to maintain a Courant number below 1, the time step must be reduced accordingly to $\Delta t = 10^{-7}$ s. The second simulation gives considerably better resolution of the pressure waves as shown in Figure 3.11.

3.5 Magnetohydrodynamic flow of a liquid

In this example we shall investigate an flow of an electrically-conducting liquid through a magnetic field. The problem is one belonging to the branch of fluid dynamics known as magnetohydrodynamics (MHD) that uses mhdFoam.

3.5.1 Problem specification

The problem is known as the Hartmann problem, chosen as it contains an analytical solution with which mhdFoam can be validated. It is defined as follows:

Solution domain The domain is 2 dimensional and consists of flow along two parallel plates as shown in Fig. 3.12.

Governing equations



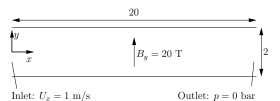


Figure 3.12: Geometry of the Hartmann problem

• Mass continuity for incompressible fluid

$$\nabla \cdot \mathbf{U} = 0 \tag{3.20}$$

• Momentum equation for incompressible fluid

$$\frac{\partial \mathbf{U}}{\partial t} + \nabla \bullet \left(\mathbf{U} \mathbf{U} \right) + \nabla \bullet \left(2 \mathbf{B} \Gamma_{\mathbf{B} \mathbf{U}} \mathbf{B} \right) + \nabla \bullet \left(\nu \mathbf{U} \right) + \nabla \left(\Gamma_{\mathbf{B} \mathbf{U}} \mathbf{B} \mathop{:}^{\bullet} \mathbf{B} \right) = - \nabla p \ \, (3.21)$$

where **B** is the magnetic flux density, $\Gamma_{\mathbf{B}\mathbf{U}} = (2\mu\rho)^{-1}$.

• Maxwell's equations

P-70

$$\nabla \times \mathbf{E} = -\frac{\partial \mathbf{B}}{\partial t} \tag{3.22}$$

where E is the electric field strength.

$$\nabla \cdot \mathbf{B} = 0 \tag{3.23}$$

$$\nabla \times \mathbf{H} = \mathbf{J} + \frac{\partial \mathbf{D}}{\partial t} = \mathbf{J} \tag{3.24}$$

assuming $\partial \mathbf{D}/\partial t \ll \mathbf{J}$. Here, \mathbf{H} is the magnetic field strength, \mathbf{J} is the current density and \mathbf{D} is the electric flux density.

• Charge continuity

$$\nabla \cdot \mathbf{J} = 0 \tag{3.25}$$

• Constitutive law

$$\mathbf{B} = \mu \mathbf{H} \tag{3.26}$$

• Ohm's law

$$\mathbf{J} = \sigma \left(\mathbf{E} + \mathbf{U} \times \mathbf{B} \right) \tag{3.27}$$

• Combining Equation 3.22, Equation 3.24, Equation 3.27, and taking the curl

$$\frac{\partial \mathbf{B}}{\partial t} + \nabla \cdot (\mathbf{U}\mathbf{B}) - \nabla \cdot (\phi_{\mathbf{B}}\mathbf{U}) - \nabla \cdot (\Gamma_{\mathbf{B}}\mathbf{B}) = 0$$
(3.28)

- inlet is specified the inlet condition with fixed velocity U = (1, 0, 0) m/s;
- outlet is specified as the outlet with with fixed pressure p = 0 Pa;
- upperWall is specified as a wall where $\mathbf{B} = (0, 20, 0) \mathrm{T}$.
- lowerWall is specified as a wall where $\mathbf{B} = (0, 20, 0) \mathrm{T}$.
- front and back boundaries are specified as empty.

Initial conditions U = 0 m/s, p = 100 Pa, B = (0, 20, 0) T.

$Transport\ properties$

- Kinematic viscosity $\nu = 1$ Pas
- Density $\rho = 1 \text{ kg m/s}$
- Electrical conductivity $\sigma = 1 \ (\Omega \, m)^{-1}$
- Permeability $\mu = 1 \text{ H/m}$

Solver name mhdFoam: an incompressible laminar magneto-hydrodynamics code.

Case name hartmann case located in the \$FOAM_TUTORIALS/mhdFoam directory.

3.5.2 Mesh generation

The geometry is simply modelled with 100 cells in the x-direction and 40 cells in the y-direction; the set of vertices and blocks are given in the mesh description file below:

```
F ield
                                            OpenFOAM: The Open Source CFD Toolbox
         //
                      O peration
                                            Version: 1.4
                      A nd
                                                         http://www.openfoam.org
                      M anipulation
      FoamFile
           version
                                2.0:
11
           instance
           class
                               dictionary; blockMeshDict:
21
23
24
25
     convertToMeters 1;
     vertices
          (0 -1 0)
(20 -1 0)
(20 1 0)
(0 1 0)
(0 -1 0.1
           (20 -1 0.1)
(20 1 0.1)
```

Open VFOAM-1.4

```
);
39
40
41
   blocks
       hex (0 1 2 3 4 5 6 7) (100 40 1) simpleGrading (1 1 1)
42
43
44
   edges
45
46
47
   patches
48
       patch inlet
50
           (0 4 7 3)
54
       patch outlet
55
           (2651)
       patch lowerWall
59
           (1540)
       patch upperWall
62
           (3 7 6 2)
65
66
       empty frontAndBack
67
68
69
70
71
72
73
   mergePatchPairs
74
75
```

3.5.3 Running the case

P-72

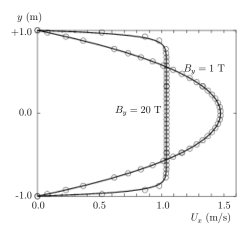
The user can run the case and view results in dxFoam. It is also useful at this stage to run the Ucomponents utility to convert the U vector field into individual scalar components. MHD flow is governed by, amongst other things, the Hartmann number which is a measure of the ratio of electromagnetic body force to viscous force

$$M = BL\sqrt{\frac{\sigma}{\rho\nu}} \tag{3.29}$$

where L is the characteristic length scale. In this case with $B_y=20$ T, M=20 and the electromagnetic body forces dominate the viscous forces. Consequently with the flow fairly steady at t=2 s the velocity profile is almost planar, viewed at a cross section midway along the domain x=10 m. The user can plot a graph of the profile of U_x in dxFoam. Now the user should reduce the magnetic flux density ${\bf B}$ to 1 Tand re-run the code and Ucomponents. In this case, M=1 and the electromagnetic body forces no longer dominate. The velocity profile consequently takes on the parabolic form, characteristic of Poiseuille flow as shown in Figure 3.13. To validate the code the analytical solution for the velocity profile U_x is superimposed in Figure 3.13, given by:

$$\frac{U_x(y)}{U_x(0)} = \frac{\cosh M - \cosh M(y/L)}{\cosh M - 1} \tag{3.30}$$

where the characteristic length L is half the width of the domain, i.e. 1 m.



3.5 Magnetohydrodynamic flow of a liquid

Figure 3.13: Velocity profile in the Hartmann problem for $B_y=1~\mathrm{T}$ and $B_y=20~\mathrm{T}$.

Index P-75 P-76 Index

Index

Symbols Numbers A B C D E F G H I J K L M N O P Q R S T U V W X Z

Symbols	access functions, P-23
*	Actor color button, U-174
tensor member function, P-25	adiabaticFlameT utility, U-94
+	adjustableRunTime
tensor member function, P-25	keyword entry, U-61, U-108
_	adjustPhi tools, U-95
tensor member function, P-25	adjustTimeStep keyword, U-61
/	agglomerator keyword, U-119
tensor member function, P-25	algorithms tools, U-95
/**/	allTime
C++ syntax, U-78	menu entry, U-131
//	analytical solution, P-45
C++ syntax, U-78	anisotropicFilter model, U-98
OpenFOAM file syntax, U-102	Annotate window panel, U-27, U-174
# include	ansysToFoam utility, U-90
C++ syntax, U-72, U-78	APIfunctions model, U-97
& &	applicationClass keyword, U-107
tensor member function, P-25	applications, U-69
&&	arbitrarily unstructured, P-31
tensor member function, P-25	arc
•	keyword entry, U-48, U-156
tensor member function, P-25	arc keyword, U-155
<pre><lesmodel>Coeffs keyword, U-191</lesmodel></pre>	ascii
<pre><delta>Coeffs keyword, U-191</delta></pre>	keyword entry, U-108
<pre><turbulencemodel>Coeffs keyword, U-191</turbulencemodel></pre>	attachMesh utility, U-90
0.000000e+00 directory, U-102	autoPatch utility, U-91
1-dimensional mesh, U-145	axes
1D mesh, U-145	right-handed, U-154
2-dimensional mesh, U-145	right-handed rectangular Cartesian, P-15
2D mesh, U-145	U-21
3D View button, U-174, U-175	axi-symmetric cases, U-150, U-160
3D view Properties	axi-symmetric mesh, U-145
menu entry, U-27, U-174-U-176	
mena energy, 0-27, 0-174 0-170	В
Numbers	background
0 directory, U-102	process, U-28, U-82
v /	backward
\mathbf{A}	keyword entry, U-116
Accept button, U-173	Backward differencing, P-39

Open∇FOAM-1.4

Open ∇ FOAM-1.4

1 . T. 1 . IM 1.1	. DI DOT ILITO
basicThermophysicalModels	symmetryPlane, P-65, U-150
library, U-96	totalPressure, U-153
binary	turbulentInlet, U-153
keyword entry, U-108	wall, U-42
BirdCarreau model, U-99	wall, P-65, P-71, U-150
blended differencing, P-38	wallBuoyantPressure, U-153
block	wedge, U-145, U-150, U-151, U-160
expansion ratio, U-157	zeroGradient, U-152
block keyword, U-155	boundary conditions, P-43
blockMesh solver, P-47	Dirichlet, P-43
blockMesh utility, U-39, U-90, U-151	inlet, P-44
blockMesh	Neumann, P-43
menu entry, U-22, U-33	no-slip impermeable wall, P-44
blockMesh executable	outlet, P-44
vertex numbering, U-156	physical, P-44
blockMeshDict	symmetry plane, P-44
dictionary, U-21, U-22, U-37, U-48, U-49,	boundary type
U-154, U-161	empty, U-134
blocks keyword, U-22, U-156	wall, U-42
bound tools, U-95	boundaryField keyword, U-106
boundaries, U-145	boundaryFoam solver, U-87
boundary, U-145	bounded
boundary	keyword entry, U-114, U-115
dictionary, U-144, U-154	boxToCell keyword, U-60
boundary condition	boxTurb utility, U-90
calculated, U-152	breaking of a dam, U-57
cyclic, U-151	bubbleFoam solver, U-88
directionMixed, U-152	buoyantFoam solver, U-89
empty, P-65, P-71, U-21, U-145, U-150	buoyantSimpleFoam solver, U-89
fixedGradient, U-152	button
fixedValue, U-152	3D View, U-174, U-175
fluxCorrectedVelocity, U-153	Accept, U-173
gammaContactAngle, U-59	Actor color, U-174
inlet, P-71	Close Case, U-33
inletOutlet, U-153	Compact, U-133
mixed, U-152	Delete, U-173
movingWallVelocity, U-153	Display Orientation Axes, U-174
outlet, P-71	Info, U-133
outletInlet, U-153	My Jobs, U-133
partialSlip, U-153	Orientation Axes, U-27
patch, U-150	Refresh Case Browser, U-41
pressureDirectedInletVelocity, U-153	Reset Range, U-28
pressureInletVelocity, U-153	Reset, U-173
pressureOutlet, P-65	Start Calculation Now, U-28
pressureTransmissive, U-153	Start Calculation, U-35
processor, U-151	Use parallel projection, U-27, U-174
setup, U-23	cont, U-133
slip, U-153	endNow, U-133
supersonicFreeStream, U-153	end, U-133
surfaceNormalFixedValue, U-153	kill, U-133
	/ = ==

Index P-77 P-78 Index

purge, U-133	cGamma keyword, U-63
read, U -133	channelOodles solver, U-88
status, U-133	checkMesh utility, U-91, U-162
suspend, U-133	checkYPlus utility, U-93
C	chemistryModel
C	library, U-97
C++ syntax	chemistryModel model, U-97
/**/, U-78	chemistrySolver model, U-97
//, U-78	chemkinMixture model, U-96, U-188
# include, U-72, U-78	chemkinToFoam utility, U-94
cacheAgglomeration keyword, U-119	Class menu, U-129
calculated	class
boundary condition, U-152	cell, P-31
Camera window panel, U-174	dimensionSet, P-26, P-32, P-33
Camera Controls window panel, U-174	face, P-31
Camera Orientation window panel, U-174	finiteVolumeCalculus, P-36
case	finiteVolumeMethod, P-36
browser, U-127	fvMesh, P-31
server, U-133	fvSchemes, P-38
case keyword, U-103	fvc, P-36
case manager	fvm, P-36
FoamX, U-123	*
Case Name text box, U-129	pointField, P-31
Case Root text box, U-129	polyBoundaryMesh, P-31
caseRoots keyword, U-19	polyMesh, P-31, U-141, U-143
cases, U-101	polyPatchList, P-31
cavitatingFoam solver, U-88	polyPatch, P-31
cavity flow, U-19	scalarField, P-29
ccm26ToFoam utility, U-90	scalar, P-24
CEI_ARCH	slice, P-31
environment variable, U-180	symmTensorField, P-29
CEI_HOME	symmTensorThirdField, P-29
environment variable, U-180	tensorField, P-29
cell	tensorThirdField, P-29
expansion ratio, U-157	tensor, P-24
cell class, P-31	vectorField, P-29
cell	vector, P-24, U-105
keyword entry, U-182	word, P-26, P-31
cellDecompFiniteElement	class keyword, U-103
library, U-95	clockTime
cellPoint	keyword entry, U-108
keyword entry, U-182	Close Case button, U-33
cellPointFace	cloud keyword, U-183
keyword entry, U-182	cmptAv
cells	tensor member function, P-25
dictionary, U-154	Co utility, U-92
cellSet utility, U-91	cofactors
central differencing, P-38	tensor member function, P-25
cfdTools	coldEngineFoam solver, U-89
library, U-95	Color by menu, U-174
cfxToFoam utility, U-90, U-161	combustionThermophysicalModels
cix for built definey, 0-50, 0-101	compastion i nermophysical Models

library, U-96	keyword entry, U-113
comments, U-78	curl, P-37
Compact button, U-133	curl
compressed	fvc member function, P-37
keyword entry, U-108	curve keyword, U-183
compressible tools, U-95	cyclic
compressibleLESmodels	boundary condition, U-151
library, U-99	cyclic
compressible Turbulence Models	keyword entry, U-150
library, U-98	cylinder
constant directory, U-101, U-187	flow around a, P-45
constLaminarFlameSpeed model, U-96	ъ
constTransport model, U-97, U-188	D
cont button, U-133	d2dt2
containers tools, U-95	fvc member function, P-37
continuum	fvm member function, P-37
mechanics, P-15	dam
control	breaking of a, U-57
of time, U-107	db tools, U-95
controlDict	ddt
dictionary, P-67, U-24, U-34, U-43, U-52,	fvc member function, P-37
U-61, U-101, U-167	fvm member function, P-37
	DeardorffDiffStress model, U-99
controlDict file, P-49	decomposePar utility, U-82, U-83, U-94
convection, see divergence, P-38	decomposeParDict
convergence, U-41	dictionary, U-82
convertToMeters keyword, U-155	decomposition
coodles solver, U-88	of field, U-82
coordinate	of mesh, U-82
system, P-15	decompression of a tank, P-63
coordinate system, U-21	defaultFieldValues keyword, U-60
CORBA, U-96, U-123	deformedGeom utility, U-91
corrected	Delete button, U-173
keyword entry, U-114, U-115	delta keyword, U-84, U-191
couplePatches utility, U-91	deltaT keyword, U-108
Courant number, P-42, U-25	dependencies, U-72
cpuTime	dependency lists, U-72
keyword entry, U-108	det
Crank Nicholson	tensor member function, P-25
temporal discretisation, P-43	determinant, see tensor, determinant
CrankNicholson	dev
keyword entry, U-116	tensor member function, P-25
createPatch utility, U-91	diag
cross product, see tensor, vector cross product	tensor member function, P-25
CrossPowerLaw	diagonal
keyword entry, U-61	keyword entry, U-119
CrossPowerLaw model, U-99	DIC
cubeRootVolDelta model, U-98	keyword entry, U-119
cubicCorrected	DICGaussSeidel
keyword entry, U-116	keyword entry, U-119
cubicCorrection	Dictionaries dictionary tree, U-135

Index P-79 P-80 Index

11.41	diMarkadirekan ana dali 11.07
dictionary	diMethylEther model, U-97
PISO, U-26	direct numerical simulation, U-62
· · · · · · · · · · · · · · · · · · ·	directionMixed
U-48, U-49, U-154, U-161	boundary condition, U-152
boundary, U-144, U-154	directory
cells, U-154	0.000000e+00, U-102 0, U-102
controlDict, P-67, U-24, U-34, U-43, U-52,	
U-61, U-101, U-167	Make, U-73 constant, U-101, U-187
decomposeParDict, U-82	fluentInterface, U-177
faces, U-143, U-154	polyMesh, U-101, U-143
fvSchemes, U-62, U-101, U-109, U-110	processorN, U-84
fvSolution, U-101, U-117	run, U-101
mechanicalProperties, U-51	system, P-49, U-101
neighbour, U-144	tutorials, P-45, U-19
owner, U-143	discretisation
points, U-143, U-154	equation, P-33
thermalProperties, U-51, U-52	Display window panel, U-27, U-28, U-172, U-174
thermophysicalProperties, U-187 transportProperties, U-24, U-41, U-43	Display Orientation Axes button, U-174
turbulenceProperties, U-24, U-41, U-45	distance
	keyword entry, U-183
dictionary tree Dictionaries, U-135	distributed keyword, U-84, U-86
Fields, U-23, U-134	div
Mesh, U-23	fvc member function, P-37
Patches, U-23	fvm member function, P-37
dieselEngineFoam solver, U-89	divergence, P-37, P-39
dieselFoam solver, U-89	divSchemes keyword, U-110
dieselMixture model, U-96, U-188	divU utility, U-92
dieselSpray	dnsFoam solver, U-88
library, U-96	double inner product, see tensor, double inner
diEthylEther model, U-97	product
differencing	dynamicMesh
Backward, P-39	library, U-95
blended, P-38	dynMixedSmagorinsky model, U-98
central, P-38	dynOneEqEddy model, U-98, U-99
Euler implicit, P-39	dynSmagorinsky model, U-98
Gamma, P-38	\mathbf{E}
MINMOD, P-38	
SUPERBEE, P-38	edgeGrading keyword, U-157 edgeMesh
upwind, P-38	library, U-95
van Leer, P-38	edges keyword, U-155
DILU	electrostaticFoam solver, U-89
keyword entry, U-119	empty
dimension	boundary condition, P-65, P-71, U-21,
checking in OpenFOAM, P-26	U-145, U-150
dimensioned <type> template class, P-26</type>	empty boundary type, U-134
dimensionedTypes tools, U-95	empty
dimensions keyword, U-106	keyword entry, U-150
dimensionSet class, P-26, P-32, P-33	end button, U-133
dimensionSet tools, U-95	endNow button, U-133

endTime keyword, U-25, U-107, U-108	Euler implicit
engine	differencing, P-39
library, U-96	temporal discretisation, P-42
engineCompRatio utility, U-93	examples
engineFoam solver, U-89	decompression of a tank, P-63
engineSwirl utility, U-90	flow around a cylinder, P-45
ensight74FoamExec utility, U-180	flow over backward step, P-54
ensight76FoamExec utility, U-92	Hartmann problem, P-69
ENSIGHT7_INPUT	supersonic flow over forward step, P-5
environment variable, U-180	explicit
ENSIGHT7_READER	temporal discretisation, P-42
environment variable, U-180	exponential model, U-97
enstrophy utility, U-92	extrudeMesh utility, U-90
environment variable	D
CELARCH, U-180	\mathbf{F}
CEI_HOME, U-180	face class, P-31
ENSIGHT7_INPUT, U-180	face keyword, U-183
ENSIGHT7_READER, U-180	faceAreaPair
FOAMX_PATH, U-139	keyword entry, U-119
FOAMX_SYSTEM_CONFIG, U-139	faceDecompFiniteElement
FOAMX_USER_CONFIG, U-139	library, U-95
FOAM_RUN, U-101, U-139	faces
JAVA_HOME, U-139	dictionary, U-143, U-154
WM_ARCH, U-76	faceSet utility, U-91
WM_COMPILER_BIN, U-76	FDIC
WM_COMPILER_DIR, U-76	keyword entry, U-119
WM_COMPILER_LIB, U-76	field
WM_COMPILER, U-76	U, U-25
WM_COMPILE_OPTION, U-76	p, U-25
	decomposition, U-82
WM_DIR, U-76	FieldField <type> template class, P-32</type>
WM_JAVAC_OPTION, U-76	Fields dictionary tree, U-23, U-134
WM_LINK_LANGUAGE, U-76	Fields window, U-28
WM_MPLIB, U-76	fields, P-29
WM_OPTIONS, U-76	mapping, U-167
WM_PROJECT_DIR, U-76	fields tools, U-95
WM_PROJECT_INST_DIR, U-76	fields keyword, U-182
WM_PROJECT_LANGUAGE, U-76	Field <type> template class, P-29</type>
WM_PROJECT_USER_DIR, U-76	fieldValues keyword, U-60
WM_PROJECT_VERSION, U-76	file
WM_PROJECT, U-76	FoamX.cfg, U-139
WM_SHELL, U-76	FoamXClient.cfg, U-124, U-138
wmake, U-76	Make/files, U-75
environmentalProperties file, U-61	controlDict, P-49
equilibriumCO utility, U-94	environmentalProperties, U-61
equilibriumFlameT utility, U-94	files, U-73
errorEstimation	options, U-73
library, U-95	transportProperties, U-60
estimateScalarError utility, U-94	file format, U-102
Euler	files file, U-73
keyword entry, U-116	financialFoam solver, U-89

Index P-81 P-82 Index

discretisation, P-27 mesh, P-31 finiteVolume tools, U-95 finiteVolumeCalculus class, P-36 finiteVolumeMethod class, P-36 finiteVolume Method class, P-36 finiteVolumeMethod class, P-36 finiteVolume Method class, P-36 finiteVolume Individual Class firation Method Class P-36 foamX_ERIT_U-124 JAVA GUI, U-125 nam server, U-124 hoat Vuility, U-90 foamX_ERIT_CoNFIG environment variable, U-139 foamX_ERIT_CoNFIG environment variable, U-139 foamX_ERIT_U-13 foamX_ERIT_U-13 foamX_ERIT_U-13 foamX_ERIT_U-13 foamX_ERIT_U-13 foamX_ERIT_U-13 foamX_ERIT_U-13 foamX_ERIT_U-13 foamX_ERIT_U	finite volume	anga browsan II 197
mesh, P-31 finiteVolume tools, U-95 finiteVolumeCalculus class, P-36 finiteVolumeMethod class, P-36 finiteVolumeMethod class, P-36 firstTime menu entry, U-131 firstTime keyword, U-107 fixed keyword entry, U-108 fixedGradient boundary condition, U-152 fixedValue boundary condition, U-152 fixetdValue boundary condition, U-152 fiattenMesh utility, U-91 flow free surface, U-57 laminar, U-19 steady, turbulent, P-54 supersonic, P-59 turbulent, U-20 flow around a cylinder, P-45 flow over backward step, P-54 flowType utility, U-92 fluentMeshToFoam utility, U-90, U-161 fluxCorrectedVelocity boundary condition, U-153 fluxRequired keyword, U-110 OpenFOAM cases, U-101 FOAM.RuN environment variable, U-101, U-139 Foam Utilities menu, U-22, U-33, U-34 foamCorrectvft script/alias, U-166 foamDataToFluent utility, U-92 foamToFieldview9 utility, U-94 foamlob script/alias, U-184 foamlog script/alias, U-185 foamMeshToFluent utility, U-92 foamToFieldview9 utility, U-92		case browser, U-127
finiteVolume tools, U-95 finiteVolumeCalculus class, P-36 finiteVolumeMethod class, P-36 finiteVolumeMethod class, P-36 firstTime menu entry, U-131 firstTime keyword, U-107 fixed keyword entry, U-108 fixedGradient boundary condition, U-152 fixedValue boundary condition, U-152 fixedValue boundary condition, U-152 flattenMesh utility, U-91 flow free surface, U-57 laminar, U-19 steady, turbulent, P-54 supersonic, P-59 turbulent, U-20 flow around a cylinder, P-45 flowType utility, U-92 fluentImterface directory, U-177 fluentMesh FOFOAM case manager, U-123 host browser, U-124 floAMX_CRIT, U-124 FoamX.cf gile, U-139 FOAMX_PATH environment variable, U-139 FOAMX_STEM_CONFIG environment variable, U-139 FOAMX_USER.CONFIG environment variable, U-139 FoamVallent.cfg file, U-124, U-138 foreground process, U-28 format keyword, U-103 fourth keyword entry, U-114, U-115 fvc class, P-36 fvc member function curl, P-37 ddt, P-37 ddt, P-37 ddt, P-37 suffrad, P-		9 ,
finiteVolumeMethod class, P-36 firistTime menu entry, U-131 firstTime keyword, U-107 fixed keyword entry, U-108 fixedGradient boundary condition, U-152 fixedValue boundary condition, U-152 fixedWalue boundary condition, U-152 fixedWalue boundary condition, U-152 fixedWalue boundary condition, U-152 fixedWalue boundary condition, U-152 flow free surface, U-57 laminar, U-19 steady, turbulent, P-54 supersonic, P-59 turbulent, U-20 flow around a cylinder, P-45 flow over backward step, P-54 flowType utility, U-92 flow around a cylinder, P-45 flow over backward step, P-54 flowType utility, U-92 flow around a cylinder, P-45 flow over backward step, P-54 flowType utility, U-92 flow around a cylinder, P-45 flow over backward step, P-54 flowType utility, U-92 flow around a cylinder, P-45 flow over backward step, P-54 flowType utility, U-92 flow around a cylinder, P-45 flow over backward step, P-54 flowType utility, U-92 flow around a cylinder, P-45 flow over backward step, P-54 flowType utility, U-92 flow around a cylinder, P-45 flow over backward step, P-54 flowType utility, U-92 flow around a cylinder, P-45 flow over backward step, P-54 flowType utility, U-92 flow around a cylinder, P-45 flow over backward step, P-54 flowType utility, U-92 flow around a cylinder, P-45 flow Type utility, U-92 flow around a cylinder, P-45 flow Type utility, U-92 flow around a cylinder, P-45 flow Type utility, U-91 flow caread, turbulent, P-37 flow around a cylinder, P-45 flow around a cylinder, P-45 flow around a cylinder, P-45 flow Type utility, U-91 for class, P-36 for mat keyword, U-103 fourth keyword entry, U-114, U-115 fvc class, P-36 fvc member function curl, P-37 datt, P-37 datt, P-37 datt, P-37 flowType utility, U-92 datt, P-37 for class, P-36 fvm member function datt, P-37 sup, P-37 fvm class, P-36 fvm class, P-36 fvm class, P-36 fvm class, P-39 fvmatk evword, U-103 for all P-24 environment variable, U-139		
finiteVolumeMethod class, P-36 firstTime menu entry, U-131 firstTime keyword, U-107 fixed keyword entry, U-108 fixedGradient boundary condition, U-152 fixedValue boundary condition, U-152 flattenMesh utility, U-91 flow free surface, U-57 laminar, U-19 steady, turbulent, P-54 supersonic, P-59 turbulent, U-20 flow around a cylinder, P-45 flow Over backward step, P-54 flow over backward step, P-54 flowType utility, U-92 fluentInterface directory, U-177 fluentMesh ToFoam utility, U-90, U-161 fluxCorrecteVelocity boundary condition, U-153 fluxRequired keyword, U-100 OpenFOAM cases, U-101 FOAM_RUN environment variable, U-101, U-139 Foam Utilities menu, U-22, U-33, U-34 foamCorrectVrt script/alias, U-166 foamDataToFluent utility, U-92 foamToFile keyword, U-103 foamInfoExec utility, U-94 foamJob script/alias, U-184 foamJop script/alias, U-185 foamMeshToFluent utility, U-92 foamToFieldview9 utility, U-92 foamUser library, U-81		
firstTime menu entry, U-131 firstTime keyword, U-107 fixed keyword entry, U-108 fixedGradient boundary condition, U-152 fixedWalue boundary condition, U-152 flattenMesh utility, U-91 flow free surface, U-57 laminar, U-19 steady, turbulent, P-54 supersonic, P-59 turbulent, U-20 flow around a cylinder, P-45 flow over backward step, P-54 flowToype utility, U-92 fluxCorrectedVelocity boundary condition, U-153 fluxRequired keyword, U-110 OpenFOAM cases, U-101 FOAM.RUN environment variable, U-101, U-139 roam Utilities menu, U-22, U-33, U-34 foamCorrectVrt script/alias, U-166 foamDataToFluent utility, U-94 FoamFile keyword, U-103 foamInfoExec utility, U-94 foamJob script/alias, U-184 foamLog script/alias, U-184 foamLog script/alias, U-184 foamLog script/alias, U-184 foamLog script/alias, U-185 foamMeshToFluent utility, U-92 foamToFieldview9 utility, U-92 foamToFX utility, U-92 foamToFX utility, U-92 foamToFX utility, U-92 foamToFX utility, U-92 foamToFieldview9 utility, U-92 foamToFie	,	
menu entry, U-131 firstTime keyword, U-107 fixed keyword entry, U-108 fixedGradient boundary condition, U-152 fixetValue boundary condition, U-152 flattenMesh utility, U-91 flow free surface, U-57 laminar, U-19 steady, turbulent, P-54 supersonic, P-59 turbulent, U-20 flow around a cylinder, P-45 flow Over backward step, P-54 flowType utility, U-92 fluentMeshToFoam utility, U-90, U-161 fluxCorrectedVelocity boundary condition, U-153 fluxRequired keyword, U-101 OpenFOAM cases, U-101 FOAM_RUN environment variable, U-101, U-139 Foam Utilities menu, U-22, U-33, U-34 foamCorrectVrt script/alias, U-166 foamDataToFluent utility, U-94 FoamFile keyword, U-103 foamInfoExec utility, U-94 foamLops script/alias, U-184 foamLog script/alias, U-185 foamMeshToFluent utility, U-92 foamToGMV utility, U-92 foamToGMV utility, U-92 foamToGMV utility, U-92 foamToFledview9 utility, U-92 foamToFledview9 utility, U-92 foamToFN utility, U-92 foamUser library, U-81 FoamX utility, U-139 FOAMX_NSYSTEM_CONFIG environment variable, U-139 FOAMX_SYSTEM_CONFIG environment variable, U-139 FOAMX_S		
firstTime keyword, U-107 fixed keyword entry, U-108 fixedGradient boundary condition, U-152 fixedWalue boundary condition, U-152 flattenMesh utility, U-91 flow free surface, U-57 laminar, U-19 steady, turbulent, P-54 supersonic, P-59 turbulent, U-20 flow around a cylinder, P-45 flow Type utility, U-92 fluentheface directory, U-177 fluentMeshToFoam utility, U-90, U-161 fluxCorrectedVelocity boundary condition, U-153 fluxRequired keyword, U-110 OpenFOAM cases, U-101 FOAM.RUN environment variable, U-101, U-139 Foam Utilities menu, U-22, U-33, U-34 foamCorrectVrt script/alias, U-166 foamDataToFluent utility, U-94 foamDataToFluent utility, U-94 foamJoats of lend tuility, U-94 foamJob script/alias, U-184 foamLog script/alias, U-185 foamWeshToFluent utility, U-92 foamToFieldview9 utility, U-92 foam		
fixed keyword entry, U-108 fixedGradient boundary condition, U-152 fixedValue boundary condition, U-152 flattenMesh utility, U-91 flow free surface, U-57 laminar, U-19 steady, turbulent, P-54 supersonic, P-59 turbulent, U-20 flow around a cylinder, P-45 flow over backward step, P-54 flowType utility, U-92 fluentInterface directory, U-177 fluentMeshToFoam utility, U-90, U-161 fluxCorrectedVelocity boundary condition, U-153 fluxRequired keyword, U-110 OpenFOAM cases, U-101 FOAM.RUN environment variable, U-101, U-139 Foam Utilities menu, U-22, U-33, U-34 foamCorrectVrt script/alias, U-186 foamDataToFluent utility, U-94 FoamFile keyword, U-103 foamInfoExec utility, U-94 foamJob script/alias, U-184 foamLog script/alias, U-184 foamLog script/alias, U-185 foamMeshToFluent utility, U-92 foamToFieldview9 utility, U-92 foamUser dictionary, U-62, U-101, U-109, U-110 fvSchemes menu entry, U-53		
keyword entry, U-108 fixedGradient boundary condition, U-152 fixedValue boundary condition, U-152 flattenMesh utility, U-91 flow free surface, U-57 laminar, U-19 steady, turbulent, P-54 supersonic, P-59 turbulent, U-20 flow around a cylinder, P-45 flow Type utility, U-92 fluxCorrectedVelocity boundary condition, U-153 fluxRequired keyword, U-110 OpenFOAM cases, U-101 FOAM.X.USER.CONFIG environment variable, U-139 Foam/Client.cfg file, U-124, U-138 foreground process, U-28 format keyword, U-103 fourth keyword entry, U-114, U-115 fvc class, P-36 fvc member function cur1, P-37 ddtt, P-37 ddtt, P-37 ddtt, P-37 ddtt, P-37 grad, P-37 laplacian, P-37 spcFadGrad, P-37		9 ,
fixedGradient boundary condition, U-152 fixedValue boundary condition, U-152 flattenMesh utility, U-91 flow free surface, U-57 laminar, U-19 steady, turbulent, P-54 supersonic, P-59 turbulent, U-20 flow around a cylinder, P-45 flowType utility, U-92 fluentMeshToFoam utility, U-90, U-161 fluxCorrectedVelocity boundary condition, U-153 fluxRequired keyword, U-110 OpenFOAM cases, U-101 FOAM.RUN environment variable, U-139 Foam Xariable, U-139 Foam Variable, U-139 Foam V		
boundary condition, U-152 fixedValue boundary condition, U-152 flattenMesh utility, U-91 flow free surface, U-57 laminar, U-19 steady, turbulent, P-54 supersonic, P-59 turbulent, U-20 flow around a cylinder, P-45 flow Type utility, U-92 fluxCorrectedVelocity boundary condition, U-153 fluxRequired keyword, U-100 CopenFOAM cases, U-101 FOAM.RUN environment variable, U-139 foamDebugSwitches utility, U-94 foamDataToFluent utility, U-94 foamDob script/alias, U-184 foamLog script/alias, U-184 foamLog script/alias, U-184 foamCorrectidity, U-92 foamToFieldview9 utility, U-92 foamToFieldview9 utility, U-92 foamToFoMW utility, U-92 foamToFMW utility, U-92 foamToFoMW utility, U-92 foamToFieldview9 utility, U-92 foamToFomWort intility, U-92 foamToFomWort intility, U-92 foamToFieldview9 utility, U-92 foamToFomWort intility, U-92 foamToFieldview9 utility, U-92 foamUser library U-81 environment variable, U-138 foremxClient.cfg file, U-124, U-138 foamXclient.cfg file, U-124, U-138 foremxClient.cfg file, U-103 foremaxClient.cfg file, U-103 format keyword, U-103		
fixedValue boundary condition, U-152 flattenMesh utility, U-91 flow free surface, U-57 laminar, U-19 steady, turbulent, P-54 supersonic, P-59 turbulent, U-20 flow around a cylinder, P-45 flow over backward step, P-54 flowType utility, U-92 fluentInterface directory, U-177 fluentMeshToFoam utility, U-90, U-161 fluxCorrectedVelocity boundary condition, U-153 fluxRequired keyword, U-110 OpenFOAM cases, U-101 FOAM.RUN environment variable, U-101, U-139 Foam Utilities menu, U-22, U-33, U-34 foamDataToFluent utility, U-94 foamJob script/alias, U-184 foamCog script/alias, U-185 foamMeshToFluent utility, U-92 foamToEMV utility, U-92 foamToFieldview9 utility, U-92 foamToToTK utility, U-92 foamToMV utility, U-92 foamUser library, U-81 FOAMX.USER.CONFIG environment variable, U-139 environment variable, U-139 formX.UseR.CONFIG environment variable, U-139 formxClient.cfg file, U-124, U-138 format keyword, U-103 format keyword, U-103 format keyword, util03 format keyword entry, U-114, U-115 fvc class, P-36 fvc member function cur1, P-37 ddt, P-37 ddt, P-37 grad, P-37 sugGradCorrection, P-37 sqcradGrad, P-37 synGradCorrection, P-37 sqcradGrad, P-37 fvm class, P-36 fvm member function d2dt2, P-37 ddt, P-37 fvm class, P-36 fvm member function d2dt2, P-37 ddt, P-37 fvm class, P-33 fvm dat, P-37 Su, P-37 fvm class, P-36 fvm member function d2dt2, P-37 ddt, P-37 fvm class, P-36 fvm member function d2dt2, P-37 ddt, P-37 fvm class, P-36 fvm member function d2dt2, P-37 ddt, P-37 fvm class, P-36 fvm member function d2dt2, P-37 ddt, P-37 fvm class, P-36 fvm member function d2dt2, P-37 fvm class, P-36 fvm member function d2dt2, P-37 ddt, P-37 fvm class, P-36 fvm member function d2dt2, P-37 fvm class, P-36 fvm member function d2dt2, P-37 ddt, P-37 fvm class, P-36 fvm environment variable, U-101, U-109 fvm class, P-36 fvm environment variable, U-103 format keyword, U-103 format keyword, U-103 format keyword, U-103 format key		
boundary condition, U-152 flattenMesh utility, U-91 flow free surface, U-57 laminar, U-19 steady, turbulent, P-54 supersonic, P-59 turbulent, U-20 flow around a cylinder, P-45 flow Type utility, U-92 fluentInterface directory, U-177 fluentMeshToFoam utility, U-90, U-161 fluxCorrectedVelocity boundary condition, U-153 fluxRequired keyword, U-110 OpenFOAM cases, U-101 FOAM.RUN environment variable, U-101, U-139 foamDataToFluent utility, U-92 foamToEMV utility, U-92 foamToFieldview9 utility, U-92 foamToFomW utility, U-92 foamToMV utility, U-92 foamToMV utility, U-92 foamToMV utility, U-92 foamOvered flow around a cylinder, P-45 format keyword, U-103 foreground process, U-28 format keyword, U-103 fourth keyword entry, U-114, U-115 fvc class, P-36 fvc member function curl, P-37 ddt, P-37 ddt, P-37 ddt, P-37 grad, P-37 sugGrad, P-37 sugGrad, P-37 sugGradGorrection, P-37 sugFradGorrection, P-37 supp. P-37 ddt, P-37 ddt, P-37 ddt, P-37 fvm class, P-36 fvm member function d2dt2, P-37 ddt, P-37 ddt, P-37 supp. P-37 fvm class, P-36 fvMesh class, P-36 fvMesh class, P-36 fvMesh class, P-36 fvMesh class, P-36 fvSchemes dictionary, U-62, U-101, U-109, U-110 fvSchemes library, U-81	,	
flattenMesh utility, U-91 flow free surface, U-57 laminar, U-19 steady, turbulent, P-54 supersonic, P-59 turbulent, U-20 flow around a cylinder, P-45 flow Type utility, U-92 fluentInterface directory, U-177 fluentMeshToFoam utility, U-90, U-161 fluxCorrectedVelocity boundary condition, U-153 fluxRequired keyword, U-110 OpenFOAM cases, U-101 FOAM.RUN environment variable, U-101, U-139 Foam Utilities menu, U-22, U-33, U-34 foamCorrectVrt script/alias, U-166 foamDataToFluent utility, U-94 foamJob script/alias, U-185 foamMeshToFoluent utility, U-94 foamJob script/alias, U-185 foamMeshToFluent utility, U-92 foamToFieldview9 utility, U-92 foamToFieldview9 utility, U-92 foamToFoMV ttility, U-92 foamToMV utility, U-92 foamToMV ttility, U-92 foamToMV ttility, U-92 foamUser library, U-81 FoamACCient.cfg file, U-124, U-138 foreground process, U-28 foamtkeyword, U-103 fourth keyword entry, U-114, U-115 fvc class, P-36 fvc member function cur1, P-37 ddt, P-37 ddt, P-37 sugrad, P-37 sugrad, P-37 sugrad, P-37 sugrad, P-37 fvm class, P-36 fvm member function ddate, P-37 ddt, P-37 ddt, P-37 ddt, P-37 ddt, P-37 sugradGrad, P-37 fvm class, P-36 fvm member function ddate, P-37 ddt, P-37 ddt, P-37 sugradGrad, P-37 fvm class, P-36 fvm drive template class, P-36 fvMesh class, P-31 fvSchemes dictionary, U-62, U-101, U-109, U-110 fvSchemes menu entry, U-53		
flow free surface, U-57 laminar, U-19 steady, turbulent, P-54 supersonic, P-59 turbulent, U-20 flow around a cylinder, P-45 flow over backward step, P-54 fluentInterface directory, U-177 fluentMeshToFoam utility, U-90, U-161 fluxCorrectedVelocity boundary condition, U-153 fluxRequired keyword, U-110 OpenFOAM cases, U-101 FOAM_RUN environment variable, U-101, U-139 Foam Utilities menu, U-22, U-33, U-34 foamCorrectVrt script/alias, U-166 foamDataToFluent utility, U-94 FoamFile keyword, U-103 foamInfoExec utility, U-94 foamLog script/alias, U-185 foamMeshToFluent utility, U-92 foamToEnsight utility, U-92 foamToGMV utility, U-92 foamToGMV utility, U-92 foamToToTK utility, U-92 foamToToTK utility, U-92 foamToToTK utility, U-92 foamToToTK utility, U-92 foamToToTX utility, U-92 foamToToTX utility, U-92 foamOuse library, U-81 format keyword, U-103 fourth keyword entry, U-114, U-115 fvc class, P-36 fvc member function cur1, P-37 ddt, P-37 ddt, P-37 grad, P-37 sqrGradGrad, P-37 syrGradGrad, P-37 fvm class, P-36 fvm member function ddt, P-37 spraddt, P-37 sqrGradGrad, P-37 fvm class, P-36 fvm member function fourut, U-114, U-115 fvc class, P-36 fvc member function fourl, P-37 grad, P-37 sqrGradCorrection, P-37 sqrGradGrad, P-37 fvm class, P-36 fvm member function ddt, P-37 sqrd, P-37 sqrGradGrad, P-37 fvm class, P-36 fvm member function ddt, P-37 sqrd, P-37 sqrGradGrad, P-37 sqrGradGrad, P-37 fvm class, P-36 fvm member function ddt, P-37 sqrd, P-37 sqrd, P-37 sqrdadGrad, P-37 sqrGradGrad, P-37 fvm class, P-36 fvm member function ddt, P-37 sqrd, P-37 sqrd, P-37 sqrd, P-37 sqrd, P-37 sqrdadgrad, P-37 sqrdadGrad, P-37 fvm class, P-36 fvm member function ddt, P-37 sqrd,		
free surface, U-57 laminar, U-19 steady, turbulent, P-54 supersonic, P-59 turbulent, U-20 flow around a cylinder, P-45 flow over backward step, P-54 flow Type utility, U-92 fluentherface directory, U-177 fluentMeshToFoam utility, U-90, U-161 fluxCorrectedVelocity boundary condition, U-153 fluxRequired keyword, U-110 OpenFOAM cases, U-101 FOAM.RUN environment variable, U-101, U-139 Foam Utilities menu, U-22, U-33, U-34 foamCorrectVrt script/alias, U-166 foamDataToFluent utility, U-94 foamJob script/alias, U-185 foamMeshToFluent utility, U-92 foamToEnsight utility, U-92 foamToGMV utility, U-92 foamToWrK utility, U-92 foamToWrK utility, U-92 foamToToTK utility, U-92 foamToToTTK utility, U-92 foamToToTK utility, U-93 foamUse library U-81 formtx keyword, U-110 keyword entry, U-114, U-113 formth keyword, U-114, U-115 foc class, P-36 fvc member function cur1, P-37 ddt, P-37 ddt, P-37 foatty, P-37 foatty, P-37 form Lesyword, U-101 fvc class, P-36 fvc member function cur1, P-37 ddt, P-37 foatty, P-37 foatty, P-37 foatty, P-37 form Lesyword, U-103 fvc member function cur1, P-37 foat Cass, P-36 fvc member function cur1, P-37 foat Cass, P-36 fvc member function cur1, P-37 foat Cass, P-36 fvc member function cur1,		9 , ,
laminar, U-19 steady, turbulent, P-54 supersonic, P-59 turbulent, U-20 flow around a cylinder, P-45 flowType utility, U-92 fluentInterface directory, U-177 fluentMeshToFoam utility, U-90, U-161 fluxCorrectedVelocity boundary condition, U-153 fluxRequired keyword, U-110 OpenFOAM cases, U-101 FOAM_RUN environment variable, U-101, U-139 Foam Utilities menu, U-22, U-33, U-34 foamCorrectVrt script/alias, U-166 foamDataToFluent utility, U-94 foamJob script/alias, U-185 foamMeshToFluent utility, U-92 foamToFieldview9 utility, U-92 foamToGMV utility, U-92 foamToVTK utility, U-92 foamToWTK utility, U-92 foamUser library, U-81 format keyword, U-103 fourth keyword entry, U-114, U-115 fvc class, P-36 fourth keyword entry, U-114, U-115 fvc class, P-36 fvc member function cur1, P-37 ddt, P-37 ddt, P-37 sufgrad, P-37 sufgradGrad, P-37 fvm class, P-36 fvm member function d2dt2, P-37 ddt, P-37 ddt, P-37 ddt, P-37 fvm class, P-36 fvm member function d2dt2, P-37 ddt, P-37 ddt, P-37 fvm class, P-36 fvm member function d2dt2, P-37 ddt, P-37 fvm class, P-36 fvm member function d2dt2, P-37 ddt, P-37 fvm class, P-36 fvm member function five member function d2dt2, P-37 ddt, P-37 fvm class, P-36 fvm member function d2dt2, P-37 fvm class, P-36 fvm member function fvx class, P-36 fvm class, P-36 fvm member function d2dt2, P-37 fvm class, P-36 fvm member function five class, P-36 fvm member function d2dt2, P-37 fvm class, P-36 fvm class, P-36 fvm member function d2dt2, P-37 fvm class, P-36 fvm class, P-36 fvm member function d2dt2, P-37 fvm class, P-36 fvm class, P-36 fvm class, P-36 fvc class, P-37 fvm class, P-36 fvm member function d2dt2, P-37 ddt, P-37 fvm class, P-36 fvm class, P-36 fvm class, P-36 fvm class, P-37 fvm class, P-36 fvm class, P-37 fvm class, P-37 fvm class, P-37 fvm class, P-37 fvm cl		9
steady, turbulent, P-54 supersonic, P-59 turbulent, U-20 flow around a cylinder, P-45 flow Over backward step, P-54 flow Type utility, U-92 fluentInterface directory, U-177 fluentMeshToFoam utility, U-90, U-161 fluxCorrectedVelocity boundary condition, U-153 fluxRequired keyword, U-110 OpenFOAM cases, U-101 FOAM_RUN environment variable, U-101, U-139 Foam Utilities menu, U-22, U-33, U-34 foamCorrectVrt script/alias, U-166 foamDataToFluent utility, U-94 foamJob script/alias, U-185 foamMeshToFluent utility, U-94 foamToGMV utility, U-92 foamToGMV utility, U-92 foamToVTK utility, U-92 foamToVTK utility, U-92 foamToWIN supersonic, P-59 keyword entry, U-114, U-115 fvc class, P-36 fvc member function cur1, P-37 ddt, P-37 ddt, P-37 sugrad, P-37 sugrad, P-37 sugrad, P-37 sugradGrad, P-37 fvm class, P-36 fvm member function d2dt2, P-37 ddt, P-37 div, P-37 sugrad, P-37 sugradGrad, P-37 fvm class, P-36 fvm member function d2dt2, P-37 ddt, P-37 div, P-37 sugrad, P-37 sugradGrad, P-37 fvm class, P-36 fvm member function d2dt2, P-37 ddt, P-37 sugradGrad, P-37 fvm class, P-36 fvm member function d2dt2, P-37 fvm lasi, P-37 div, P-37 sugrad, P-37 fvm class, P-36 fvm member function d2dt2, P-37 ddt, P-37 fvm class, P-36 fvm class, P-36 fvm member function d2dt2, P-37 ddt, P-37 fvm class, P-36 fvm class, P-36 fvm member function d2dt2, P-37 ddt, P-37 fvm class, P-36 fvm member function d2dt2, P-37 d2t2, P-37 fvm class, P-36 fvm class, P-36 fvm member function d2dt2, P-37 d2t2, P-37 fvm class, P-36 fvm class, P-36 fvm member function d2dt2, P-37 d2t2 fvm class, P-36 fvm member function d2dt2, P-37 d2t2 fvm c		
supersonic, P-59 turbulent, U-20 flow around a cylinder, P-45 flow over backward step, P-54 flowType utility, U-92 fluentInterface directory, U-177 fluentMeshToFoam utility, U-90, U-161 fluxCorrectedVelocity boundary condition, U-153 fluxRequired keyword, U-110 OpenFOAM cases, U-101 FOAM_RUN environment variable, U-101, U-139 Foam Utilities menu, U-22, U-33, U-34 foamCorrectVrt script/alias, U-166 foamDataToFluent utility, U-94 foamJob script/alias, U-184 foamLog script/alias, U-185 foamMeshToFluent utility, U-92 foamToEnsight utility, U-92 foamToGMV utility, U-92 foamToGMV utility, U-92 foamToVTK utility, U-92 foamUser library, U-81 keyword entry, U-114, U-114 fvc class, P-36 fvc member function cur1, P-37 ddt, P-37 ddt, P-37 ddt, P-37 grad, P-37 grad, P-37 laplacian, P-37 sqradGrad, P-37 fvm class, P-36 fvm member function d2dt2, P-37 ddt, P-37 fvm class, P-36 fvm member function d2dt2, P-37 ddt, P-37 fvm class, P-36 fvm member function d2dt2, P-37 ddt, P-37 fvm class, P-36 fvm member function d2dt2, P-37 ddt, P-37 fvm class, P-36 fvm member function d2dt2, P-37 div, P-37 fvm class, P-36 fvm member function d2dt2, P-37 fvm class, P-36 fvm class, P-37 fvm class,		,
turbulent, U-20 flow around a cylinder, P-45 flow over backward step, P-54 flowType utility, U-92 fluentInterface directory, U-177 fluentMeshToFoam utility, U-90, U-161 fluxCorrectedVelocity boundary condition, U-153 fluxRequired keyword, U-110 OpenFOAM cases, U-101 FOAM_RUN environnment variable, U-101, U-139 Foam Utilities menu, U-22, U-33, U-34 foamCorrectVrt script/alias, U-166 foamDataToFluent utility, U-94 FoamFile keyword, U-103 foamInfoExec utility, U-94 foamJob script/alias, U-185 foamMeshToFluent utility, U-92 foamToGMV utility, U-92 foamToGMV utility, U-92 foamToGMV utility, U-92 foamToFiledview9 utility, U-92 foamToFiledview9 utility, U-92 foamToFILE keyword, U-103 foamUser library, U-81 fvc class, P-36 fvc member function cur1, P-37 ddt, P-37 ddt, P-37 laplacian, P-37 sqrGradCorrection, P-37 fvm class, P-36 fvm		
flow around a cylinder, P-45 flow over backward step, P-54 flowType utility, U-92 fluentInterface directory, U-177 fluentMeshToFoam utility, U-90, U-161 fluxCorrectedVelocity boundary condition, U-153 fluxRequired keyword, U-110 OpenFOAM cases, U-101 FOAM_RUN environment variable, U-101, U-139 Foam Utilities menu, U-22, U-33, U-34 foamCorrectVrt script/alias, U-166 foamDataToFluent utility, U-94 FoamFile keyword, U-103 foamInfoExec utility, U-94 foamJob script/alias, U-185 foamMeshToFluent utility, U-92 foamToGMV utility, U-92 foamUser library, U-81 fvc member function cur1, P-37 ddt, P-37 ddt, P-37 ddt, P-37 snGrad, P-37 snGradCorrection, P-37 sqrGradGrad, P-37 fvm class, P-36 fvm member function div, P-37 laplacian, P-37 sqrGrad, P-37 sqrGradGrad, P-37 fvm class, P-36 fvm member function div, P-37 laplacian, P-37 sqrGradGrad, P-37 sqrGradGrad, P-37 fvm class, P-36 fvm member function div, P-37 laplacian, P-37 sqrGradGrad, P-37 sqrGradGrad, P-37 fvm class, P-36 fvm ember function fvschemes function fvschemes function cur1, P-37 ddt, P-37 ddt, P-37 sqrdradGrad, P-37 sqrGradGrad, P-37 sqrGradCorrection, P-37 sqrGradC	*	
flow over backward step, P-54 flowType utility, U-92 fluentInterface directory, U-177 fluentMeshToFoam utility, U-90, U-161 fluxCorrectedVelocity boundary condition, U-153 fluxRequired keyword, U-110 OpenFOAM cases, U-101 FOAM_RUN environment variable, U-101, U-139 environment variable, U-101, U-139 foamCorrectVrt script/alias, U-166 foamDataToFluent utility, U-92, U-177 foamDebugSwitches utility, U-94 foamJob script/alias, U-184 foamCog script/alias, U-184 foamCog script/alias, U-184 foamToFluent utility, U-92 foamToGMV utility, U-92 foamToFNUTK utility, U-92 foamToVTK utility, U-92 foamUser library, U-81 cur1, P-37 ddt, P-37 ddt, P-37 foalt, P-37 sugradCara, P-37 fom class, P-36 form member function d2dt2, P-37 fom class, P-36 form member function d2dt2, P-37 ddt, P-37 fom class, P-36 form member function d2dt2, P-37 ddt, P-37 fom class, P-36 form class, P-37 form		· · · · · · · · · · · · · · · · · · ·
flowType utility, U-92 fluentInterface directory, U-177 fluentMeshToFoam utility, U-90, U-161 fluxCorrectedVelocity boundary condition, U-153 fluxRequired keyword, U-110 OpenFOAM cases, U-101 FOAM_RUN environment variable, U-101, U-139 Foam Utilities menu, U-22, U-33, U-34 foamCorrectVrt script/alias, U-166 foamDataToFluent utility, U-94 FoamFile keyword, U-103 foamInfoExec utility, U-94 foamJob script/alias, U-184 foamCog script/alias, U-185 foamMeshToFluent utility, U-92 foamToEnsight utility, U-92 foamToGMV utility, U-92 foamToGMV utility, U-92 foamToFNVTK utility, U-92 foamToVTK utility, U-92 foamUser library, U-81 ddt, P-37 ddt, P-37 fory grad, P-37 snGrad, P-37 snGradCorrection, P-37 sqrGradGrad, P-37 fwm class, P-36 fwm member function d2dt2, P-37 ddt, P-37 fwm class, P-36 fwm member function d2dt2, P-37 fwm class, P-37 sqrGradGrad, P-37 sqrGradGrad, P-37 fwm class, P-36 fwm member function d2dt2, P-37 fwm class, P-36 fwm member function d2dt2, P-37 fwm class, P-37 fwm class, P-36 fwm member function d2dt2, P-37 fwm class, P-37 fwm class, P-36 fwm member function d2dt2, P-37 fwm class, P-37 fwm class, P-36 fwm member function d2dt2, P-37 fwm class, P-37 fwm class, P-36 fwm class, P-36 fwm member function d2dt2, P-37 fwm class, P-37 fwm class, P-36 fwm class, P-36 fwm class, P-36 fwm class, P-36 fwm class, P-37 fwm class, P-3		
fluentInterface directory, U-177 ddt, P-37 fluentMeshToFoam utility, U-90, U-161 div, P-37 fluxCorrectedVelocity gGrad, P-37 boundary condition, U-153 grad, P-37 fluxRequired keyword, U-110 laplacian, P-37 OpenFOAM lsGrad, P-37 cases, U-101 snGradCorrection, P-37 FOAM_RUN snGradCorrection, P-37 environment variable, U-101, U-139 snGradCorrection, P-37 Foam Utilities menu, U-22, U-33, U-34 foamCorrectVrt script/alias, U-166 fvm class, P-36 foamDataToFluent utility, U-92, U-177 ddt, P-37 ddt, P-37 foamPobugSwitches utility, U-94 ddt, P-37 ddt, P-37 foamJob script/alias, U-184 Su, P-37 suSp, P-37 foamLog script/alias, U-185 SuSp, P-37 suSp, P-37 foamMeshToFluent utility, U-92 fvMatrix template class, P-36 foamToEnsight utility, U-92 fvSchemes foamToGMV utility, U-92 dictionary, U-62, U-101, U-109, U-110 fvSchemes dictionary, U-62, U-101, U-109, U-110 fvSchemes menu entry, U-53	* /	
fluentMeshToFoam utility, U-90, U-161 fluxCorrectedVelocity boundary condition, U-153 fluxRequired keyword, U-110 OpenFOAM cases, U-101 FOAM_RUN environment variable, U-101, U-139 Foam Utilities menu, U-22, U-33, U-34 foamCorrectVrt script/alias, U-166 foamDataToFluent utility, U-92 foamJob script/alias, U-185 foamMeshToFluent utility, U-94 foamCospiely dias, U-185 foamMeshToFluent utility, U-90, U-177 foamDebug script/alias, U-185 foamMeshToFluent utility, U-92 foamToEnsight utility, U-92 foamToGMV utility, U-92 foamToFINGMV ut		,
fluxCorrectedVelocity boundary condition, U-153 fluxRequired keyword, U-110 OpenFOAM cases, U-101 FOAM_RUN environment variable, U-101, U-139 Foam Utilities menu, U-22, U-33, U-34 foamCorrectVrt script/alias, U-166 foamDataToFluent utility, U-94 FoamFile keyword, U-103 foamJob script/alias, U-184 foamLog script/alias, U-185 foamMeshToFluent utility, U-92 foamToEnsight utility, U-92 foamToGMV utility, U-92 foamToGMV utility, U-92 foamToVTK utility, U-92 foamUser library, U-81 gGrad, P-37 grad, P-37 grad, P-37 snGrad, P-37 snGrad, P-37 snGrad, P-37 snGrad, P-37 formGradGrad, P-37 formCass, P-36 form member function d2dt2, P-37 ddt, P-37 ddt, P-37 form member function d2dt2, P-37 div, P-37 su, P-37 fowMatrix template class, P-36 fofMesh class, P-36 fofMesh class, P-31 fvSchemes dictionary, U-62, U-101, U-109, U-110 fvSchemes menu entry, U-53		
boundary condition, U-153 fluxRequired keyword, U-110 OpenFOAM cases, U-101 FOAM_RUN environment variable, U-101, U-139 Foam Utilities menu, U-22, U-33, U-34 foamCorrectVrt script/alias, U-166 foamDataToFluent utility, U-94 FoamFile keyword, U-103 foamIog script/alias, U-184 foamLog script/alias, U-185 foamMeshToFluent utility, U-92 foamToFieldview9 utility, U-92 foamToGMV utility, U-92 foamToVTK utility, U-92 foamToVTK utility, U-92 foamUser library, U-81 grad, P-37 laplacian, P-37 snGradCorrection, P-37 sufficiently is snGradCorrection, P-37 sufficiently in Leas, P-37 ddt, P-37 ddt, P-37 ddt, P-37 ddt, P-37 sufficiently in Laplacian, P-37 sufficiently i		
fluxRequired keyword, U-110 OpenFOAM cases, U-101 FOAM_RUN environment variable, U-101, U-139 Foam Utilities menu, U-22, U-33, U-34 foamCorrectVrt script/alias, U-166 foamDataToFluent utility, U-94 FoamFile keyword, U-103 foamInfoExec utility, U-94 foamJob script/alias, U-185 foamMeshToFluent utility, U-90, U-177 foamToEnight utility, U-92 foamToGMV utility, U-92 foamToVTK utility, U-92 foamToVTK utility, U-92 foamUser library, U-81 laplacian, P-37 snGradCorrection, P-37 sqrGradGrad, P-37 fvm class, P-36 fvm member function d2dt2, P-37 ddt, P-37 ddt, P-37 ddt, P-37 SuSp, P-37 suSp, P-37 fvMesh class, P-36 fvMesh class, P-36 fvMesh class, P-31 fvSchemes dictionary, U-62, U-101, U-109, U-110 fvSchemes menu entry, U-53	•	
OpenFOAM cases, U-101 FOAM_RUN environment variable, U-101, U-139 Foam Utilities menu, U-22, U-33, U-34 foamCorrectVrt script/alias, U-166 foamDataToFluent utility, U-94 FoamFile keyword, U-103 foamlog script/alias, U-185 foamMeshToFluent utility, U-92 foamToEnight utility, U-92 foamToGMV utility, U-92 foamToFIVTK utility, U-92 foamUser library, U-81 1sGrad, P-37 snGradCorrection, P-37 sqrGradGrad, P-37 fvm class, P-36 fvm member function d2dt2, P-37 ddt, P-37 ddt, P-37 ddt, P-37 SuSp, P-37 SuSp, P-37 fvMatrix template class, P-36 fvMesh class, P-31 fvSchemes dictionary, U-62, U-101, U-109, U-110 fvSchemes menu entry, U-53	,	9 /
cases, U-101 FOAM_RUN environment variable, U-101, U-139 Foam Utilities menu, U-22, U-33, U-34 foamCorrectVrt script/alias, U-166 foamDataToFluent utility, U-94 FoamFile keyword, U-103 foamInfoExec utility, U-94 foamJob script/alias, U-184 foamLog script/alias, U-185 foamMeshToFluent utility, U-90 foamToEnsight utility, U-92 foamToGMV utility, U-92 foamToGMV utility, U-92 foamToVTK utility, U-92 foamUser library, U-81 snGrad, P-37 snGradCorrection, P-37 sqradGrad, P-37 fvm class, P-36 fvm member function d2dt2, P-37 ddt, P-37 ddt, P-37 ddt, P-37 suSp, P-37 suSp, P-37 fvMatrix template class, P-36 fvMesh class, P-31 fvSchemes dictionary, U-62, U-101, U-109, U-110 fvSchemes menu entry, U-53		• '
FOAM_RÜN environment variable, U-101, U-139 Foam Utilities menu, U-22, U-33, U-34 foamCorrectVrt script/alias, U-166 foamDataToFluent utility, U-92 foamFile keyword, U-103 foamInfoExec utility, U-94 foamJob script/alias, U-184 foamLog script/alias, U-185 foamMeshToFluent utility, U-90, U-177 foamToEnsight utility, U-92 foamToGMV utility, U-92 foamToGMV utility, U-92 foamToVTK utility, U-92 foamUser library, U-81 snGradCorrection, P-37 sqrGradGrad, P-37 fwm class, P-36 fwm member function d2dt2, P-37 ddt, P-37 ddt, P-37 laplacian, P-37 suSp, P-37 suSp, P-37 fwMatrix template class, P-36 fwMesh class, P-31 fvSchemes dictionary, U-62, U-101, U-109, U-110 fvSchemes menu entry, U-53	OpenFOAM	*
environment variable, U-101, U-139 Foam Utilities menu, U-22, U-33, U-34 foamCorrectVrt script/alias, U-166 foamDataToFluent utility, U-92, U-177 foamDebugSwitches utility, U-94 foamInoExec utility, U-94 foamJob script/alias, U-184 foamLog script/alias, U-185 foamMeshToFluent utility, U-90, U-177 foamToEnsight utility, U-92 foamToGMV utility, U-92 foamToVTK utility, U-92 foamUser library, U-81 sqrGradGrad, P-37 fwm class, P-36 fwm member function d2dt2, P-37 ddt, P-37 div, P-37 laplacian, P-37 suSp, P-37 fwMatrix template class, P-36 fwMesh class, P-36 fwSchemes dictionary, U-62, U-101, U-109, U-110 fvSchemes menu entry, U-53		,
Foam Utilities menu, U-22, U-33, U-34 foamCorrectVrt script/alias, U-166 foamDataToFluent utility, U-92, U-177 foamDebugSwitches utility, U-94 foamInoExec utility, U-94 foamJob script/alias, U-184 foamLog script/alias, U-185 foamMeshToFluent utility, U-90, U-177 foamToEnsight utility, U-92 foamToGMV utility, U-92 foamToVTK utility, U-92 foamUser library, U-81 fvm class, P-36 fvm member function d2dt2, P-37 ddt, P-37 div, P-37 SuSp, P-37 suSp, P-37 foMatrix template class, P-36 fvMesh class, P-36 fvMesh class, P-36 fvMesh class, P-31 fvSchemes dictionary, U-62, U-101, U-109, U-110 fvSchemes menu entry, U-53		
foamCorrectVrt script/alias, U-166 foamDataToFluent utility, U-92, U-177 foamDebugSwitches utility, U-94 FoamFile keyword, U-103 foamInfoExec utility, U-94 foamJob script/alias, U-184 foamLog script/alias, U-185 foamMeshToFluent utility, U-90, U-177 foamToEnsight utility, U-92 foamToGMV utility, U-92 foamToVTK utility, U-92 foamUser library, U-81 fome MeshToFluent utility, U-92 foamToFieldview9 foamToVTK utility, U-92 foamToFluent utility, U-92 foamToFluent utility, U-92 foamToFluent utility, U-92 foamToVTK utility, U-92 foamToVTK utility, U-92 foamToVTK utility, U-92 foamUser free function d2dt2, P-37 ddt, P-37 ddt, P-37 suSp, P-37 foamLatic template class, P-36 forMesh class, P-36 fvSchemes dictionary, U-62, U-101, U-109, U-110 fvSchemes menu entry, U-53		- · · · · · · · · · · · · · · · · · · ·
foamDataToFluent utility, U-92, U-177 foamDebugSwitches utility, U-94 FoamFile keyword, U-103 foamInfoExec utility, U-94 foamJob script/alias, U-184 foamLog script/alias, U-185 foamMeshToFluent utility, U-90, U-177 foamToEnsight utility, U-92 foamToGMV utility, U-92 foamToGMV utility, U-92 foamToVTK utility, U-92 foamUser library, U-81 ddt, P-37 ddt, P-37 ddt, P-37 susp, P-37 fomVatrix template class, P-36 fvMesh class, P-36 fvSchemes dictionary, U-62, U-101, U-109, U-110 fvSchemes menu entry, U-53		,
foamDebugSwitches utility, U-94 FoamFile keyword, U-103 foamInfoExec utility, U-94 foamJob script/alias, U-184 foamLog script/alias, U-185 foamMeshToFluent utility, U-90, U-177 foamToEnsight utility, U-92 foamToFieldview9 utility, U-92 foamToGMV utility, U-92 foamToVTK utility, U-92 foamUser library, U-81 ddt, P-37 div, P-37 foav, P-37 SuSp, P-37 fwMatrix template class, P-36 fwMesh class, P-31 fvSchemes dictionary, U-62, U-101, U-109, U-110 fvSchemes fvSchemes menu entry, U-53		
FoamFile keyword, U-103 div, P-37 foamInfoExec utility, U-94 laplacian, P-37 foamJob script/alias, U-184 Su, P-37 foamLog script/alias, U-185 SuSp, P-37 foamMeshToFluent utility, U-90, U-177 foamToEnsight utility, U-92 fvMesh class, P-31 foamToFieldview9 utility, U-92 fvSchemes foamToGMV utility, U-92 dictionary, U-62, U-101, U-109, U-110 foamToVTK utility, U-92 fvSchemes foamUser fvSchemes library, U-81 menu entry, U-53		
foamInfoExec utility, U-94 foamJob script/alias, U-184 foamLog script/alias, U-185 foamMeshToFluent utility, U-90, U-177 foamToEnsight utility, U-92 foamToGMV utility, U-92 foamToGMV utility, U-92 foamToVTK utility, U-92 foamToVTK utility, U-92 foamUser library, U-81 laplacian, P-37 SuSp, P-37 foMatrix template class, P-36 fvMesh class, P-31 fvSchemes dictionary, U-62, U-101, U-109, U-110 fvSchemes class, P-38 fvSchemes menu entry, U-53		
foamJob script/alias, U-184 foamLog script/alias, U-185 foamMeshToFluent utility, U-90, U-177 foamToEnsight utility, U-92 foamToGMV utility, U-92 foamToGMV utility, U-92 foamToVTK utility, U-92 foamUser library, U-81 Su, P-37 SuSp, P-37 fvMatrix template class, P-36 fvMesh class, P-31 fvSchemes dictionary, U-62, U-101, U-109, U-110 fvSchemes class, P-38 fvSchemes menu entry, U-53	,	*
foamLog script/alias, U-185 foamMeshToFluent utility, U-90, U-177 foamToEnsight utility, U-92 foamToFieldview9 utility, U-92 foamToGMV utility, U-92 foamToVTK utility, U-92 foamToVTK utility, U-92 foamUser library, U-81 SuSp, P-37 fvMatrix template class, P-36 fvMesh class, P-31 fvSchemes dictionary, U-62, U-101, U-109, U-110 fvSchemes class, P-38 fvSchemes menu entry, U-53		laplacian, P-37
foamMeshToFluent utility, U-90, U-177 foamToEnsight utility, U-92 foamToFieldview9 utility, U-92 foamToGMV utility, U-92 foamToVTK utility, U-92 foamToVTK utility, U-92 foamUser library, U-81 foamToVTK utility, U-92 fvSchemes fvSchemes fvSchemes menu entry, U-53	± / /	
foamToEnsight utility, U-92 fvMesh class, P-31 foamToFieldview9 utility, U-92 fvSchemes foamToGMV utility, U-92 dictionary, U-62, U-101, U-109, U-110 foamToVTK utility, U-92 fvSchemes class, P-38 foamUser fvSchemes menu entry, U-53	9 1 /	SuSp, P-37
foamToFieldview9 utility, U-92 foamToGMV utility, U-92 foamToVTK utility, U-92 foamUser library, U-81 foamToVTK utility, U-92 fvSchemes class, P-38 fvSchemes menu entry, U-53		* '
foamToGMV utility, U-92 dictionary, U-62, U-101, U-109, U-110 foamToVTK utility, U-92 fvSchemes class, P-38 foamUser fvSchemes menu entry, U-53	0 0,	,
foamToVTK utility, U-92 fvSchemes class, P-38 foamUser fvSchemes menu entry, U-53		
foamUser fvSchemes library, U-81 menu entry, U-53		The state of the s
library, U-81 menu entry, U-53		*
FoamX fvSolution		
	FoamX	fvSolution

dictionary, U-101, U-117	host, U-20
C	browser, U-124
G	hThermo model, U-96, U-188
gambitToFoam utility, U-90, U-161 GAMG	I
keyword entry, U-118, U-119	I
Gamma	tensor member function, P-25
keyword entry, U-113	icoDyMFoam solver, U-87
Gamma differencing, P-38	icoErrorEstimate utility, U-94
gammaContactAngle	icoFoam solver, U-19, U-24, U-25, U-28, U-87
boundary condition, U-59	icoMomentError utility, U-94
Gauss	ideasToFoam utility, U-161
keyword entry, U-114	ideasUnvToFoam utility, U-90
Gauss's theorem, P-36	identities, see tensor, identities
GaussSeidel	identity, see tensor, identity
keyword entry, U-119	incompressible tools, U-95
General window panel, U-174	incompressibleLESmodels
general model, U-97	library, U-98
general	incompressiblePostProcessing
keyword entry, U-108	library, U-95
geometric-algebraic multi-grid, U-119	in compressible Transport Models
GeometricBoundaryField template class, P-32	library, P-55, U-99
geometricField <type> template class, P-32</type>	incompressibleTurbulenceModels
gGrad	library, P-55, U-97
fvc member function, P-37	index
global tools, U-95	notation, P-16, P-17
gmshToFoam utility, U-90	Info button, U-133
gnuplot	Information window panel, U-172
keyword entry, U-109, U-182	inhomogeneousMixture model, U-96, U-188
grad	inlet
fvc member function, P-37	boundary condition, P-71
(Grad Grad) squared, P-37	inletOutlet
gradient, P-37, P-40	boundary condition, U-153
Gauss scheme, P-40	inner product, see tensor, inner product
Gauss's theorem, U-53	insideCells utility, U-91
least square fit, U-53	instance keyword, U-103
least squares method, P-40, U-53	interFoam solver, U-88
surface normal, P-40	internalField keyword, U-106, U-135
gradSchemes keyword, U-110	interpolationScheme keyword, U-182
graphFormat keyword, U-109	interpolations tools, U-95
Gstream	interpolationSchemes keyword, U-110
library, U-96	inv
guldersLaminarFlameSpeed model, U-96	tensor member function, P-25
Н	isoOctane model, U-97
	т
hConstThermo model, U-97, U-187	J
hhuMixtureThermo model, U-96, U-188	janafThermo model, U-97, U-187
hierarchical	JAVA_HOME
keyword entry, U-83, U-84	environment variable, U-139
hMixtureThermo model, U-96, U-188	jplot
homogeneousMixture model, U-96, U-188	keyword entry, U-109, U-182

K	maxDeltaT, U-61
kappa keyword, U-191	mergeLevels, U-120
kEpsilon model, U-98	method, U-84
keyword	metisCoeffs, U-84
FoamFile, U-103	midPointAndFace, U-183
LESmodel, U-191	midPoint, U-183
adjustTimeStep, U-61	nFaces, U-144
agglomerator, U-119	nFinestSweeps, U-120
applicationClass, U-107	nGammaSubCycles, U-63
arc, U-155	nPostSweeps, U-120
blocks, U-22, U-156	nPreSweeps, U-120
block, U-155	numberOfSubdomains, U-84
boundaryField, U-106	n, U-84
boxToCell, U-60	object, U-103
cGamma, U-63	order, U-84
cacheAgglomeration, U-119	outputFormat, U-182
caseRoots, U-19	pRefCell, U-26, U-121
case, U-103	pRefValue, U-26, U-121
class, U-103	patchMap, U-168
cloud, U-183	patches, U-155, U-157
convertToMeters, U-155	pdRefCell, U-121
curve, U-183	pdRefValue, U-121
defaultFieldValues, U-60	physicalType, U-144, U-148
deltaT, U-108	preconditioner, U-117, U-118
delta, U-84, U-191	processorWeights, U-84
dimensions, U-106	purgeWrite, U-108
distributed, U-84, U-86	refGradient, U-152
divSchemes, U-110	referenceLevel, U-106, U-135
edgeGrading, U-157	regions, U-60
edges, U-155	relTol, U-54, U-117, U-118
endTime, U-25, U-107, U-108	roots, U-84, U-86
face, U-183	root, U-103
fieldValues, U-60	runTimeModifiable, U-109
fields, U-182	sampleSets, U-182
firstTime, U-107	simpleGrading, U-157
fluxRequired, U-110	smoother, U-120
format, U-103	snGradSchemes, U-110
gradSchemes, U-110	solvers, U-117
graphFormat, U-109	spline, U-155
instance, U-103	startFace, U-144
internalField, U-106, U-135	startFrom, U-25, U-107
interpolationSchemes, U-110	startTime, U-25, U-107
interpolationScheme, U-182	stopAt, U-107
kappa, U-191	thermoType, U-187
laplacianSchemes, U-110	timeFormat, U-108
latestTime, U-41	timePrecision, U-109
leastSquares, U-53	timeScheme, U-110
local, U-103	tolerance, U-54, U-117, U-118
manualCoeffs, U-84	topoSetSource, U-60
maxCo, U-61	turbulenceModel, U-191
	- 31 5 41 511 511 54 51, 0 101

turbulence, U-191
type, U-148
uniform, U-183
valueFraction, U-152
value, U-152
version, U-103
vertices, U-22, U-155
wallFunctionCoeffs, U-191
writeCompression, U-108
writeControl, U-25, U-61, U-108
writeFormat, U-55, U-108
writeInterval, U-25, U-35, U-108
writePrecision, U-108
<lesmodel>Coeffs, U-191</lesmodel>
<delta>Coeffs, U-191</delta>
<pre><turbulencemodel>Coeffs, U-191</turbulencemodel></pre>
keyword entry
CrankNicholson, U-116
CrossPowerLaw, U-61
DICGaussSeidel, U-119
DIC, U-119
DILU, U-119
Euler, U-116
FDIC, U-119
GAMG, U-118, U-119
Gamma, U-113
GaussSeidel, U-119
Gauss, U-114
MGridGen, U-119
MUSCL, U-113
Newtonian, U-61
PBiCG, U-118
PCG, U-118
QUICK, U-113, U-116
SFCD, U-113, U-116
UMIST, U-111
adjustableRunTime, U-61, U-108
arc, U-48, U-156
ascii, U-108
backward, U-116
binary, U-108
bounded, U-114, U-115
cellPointFace, U-182
cellPoint, U-182
cell, U-182
clockTime, U-108
compressed, U-108
corrected, $U-114$, $U-115$
cpuTime, U-108
cubicCorrected, U-116

 $\mathtt{cyclic},\, \textcolor{red}{U\text{-}150}$ diagonal, U-119 distance, U-183 empty, U-150 ${\tt faceAreaPair},\, {\color{red}U\textbf{-}119}$ fixed, U-108 fourth, U-114, U-115 general, U-108 gnuplot, U-109, U-182 hierarchical, U-83, U-84 jplot, U-109, U-182 latestTime, U-107 leastSquares, U-114 limitedCubic, U-113 limitedLinear, U-113 limited, U-114, U-115 linearUpwind, U-113, U-116 linear, U-113, U-116 line, U-156 manual, U-83, U-84 metis, U-83, U-84 midPoint, U-113 nextWrite, U-108 ${\tt noWriteNow},\, {\color{red}U\text{-}108}$ none, U-111, U-119 patch, U-150 polyLine, U-156 polySpline, U-156 processor, U-150 raw, U-109, U-182 runTime, U-35, U-108 scientific, U-108 simpleSpline, U-156 simple, U-83, U-84 skewLinear, U-113, U-116 smoothSolver, U-118 startTime, U-25, U-107 steadyState, U-116 symmetryPlane, U-150 $\mathtt{timeStep},\ U\text{-}25,\ U\text{-}35,\ U\text{-}108$ uncompressed, U-108 ${\tt uncorrected},\, U\text{-}114,\, U\text{-}115$ upwind, U-113, U-116 vanLeer, U-113 wall, U-150 wedge, U-150 writeControl, U-108 writeNow, U-107 xmgr, U-109, U-182

cubicCorrection, U-113

Index P-85 P-86 Index

xyz, U-183	cellDecompFiniteElement, U-95
x, U-183	cfdTools, U-95
y, U-183	chemistryModel, U-97
z, U-183	combustionThermophysicalModels, U-96
kill button, U-133	compressibleLESmodels, U-99
kivaToFoam utility, U-90	compressibleTurbulenceModels, U-98
Kronecker delta, P-21	dieselSpray, U-96
	dynamicMesh, U-95
${f L}$	edgeMesh, U-95
lagrangian	engine, U-96
library, U-96	errorEstimation, U-95
LAM	faceDecompFiniteElement, U-95
message passing interface, U-84	foamUser, U-81
MPI, U-84	
Lambda2 utility, U-92	incompressible LES models, U-98
LamBremhorstKE model, U-98	incompressiblePostProcessing, U-95
laminar model, U-97, U-98	incompressibleTransportModels, P-55, U-99
laminarFlameSpeedModels	incompressibleTurbulenceModels, P-55, U-97
library, U-96	lagrangian, U-96
laplaceFilter model, U-98	laminarFlameSpeedModels, U-96
Laplacian, P-38	liquids, U-97
laplacian, P-37	meshTools, U-95
laplacian	mico-2.3.13, U-96
fvc member function, P-37	mpich-1.2.4, U-96
fvm member function, P-37	pdf, U-97
laplacianFoam solver, U-87	primitive, P-23
laplacianSchemes keyword, U-110	randomProcesses, U-96
latestTime	sampling, U-95
keyword entry, U-107	shapeMeshTools, U-95
menu entry, U-131	specie, U-97
latestTime keyword, U-41	thermophysicalFunctions, U-97
LaunderGibsonRSTM model, U-98	thermophysical, U-187
LaunderSharmaKE model, U-98	triSurface, U-95
leastSquares	vtkFoam, U-171
keyword entry, U-114	zlib-1.2.1, U-96
leastSquares keyword, U-53	lid-driven cavity flow, U-19
LESdeltas	LienCubicKE model, U-98
library, U-98	LienCubicKELowRE model, U-98
LESfilters	LienLeschzinerLowRE model, U-98
library, U-98	limited
lesInterFoam solver, U-88	keyword entry, U-114, U-115
LESmodel keyword, U-191	limitedCubic
libraries, U-69	keyword entry, U-113
library	limitedLinear
Gstream, U-96	keyword entry, U-113
LESdeltas, U-98	line
LESfilters, U-98	keyword entry, U-156
ODE, U-95	linear
OpenFOAM, U-95	keyword entry, U-113, U-116
PVFoamReader, U-171	linearUpwind
basicThermophysicalModels, $U-96$	keyword entry, U-113, U-116

liquid	Property, U-174
electrically-conducting, P-69	Read Mesh&Fields, U-23, U-45, U-50
liquids	Refresh Case Browser, U-41
library, U-97	Source, U-28, U-175
lists, P-29	Wireframe, U-174
List <type> template class, P-29</type>	allTime, U-131
local keyword, U-103	blockMesh, U-22, U-33
locDynOneEqEddy model, U-98	firstTime, U-131
Lower and Upper Times text box, U-173	fvSchemes, U-53
lowReOneEqEddy model, U-99	latestTime, U-131
LRDDiffStress model, U-98	mapFields, U-34
LRR model, U-98	noTime, U-131
lsGrad	preProcessing, U-34
fvc member function, P-37	sample, U-55
3.5	mergeLevels keyword, U-120
\mathbf{M}	mergeMeshes utility, U-91
Mach utility, U-92	Mesh dictionary tree, U-23
mag	Mesh menu, U-50
tensor member function, P-25	mesh
magGradU utility, U-92	
magnetohydrodynamics, P-69	1-dimensional, U-145
magSqr	1D, U-145
tensor member function, P-25	2-dimensional, U-145
magU utility, U-36, U-92	2D, U-145
Make directory, U-73	axi-symmetric, U-145
make script/alias, U-71	basic, P-31
Make/files file, U-75	block structured, U-151
manual	decomposition, U-82
keyword entry, U-83, U-84	description, U-141
manualCoeffs keyword, U-84	finite volume, P-31
mapFields utility, U-34, U-40, U-43, U-56, U-90,	generation, U-151
U-167	grading, U-151, U-157
mapFields	grading, example of, P-54
menu entry, U-34	non-orthogonal, P-45
mapping	refinement, P-63
fields, U-167	resolution, U-33
matrices tools, U-95	specification, U-141
max	validity constraints, U-141
tensor member function, P-25	meshes tools, U-95
maxCo keyword, U-61	meshTools
maxDeltaT keyword, U-61	library, U-95
mechanicalProperties	message passing interface
dictionary, U-51	LAM, U-84
menu	MPICH, U-193
Class, U-129	method keyword, U-84
	metis
Color by, U-174	keyword entry, U-83, U-84
Foam Utilities, U-22, U-33, U-34	metisCoeffs keyword, U-84
Mesh, U-50	MGridGen
View, U-28, U-174, U-175	
menu entry	keyword entry, U-119
3D view Properties, U-27, U-174-U-176	mhdFoam solver, P-71, U-89

Index P-87 P-88 Index

mico-2.3.13	exponential, U-97
library, U-96	general, U-97
midPoint	guldersLaminarFlameSpeed, U-96
keyword entry, U-113	hConstThermo, U-97, U-187
midPoint keyword, U-183	hMixtureThermo, U-96, U-188
midPointAndFace keyword, U-183	hThermo, U-96, U-188
min	hhuMixtureThermo, U-96, U-188
tensor member function, P-25	homogeneousMixture, U-96, U-188
MINMOD differencing, P-38	inhomogeneousMixture, U-96, U-188
mirrorMesh utility, U-91	isoOctane, U-97
mixed	janafThermo, U-97, U-187
boundary condition, U-152	kEpsilon, U-98
mixedSmagorinsky model, U-98	laminar, U-97, U-98
mixtureAdiabaticFlameT utility, U-94	laplaceFilter, U-98
model	locDynOneEqEddy, U-98
APIfunctions, U-97	lowReOneEgEddy, U-99
BirdCarreau, U-99	mixedSmagorinsky, U-98
CrossPowerLaw, U-99	multiComponentMixture, U-96, U-188
DeardorffDiffStress, U-99	nDecane, U-97
LRDDiffStress, U-98	nDodecane, U-97
LRR, U-98	nHeptane, U-97
LamBremhorstKE, U-98	nOctane, U-97
LaunderGibsonRSTM, U-98	normal, U-97
LaunderSharmaKE, U-98	oneEqEddy, U-98, U-99
LienCubicKELowRE, U-98	perfectGas, U-97, U-187
LienCubicKE, U-98	pureMixture, U-96, U-188
LienLeschzinerLowRE, U-98	scaleSimilarity, U-98
NSRDSfunctions, U-97	simpleFilter, U-98
Newtonian, U-99	smoothDelta, U-98
NonlinearKEShih, U-98	specieThermo, U-97, U-187
PrandtlDelta, U-98	spectEddyVisc, U-98
QZeta, U-98	sutherlandTransport, U-97, U-188
RNGkEpsilon, U-98	uniform, U-97
RosinRammler, U-97	veryInhomogeneousMixture, U-96, U-188
Smagorinsky2, U-98	water, U-97
Smagorinsky, U-98, U-99	momentScalarError utility, U-94
SpalartAllmaras, U-98, U-99	moveDynamicMesh utility, U-91
anisotropicFilter, U-98	moveEngineMesh utility, U-91
chemistryModel, U-97	moveMesh utility, U-91
chemistrySolver, U-97	movingWallVelocity
chemkinMixture, U-96, U-188	boundary condition, U-153
constLaminarFlameSpeed, $U-96$	MPI
constTransport, U-97, U-188	LAM, U-84
cubeRootVolDelta, U-98	MPICH, U-193
diEthylEther, U-97	MPICH
diMethylEther, U-97	message passing interface, U-193
dieselMixture, U-96, U-188	MPI, U-193
dynMixedSmagorinsky, U-98	mpich-1.2.4
dynOneEqEddy, U-98, U-99	library, U-96
dynSmagorinsky, U-98	mshToFoam utility, U-90

multiComponentMixture model, U-96, U-188	Opacity text box, U-174
multigrid	OpenFOAM
geometric-algebraic, U-119	applications, U-69
multiphaseInterFoam solver, U-88	file format, U-102
MUSCL	libraries, U-69
keyword entry, U-113	OpenFOAM
My Jobs button, U-133	library, U-95
N	OpenFOAM file syntax
n keyword, U-84	//, U-102
nabla	operator
operator, P-27	scalar, P-28
name	vector, P-27
server, U-124	options file, U-73
nDecane model, U-97	order keyword, U-84
nDodecane model, U-97	Orientation Axes button, U-27
neighbour	outer product, see tensor, outer product
dictionary, U-144	outlet
netgenNeutralToFoam utility, U-90	boundary condition, P-71
Newtonian	outletInlet
	boundary condition, U-153
keyword entry, U-61	outputFormat keyword, U-182
Newtonian model, U-99 nextWrite	owner
	dictionary, U-143
keyword entry, U-108	Р
nFaces keyword, U-144	p field, U-25
nFinestSweeps keyword, U-120	paraFoam, U-26, U-171
nGammaSubCycles keyword, U-63	paraFoam utility, U-92
nHeptane model, U-97	parallel
nOctane model, U-97	running, U-82
non-orthogonal mesh, P-45	Parameters window panel, U-28, U-172, U-173
none	partialSlip
keyword entry, U-111, U-119	boundary condition, U-153
NonlinearKEShih model, U-98	boundary condition, 0-155
nonNewtonianIcoFoam solver, U-87	natch
	patch
normal model, U-97	boundary condition, U-150
noTime	boundary condition, U-150 patch
noTime menu entry, U-131	boundary condition, U-150 patch keyword entry, U-150
noTime menu entry, U-131 noWriteNow	boundary condition, U-150 patch keyword entry, U-150 patchAverage utility, U-93
noTime menu entry, U-131 noWriteNow keyword entry, U-108	boundary condition, U-150 patch keyword entry, U-150 patchAverage utility, U-93 Patches dictionary tree, U-23
noTime menu entry, U-131 noWriteNow keyword entry, U-108 nPostSweeps keyword, U-120	boundary condition, U-150 patch keyword entry, U-150 patchAverage utility, U-93 Patches dictionary tree, U-23 patches keyword, U-155, U-157
noTime menu entry, U-131 noWriteNow keyword entry, U-108 nPostSweeps keyword, U-120 nPreSweeps keyword, U-120	boundary condition, U-150 patch keyword entry, U-150 patchAverage utility, U-93 Patches dictionary tree, U-23 patches keyword, U-155, U-157 patchIntegrate utility, U-93
noTime menu entry, U-131 noWriteNow keyword entry, U-108 nPostSweeps keyword, U-120 nPreSweeps keyword, U-120 NSRDSfunctions model, U-97	boundary condition, U-150 patch keyword entry, U-150 patchAverage utility, U-93 Patches dictionary tree, U-23 patches keyword, U-155, U-157 patchIntegrate utility, U-93 patchMap keyword, U-168
noTime menu entry, U-131 noWriteNow keyword entry, U-108 nPostSweeps keyword, U-120 nPreSweeps keyword, U-120	boundary condition, U-150 patch keyword entry, U-150 patchAverage utility, U-93 Patches dictionary tree, U-23 patches keyword, U-155, U-157 patchIntegrate utility, U-93 patchMap keyword, U-168 patchTool utility, U-91
noTime menu entry, U-131 noWriteNow keyword entry, U-108 nPostSweeps keyword, U-120 nPreSweeps keyword, U-120 NSRDSfunctions model, U-97 numberOfSubdomains keyword, U-84	boundary condition, U-150 patch keyword entry, U-150 patchAverage utility, U-93 Patches dictionary tree, U-23 patches keyword, U-155, U-157 patchIntegrate utility, U-93 patchMap keyword, U-168 patchTool utility, U-91 PBiCG
noTime menu entry, U-131 noWriteNow keyword entry, U-108 nPostSweeps keyword, U-120 nPreSweeps keyword, U-120 NSRDSfunctions model, U-97 numberOfSubdomains keyword, U-84	boundary condition, U-150 patch keyword entry, U-150 patchAverage utility, U-93 Patches dictionary tree, U-23 patches keyword, U-155, U-157 patchIntegrate utility, U-93 patchMap keyword, U-168 patchTool utility, U-91 PBiCG keyword entry, U-118
noTime menu entry, U-131 noWriteNow keyword entry, U-108 nPostSweeps keyword, U-120 nPreSweeps keyword, U-120 NSRDSfunctions model, U-97 numberOfSubdomains keyword, U-84 O object keyword, U-103	boundary condition, U-150 patch keyword entry, U-150 patchAverage utility, U-93 Patches dictionary tree, U-23 patches keyword, U-155, U-157 patchIntegrate utility, U-93 patchMap keyword, U-168 patchTool utility, U-91 PBiCG keyword entry, U-118 PCG
noTime menu entry, U-131 noWriteNow keyword entry, U-108 nPostSweeps keyword, U-120 nPreSweeps keyword, U-120 NSRDSfunctions model, U-97 numberOfSubdomains keyword, U-84 O object keyword, U-103 objToVTK utility, U-91	boundary condition, U-150 patch keyword entry, U-150 patchAverage utility, U-93 Patches dictionary tree, U-23 patches keyword, U-155, U-157 patchIntegrate utility, U-93 patchMap keyword, U-168 patchTool utility, U-91 PBiCG keyword entry, U-118 PCG keyword entry, U-118
noTime menu entry, U-131 noWriteNow keyword entry, U-108 nPostSweeps keyword, U-120 nPreSweeps keyword, U-120 NSRDSfunctions model, U-97 numberOfSubdomains keyword, U-84 O object keyword, U-103 objToVTK utility, U-91 ODE	boundary condition, U-150 patch keyword entry, U-150 patchAverage utility, U-93 Patches dictionary tree, U-23 patches keyword, U-155, U-157 patchIntegrate utility, U-93 patchMap keyword, U-168 patchTool utility, U-91 PBiCG keyword entry, U-118 PCG keyword entry, U-118 pdf
noTime menu entry, U-131 noWriteNow keyword entry, U-108 nPostSweeps keyword, U-120 nPreSweeps keyword, U-120 NSRDSfunctions model, U-97 numberOfSubdomains keyword, U-84 O object keyword, U-103 objToVTK utility, U-91 ODE library, U-95	boundary condition, U-150 patch keyword entry, U-150 patchAverage utility, U-93 Patches dictionary tree, U-23 patches keyword, U-155, U-157 patchIntegrate utility, U-93 patchMap keyword, U-168 patchTool utility, U-91 PBiCG keyword entry, U-118 PCG keyword entry, U-118 pdf library, U-97
noTime menu entry, U-131 noWriteNow keyword entry, U-108 nPostSweeps keyword, U-120 nPreSweeps keyword, U-120 NSRDSfunctions model, U-97 numberOfSubdomains keyword, U-84 O object keyword, U-103 objToVTK utility, U-91 ODE	boundary condition, U-150 patch keyword entry, U-150 patchAverage utility, U-93 Patches dictionary tree, U-23 patches keyword, U-155, U-157 patchIntegrate utility, U-93 patchMap keyword, U-168 patchTool utility, U-91 PBiCG keyword entry, U-118 PCG keyword entry, U-118 pdf

Index P-89 P-90 Index

Pe utility, U-92	foreground, U-28
perfectGas model, U-97, U-187	processor
permutation symbol, P-20	boundary condition, U-151
physicalType keyword, U-144, U-148	processor
PISO	keyword entry, U-150
dictionary, U-26	processorN directory, U-84
plot3dToFoam utility, U-90	processorWeights keyword, U-84
pointField class, P-31	Property
pointField <type> template class, P-33</type>	menu entry, U-174
points	ptot utility, U-93
dictionary, U-143, U-154	pureMixture model, U-96, U-188
pointSet utility, U-91	purge button, U-133
polyBoundaryMesh class, P-31	purgeWrite keyword, U-108
polyDualMesh utility, U-90	PVFoamReader
polyLine	library, U-171
keyword entry, U-156	Q
polyMesh directory, U-101, U-143	Q utility, U-92
polyMesh class, P-31, U-141, U-143	QUICK
polyPatch class, P-31	keyword entry, U-113, U-116
polyPatchList class, P-31	QZeta model, U-98
polySpline	QZeta model, 0-30
keyword entry, U-156	\mathbf{R}
post-processing, U-171	R utility, U-93
post-processing	randomProcesses
paraFoam, U-171	library, U-96
postChannel utility, U-93	rasInterFoam solver, U-88
potentialFoam solver, P-46, U-87	raw
pow	keyword entry, U-109, U-182
tensor member function, P-25	Rcomponents utility, U-93
PrandtlDelta model, U-98	reactingFoam solver, U-89
preconditioner keyword, U-117, U-118	read button, U-133
pRefCell keyword, U-26, U-121	Read Mesh&Fields
pRefValue keyword, U-26, U-121	menu entry, U-23, U-45, U-50
preProcessing	reconstructPar utility, U-87, U-94
menu entry, U-34	reconstructParMesh utility, U-94
pressure waves	referenceLevel keyword, U-106, U-135
in liquids, P-64	refGradient keyword, U-152
pressureDirectedInletVelocity	refineMesh utility, U-91
boundary condition, U-153	Refresh Case Browser button, U-41
pressureInletVelocity	Refresh Case Browser
boundary condition, U-153	menu entry, U-41
pressureOutlet	Region window, U-28
boundary condition, P-65	regions keyword, U-60
pressureTransmissive	relative tolerance, U-118
boundary condition, U-153	relTol keyword, U-54, U-117, U-118
primitive	renumberMesh utility, U-91
library, P-23	Reset button, U-173
primitives tools, U-95	Reset Range button, U-28
process background, U-28, U-82	restart, U-41 Reynolds number, U-20, U-24

rhopSonicFoam solver, U-88
rhoSimpleFoam solver, U-88
rhoSonicFoam solver, U-88
rhoTurbFoam solver, U-88
rmdepall script/alias, U-77
RNGkEpsilon model, U-98
root keyword, U-103
roots keyword, U-84, U-86
RosinRammler model, U-97
rotateMesh utility, U-91
run
parallel, U-82
run directory, U-101
runFoamX script/alias, U-123-U-125
$runFoamXHB\ \mathrm{script/alias},\ U\text{-}123,\ U\text{-}124$
runTime
keyword entry, U-35, U-108
runTimeModifiable keyword, U-109
C
S
sammToFoam utility, U-90
sample utility, U-93, U-181
sample
menu entry, U-55
sampleSets keyword, U-182
sampleSurface utility, U-93
sampling
library, U-95
scalar, P-16
operator, P-28
scalar class, P-24
scalarField class, P-29
scalarTransportFoam solver, U-87
scale
tensor member function, P-25
scalePoints utility, U-165
scaleSimilarity model, U-98
scientific
keyword entry, U-108
script/alias
foamCorrectVrt, U-166
foamJob, U-184
foamLog, U-185
make, U-71
rmdepall, U-77
runFoamXHB, U-123, U-124
runFoamX, U-123-U-125
wclean, U-76
wmake, U-71
second time derivative, P-37
Seed window, U-177

```
setFields utility, U-59, U-60, U-90
settlingFoam solver, U-88
SFCD
    keyword entry, U-113, U-116
shape, U-156
shapeMeshTools
   library, U-95
simple
    keyword entry, U-83, U-84
simpleFilter model, U-98
simpleFoam solver, P-55, U-87
simpleGrading keyword, U-157
simpleSpline
    keyword entry, U-156
skew
    tensor member function, P-25
skewLinear
    keyword entry, U-113, U-116
slice class, P-31
    boundary condition, U-153
Smagorinsky model, U-98, U-99
Smagorinsky2 model, U-98
smapToFoam utility, U-92
smoothDelta model, U-98
smoother keyword, U-120
smoothSolver
    keyword entry, U-118
snGrad
    fvc member function, P-37
snGradCorrection
    fvc member function, P-37
snGradSchemes keyword, U-110
solidDisplacementFoam solver, U-51, U-89
{\sf solidEquilibriumDisplacementFoam\ solver,\ U-89}
solver
    XiFoam, U-89
    Xoodles, U-89
    blockMesh, P-47
    boundaryFoam, U-87
    bubbleFoam, U-88
    buoyantFoam, U-89
    buoyantSimpleFoam, U-89
    cavitatingFoam, U-88
    channelOodles, U-88
    coldEngineFoam, U-89
    coodles, U-88
    dieselEngineFoam, U-89
    dieselFoam, U-89
```

Selection Window window, U-27, U-172

Index P-91 P-92 Index

dnsFoam, U-88	sqr
electrostaticFoam, U-89	tensor member function, P-25
engineFoam, U-89	sqrGradGrad
financialFoam, U-89	fvc member function, P-37
icoDyMFoam, U-87	Standard Views window panel, U-174
icoFoam, U-19, U-24, U-25, U-28, U-87	Start Calculation button, U-35
interFoam, U-88	Start Calculation Now button, U-28
laplacianFoam, U-87	startFace keyword, U-144
lesInterFoam, U-88	startFrom keyword, U-25, U-107
mhdFoam, P-71, U-89	starToFoam utility, U-90, U-161
multiphaseInterFoam, U-88	startTime
nonNewtonianIcoFoam, U-87	keyword entry, U-25, U-107
oodles, U-88	startTime keyword, U-25, U-107
potentialFoam, P-46, U-87	status button, U-133
rasInterFoam, U-88	steady flow
reactingFoam, U-89	turbulent, P-54
rhoSimpleFoam, U-88	steadyState
rhoSonicFoam, U-88	keyword entry, U-116
rhoTurbFoam, U-88	stitchMesh utility, U-91
rhopSonicFoam, U-88	stopAt keyword, U-107
scalarTransportFoam, U-87	Stored Camera Position window panel, U-174
settlingFoam, U-88	streamFunction utility, U-92
simpleFoam, P-55, U-87	stress analysis of plate with hole, U-45
solidDisplacementFoam, U-51, U-89	stressComponents utility, U-93
solidEquilibriumDisplacementFoam, U-89	Su
sonicFoamAutoMotion, U-88	fvm member function, P-37
sonicFoam, P-61, U-88	subsetMesh utility, U-91
sonicLiquidFoam, P-65, U-88	summation convention, P-17
sonicTurbFoam, U-88	SUPERBEE differencing, P-38
turbFoam, U-20, U-88	supersonic flow, P-59
twoLiquidMixingFoam, U-88	supersonic flow over forward step, P-59
twoPhaseEulerFoam, U-88	supersonicFreeStream
solver relative tolerance, U-118	boundary condition, U-153
solver tolerance, U-118	surfaceField <type> template class, P-33</type>
solvers keyword, U-117	surfaceNormalFixedValue
sonicFoam solver, P-61, U-88 sonicFoamAutoMotion solver, U-88	boundary condition, U-153
sonicLiquidFoam solver, P-65, U-88	SuSp
sonicTurbFoam solver, U-88	fvm member function, P-37
Source	suspend button, U-133
menu entry, U-28, U-175	sutherlandTransport model, U-97, U-188
source, P-37	symm
SpalartAllmaras model, U-98, U-99	tensor member function, P-25
specie	symmetryPlane
library, U-97	boundary condition, P-65, U-150
specieThermo model, U-97, U-187	symmetryPlane
spectEddyVisc model, U-98	keyword entry, U-150
spline keyword, U-155	symmTensorField class, P-29
splitMesh utility, U-91	symmTensorThirdField class, P-29
splitMeshRegions utility, U-91	system directory, P-49, U-101
-r	-7

${f T}$	scalar division, P-18
T()	scalar multiplication, P-18
tensor member function, P-25	scale function, P-20
template class	second rank, P-16
GeometricBoundaryField, P-32	skew, P-22
fvMatrix, P-36	square of, P-20
dimensioned <type>, P-26</type>	subtraction, P-18
FieldField <type>, P-32</type>	symmetric, P-22
Field <type>, P-29</type>	symmetric rank 2, P-16
geometricField <type>, P-32</type>	symmetric rank 3, P-17
List <type>, P-29</type>	trace, P-22
pointField <type>, P-33</type>	transformation, P-21
surfaceField <type>, $P-33$</type>	transpose, P-16, P-22
volField <type>, P-33</type>	triple inner product, P-19
temporal discretisation, P-42	vector cross product, P-20
Crank Nicholson, P-43	tensor class, P-24
Euler implicit, P-42	tensor member function
explicit, P-42	*, P-25
in OpenFOAM, P-43	+, P-25
tensor, P-15	-, P-25
addition, P-18	/, P-25
algebraic operations, P-18	&, P-25
algebraic operations in OpenFOAM, P-24	&&, P-25
antisymmetric, see tensor, skew	^, P-25
calculus, P-27	cmptAv, P-25
classes in OpenFOAM, P-23	cofactors, $P-25$
cofactors, P-22	det, P-25
component average, P-20	dev, P-25
component maximum, P-20	diag, P-25
component minimum, P-20	I, P-25
determinant, P-22	inv, P-25
deviatoric, P-22	mag, P-25
diagonal, P-22	magSqr, P-25
dimension, P-16	max, P-25
double inner product, P-19	min, P-25
geometric transformation, P-21	pow, P-25
Hodge dual, P-23 hydrostatic, P-22	scale, P-25 skew, P-25
identities, P-21	sqr, P-25
identities, P-21	sqr, 1-25 symm, P-25
inner product, P-18	T(), P-25
inverse, P-23	tr. P-25
magnitude, P-20	transform, P-25
magnitude squared, P-20	tensorField class, P-29
mathematics, P-15	tensorThirdField class, P-29
notation, P-17	tetDecomposition utility, U-91
nth power, P-20	tetgenToFoam utility, U-90
outer product, P-19	text box
rank, P-16	Case Name, U-129
rank 3, P-17	Case Root, U-129

Index P-93 P-94 Index

Lower and Upper Times, U-173	totalPressure
Opacity, U-174	boundary condition, U-153
Time step, U-173	tr
times, U-33	tensor member function, P-25
thermalProperties	trace, see tensor, trace
dictionary, U-51, U-52	transform
thermophysical	tensor member function, P-25
library, U-187	transformPoints utility, U-91
thermophysicalFunctions	transportProperties
library, U-97	dictionary, U-24, U-41, U-43
thermophysicalProperties	transportProperties file, U-60
dictionary, U-187	triple inner product, P-19
thermoType keyword, U-187	triSurface
Time window, U-28	library, U-95
time	turbFoam solver, U-20, U-88
control, U-107	turbulence
time derivative, P-37	dissipation, U-42
first, P-39	kinetic energy, U-42
second, P-37, P-39	length scale, U-42
Time step text box, U-173	model, U-43
time step, U-25	turbulence keyword, U-191 turbulence model, U-42
timeFormat keyword, U-108	turbulence model, U-42 turbulenceModel keyword, U-191
timePrecision keyword, U-109	turbulenceProperties
times text box, U-33	dictionary, U-43, U-190
timeScheme keyword, U-110	turbulent flow
timeStep	steady, P-54
keyword entry, U-25, U-35, U-108	turbulentlnlet
tolerance	boundary condition, U-153
solver, U-118	tutorials
solver relative, U-118	breaking of a dam, U-57
tolerance keyword, U-54, U-117, U-118	lid-driven cavity flow, U-19
tools	stress analysis of plate with hole, U-45
adjustPhi, U-95	tutorials directory, P-45, U-19
algorithms, U-95	twoLiquidMixingFoam solver, U-88
bound, U-95	twoPhaseEulerFoam solver, U-88
compressible, U-95	type keyword, U-148
containers, U-95	TT
db, U-95 dimensionSet, U-95	U
dimensionedTypes, U-95	U field, U-25
fields, U-95	Ucomponents utility, P-72, U-36, U-93
finiteVolume, U-95	UMIST
global, U-95	keyword entry, U-111
incompressible, U-95	uncompressed
interpolations, U-95	keyword entry, U-108 uncorrected
matrices, U-95	keyword entry, U-114, U-115
meshes, U-95	uniform model, U-97
primitives, U-95	uniform keyword, U-183
wallDist, U-95	units
topoSetSource keyword, U-60	of measurement, P-26
soposossource keyword, 0-00	or measurement, 1 -20

S.I. base, P-26	foamToEnsight, U-92
uprime utility, U-93	foamToFieldview9, U-92
upwind	foamToGMV, U-92
keyword entry, U-113, U-116	foamToVTK, $U-92$
upwind differencing, P-38, U-62	gambitToFoam, U-90, U-161
Use parallel projection button, U-27, U-174	gmshToFoam, U-90
utility	icoErrorEstimate, U-94
Co, U-92	icoMomentError, U-94
FoamX, U-90	ideasToFoam, U-161
Lambda2, U-92	ideasUnvToFoam, U-90
Mach, U-92	insideCells, U-91
Pe, U-92	kivaToFoam, U-90
Q, U-92	magGradU, U-92
Rcomponents, U-93	magU, U-36, U-92
R, U-93	mapFields, U-34, U-40, U-43, U-56, U-90.
Ucomponents, P-72, U-36, U-93	U-167
adiabaticFlameT, U-94	mergeMeshes, U-91
ansys ToFoam, U-90	mirrorMesh, U-91
attachMesh, U-90	mixtureAdiabaticFlameT, U-94
autoPatch, U-91	momentScalarError, U-94
blockMesh, U-39, U-90, U-151	moveDynamicMesh, U-91
boxTurb, U-90	moveEngineMesh, U-91
	9
ccm26ToFoam, U-90	moveMesh, U-91
cellSet, U-91	mshToFoam, U-90
cfxToFoam, U-90, U-161	netgenNeutralToFoam, U-90
checkMesh, U-91, U-162	objToVTK, U-91
checkYPlus, U-93	paraFoam, U-92
chemkinToFoam, U-94	patchAverage, U-93
couplePatches, U-91	patchIntegrate, U-93
createPatch, U-91	patchTool, U-91
decomposePar, U-82, U-83, U-94	plot3dToFoam, U-90
deformedGeom, U-91	pointSet, U-91
divU, U-92	polyDualMesh, U -90
engineCompRatio, U-93	postChannel, U-93
engineSwirl, U-90	ptot, U-93
ensight74FoamExec, U-180	reconstructParMesh, U-94
ensight76FoamExec, U -92	reconstructPar, U-87, U-94
enstrophy, U-92	refineMesh, U-91
equilibriumCO, U-94	renumberMesh, U-91
equilibriumFlameT, U -94	rotateMesh, U-91
estimateScalarError, U-94	sammToFoam, U-90
extrudeMesh, U-90	sampleSurface, U-93
faceSet, U-91	sample, U-93, U-181
flattenMesh, U-91	scalePoints, U-165
flowType, U-92	setFields, U-59, U-60, U-90
fluentMeshToFoam, U-90, U-161	smapToFoam, U-92
foamDataToFluent, U-92, U-177	splitMeshRegions, U-91
foamDebugSwitches, U-94	splitMesh, U-91
foamInfoExec, U-94	starToFoam, U-90, U-161
foamMeshToFluent, U-90, U-177	stitchMesh, U-91
	the state of the s

Index P-95

stroomEunstion II 02	wallCradH utility II 02
streamFunction, U-92 stressComponents, U-93	wallGradU utility, U-93 wallHeatFlux utility, U-93
subsetMesh, U-91	
tetDecomposition, U-91	wallShearStress utility, U-93 water model, U-97
tetgenToFoam, U-90	
	wclean script/alias, U-76
transformPoints, U-91	wdot utility, U-93
uprime, U-93	wedge
vorticity, U-93	boundary condition, U-145, U-150, U-151,
wallGradU, U-93	U-160
wallHeatFlux, U-93	wedge
wallShearStress, U-93	keyword entry, U-150
wdot, U-93	window
writeCellCentres, U-93	Fields, U-28
writeMeshObj, U-90	Region, U-28
yPlusLES, U-93	Seed, U-177
zipUpMesh, U-91	Selection Window, U-27, U-172
\mathbf{V}	Time, U-28
value keyword, U-152	window panel
valueFraction keyword, U-152	Annotate, U-27, U-174
van Leer differencing, P-38	Camera Controls, U-174
vanLeer	Camera Orientation, U-174
keyword entry, U-113	Camera, U-174
vector, P-16	Display, U-27, U-28, U-172, U-174
operator, P-27	General, U-174
unit, P-20	Information, U-172
vector class, P-24, U-105	Parameters, U-28, U-172, U-173
vector product, see tensor, vector cross product	Standard Views, U-174
vectorField class, P-29	Stored Camera Position, U-174
version keyword, U-103	Wireframe
vertices keyword, U-22, U-155	menu entry, U-174
veryInhomogeneousMixture model, U-96, U-188	WM_ARCH
View menu, U-28, U-174, U-175	environment variable, U-76
viscosity	WM_COMPILE_OPTION
kinematic, U-24, U-43	environment variable, U-76
volField <type> template class, P-33</type>	WM_COMPILER
vorticity utility, U-93	environment variable, U-76
vtkFoam	WM_COMPILER_BIN
library, U-171	environment variable, U-76
TT 7	WM_COMPILER_DIR
\mathbf{W}	environment variable, U-76
wall	WM_COMPILER_LIB
boundary condition, P-65, P-71, U-150	environment variable, U-76
wall boundary type, U-42	WM_DIR
wall	environment variable, U-76
keyword entry, U-150	WM_JAVAC_OPTION
wall function, U-98	environment variable, U-76
wallBuoyantPressure	WM_LINK_LANGUAGE
boundary condition, U-153	environment variable, U-76
wallDist tools, U-95	WM_MPLIB
wallFunctionCoeffs keyword, U-191	environment variable, U-76

P-96 Index

WM_OPTIONS environment variable, U-76 WM_PROJECT environment variable, U-76 WM_PROJECT_DIR environment variable, U-76 WM_PROJECT_INST_DIR environment variable, U-76 WM_PROJECT_LANGUAGE environment variable, U-76 WM_PROJECT_USER_DIR environment variable, U-76 WM_PROJECT_VERSION environment variable, U-76 WM_SHELL environment variable, U-76	writeMeshObj utility, U-90 writeNow keyword entry, U-107 writePrecision keyword, U-100 X x keyword entry, U-183 XiFoam solver, U-89 xmgr keyword entry, U-109, U-182 Xoodles solver, U-89 xyz keyword entry, U-183 Y
vmake platforms, U-73 vmake script/alias, U-71	keyword entry, U-183 yPlusLES utility, U-93
word class, P-26, P-31 writeCellCentres utility, U-93 writeCompression keyword, U-108 writeControl keyword entry, U-108 writeControl keyword, U-25, U-61, U-108 writeFormat keyword, U-55, U-108	z keyword entry, U-183 zeroGradient boundary condition, U-152 zipUpMesh utility, U-91 zlib-1.2.1
writeInterval keyword, U-25, U-35, U-108	library, U-96