

A Comparison of Numerical Schemes on Triangular and Quadrilateral Meshes

by

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Finite volume node based Jameson and Ni schemes are compared on both triangular and quadrilateral meshes. The triangular Ni scheme is developed as an extension to the quadrilateral Ni scheme which is described as well as the triangular and quadrilateral Jameson schemes. Two different conservative freestream numerical smoothing methods are used, one of which is high-accuracy and the other low-accuracy. An extensive numerical accuracy study is performed to investigate the accuracy of these schemes and the effect of the different numerical smoothing methods. When the high-accuracy numerical smoothing is used, the accuracy study shows that all the schemes are indeed second order accurate. The low-accuracy smoothing reduces the accuracy of the schemes as well as producing less smooth solutions on irregular meshes. Both triangular and quadrilateral schemes can be used for spatial adaptation, but triangular meshes are more easily fit around complex geometries and do not require modifications in the flow solver.

Thesis Supervisor: Michael B. Giles,

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Nomenclature

a	speed of sound
A	area
E	total energy
F, G	flux vectors in x and y directions
H	total enthalpy
M	Mach number
p	pressure
s	entropy
t	time
U	state vector at nodes
U_p	primitive state vector at node
u	velocity in the x direction
v	velocity in the y direction
w	velocity magnitude
α	flow angle
γ	specific heat ratio
ρ	density
ϕ	characteristic variable

Chapter 1

Introduction

Over the last decade more and more interest has been taken in using Computational Fluid Dynamics (CFD) as a design and research tool. A productive tool must be affordable as well as accurate. Since CFD methods have become an important tool in aerodynamic design, it is important that we understand the numerical methods being used by knowing their strong points and limitations. The goal of this work is to compare the use of finite volume methods which solve the Euler equations on quadrilateral and triangular meshes to better understand them.

1.1 Background

The use of numerical solution procedures with quadrilateral meshes has been extensively studied as well as procedures such as multi-grid and spatial adaptation which reduce the computer time required to compute a solution. Multi-grid is a technique which is limited to steady state calculations in which iterations in the solution procedure are successively computed on several meshes of different node densities, and flow changes are interpolated from coarser meshes to finer meshes. In this manner flow changes are greatly accelerated and so the computational time required is on the same order as for the coarsest mesh and the accuracy is that of the finest mesh. For quadrilaterals, the finer meshes are typically found by dividing a coarse cell into four finer cells. The first multi-grid method for the Euler equations on a quadrilateral mesh was developed by Ni [15] in 1981. In 1983 another method was developed by Jameson [9]. Methods which spatially adapt automatically during the solution procedure have also been developed. These methods place small cells where the physical characteristic length is small, such as in shocks, and large cells where the physical characteristic length

is large. Dannenhoffer and Baron [2] discuss the details by which a cell can be chosen for adaptation. By using adaptation, the computational time is reduced, since to achieve a desired refinement in one portion of a mesh, a globally refined mesh is not required. The method of Dannenhoffer and Baron combines both multi-grid and adaptation to obtain the benefits of both methods.

To use CFD methods to compute the flow around complex geometries, several methods have been tried. The difficult task is placing a mesh around the geometry. One approach is to divide the flow field into several coarse blocks and a finer mesh is created in each block. This method requires a large amount of interaction between the user and the mesh generation program, but is widely used today. Another approach uses meshes which surround different parts of a geometry and overlap to form a completely meshed region. The complexity of this approach is in the need to interpolate between each mesh. A third approach involves using unstructured triangles to completely mesh the region. One particular method to create this mesh is based on Delaunay [4] triangulation which allows the mesh generation process to be automatic. Baker [1] describes the generation of a tetrahedral mesh about an entire aircraft using this method.

Interest in the third kind of mesh has prompted research in the use of triangular meshes to solve the Euler and Navier-Stokes equations. In 1985 Löhner, Morgan, Peraire and Zienkiewicz [12] presented a finite element procedure for solving the Navier-Stokes equations on a triangular mesh. A flow solver was developed by Jameson and Mavriplis [14] which solves the Euler equations for a mesh composed of triangles and a similar method was developed by Jameson, Baker and Weatherill [10] for use with tetrahedra. As with quadrilateral meshes, multi-grid and spatial adaptation methods have been developed for use with triangular meshes. A triangular multi-grid method was developed by Mavriplis and Jameson [14] which uses completely unrelated grids at each level and interpolates the solution for transfer from one mesh to another. Several spatial adaptation procedures have been developed to refine the mesh where the physical characteristic length is small. Unstructured triangular meshes may be refined without creating interfaces between regions or coarse and fine cells. Substantial work has been done by Holmes, Lamson and Connell [8] using the Delaunay triangulation method to

add new points. Another method of adaptation has been developed by Peraire, Vahdeti, Morgan and Zienkiewicz [16] which completely remeshes the region putting finer cells where the physical characteristic length is small.

Controversy exists in the CFD community as to the relative benefits of quadrilateral and triangular meshes. In particular a recent paper by Roe [19] proved that the local truncation error is only first order and he conjectured that this implies that the solution is only first order accurate. This contrasts with the demonstrated second order accuracy of finite volume quadrilateral schemes. Another paper by Giles [5] argues that on the contrary these triangular schemes can be globally second order accurate. By being more widely used for a longer period of time, quadrilateral schemes are better understood and accepted, therefore the important part of the controversy is focused on how good triangular schemes are.

The goal of this study is to address this question of how well a given computational method can perform on a triangular mesh as compared to the more commonly used quadrilateral meshes. In particular, two schemes will be examined: the node-based quadrilateral cell Jameson scheme which has been modified for triangular meshes by Mavriplis and Jameson [14], and the quadrilateral cell Ni scheme [15] which has been modified here for use on triangular meshes. Care has been taken to keep the triangular and quadrilateral versions of a scheme similar to provide a fair basis for comparison.

1.2 Overview

First a description of the numerical schemes which solve the Euler equations examined is given in Chapter 2. Next the numerical smoothing required to make these schemes stable and suppress unwanted spurious modes as well as capture shocks is described in Chapter 3. Chapter 4 presents the meshes and the pointer system which is used to describe them. These chapters complete the description of the theory required to write a computer program to implement the schemes described in Chapter 2. Flows computed with these schemes are described in Chapter 5. In Chapter 6 mathematical

and numerical methods of determining the accuracy of these schemes are described and numerical results are given. Finally a short discussion on spatial adaptation is given in Chapter 7 and some conclusions of this study are given in Chapter 8.

Chapter 2

Numerical Schemes

In this chapter the governing equations for an inviscid gas, known as the Euler equations, are presented along with two methods for numerically solving these equations. Both methods are node based, finite volume numerical schemes. The first method is a Lax-Wendroff scheme which was originally developed for quadrilateral cell meshes by Ni [15] and further expanded upon by Giles [7]. This scheme will be referred to as a “Ni Scheme” in the future. A triangular cell mesh extension of the Ni scheme which was developed by the author will also be discussed. The second method is a four step scheme which was developed by Jameson [11] for quadrilateral cell meshes and extended for triangular cell meshes by Mavriplis and Jameson [14]. This scheme will be referred to as a “Jameson Scheme”.

2.1 Governing Equations

The flows examined here are steady. To reach a steady state, unsteady equations will be integrated in time from some initial condition until there is no change in the state of the flow field.

For flows with sufficiently large Reynolds numbers the effect of viscosity is confined to a thin region near solid walls where a boundary layer exists. The governing equations used in this study neglect the viscous terms and the heat transfer terms from the full Navier-Stokes equations and are referred to as the Euler equations. These equations are first-order hyperbolic partial differential equations which can be written as

$$\frac{\partial U}{\partial t} = - \left(\frac{\partial F}{\partial x} + \frac{\partial G}{\partial y} \right) \quad (2.1)$$

where U is a state vector of dependent variables and F and G are flux vectors in the x and y directions respectively. F and G are functions of the state vector U .

$$U = \begin{bmatrix} \rho \\ \rho u \\ \rho v \\ \rho E \end{bmatrix}, \quad F = \begin{bmatrix} \rho u \\ \rho u^2 + p \\ \rho uv \\ \rho uH \end{bmatrix}, \quad G = \begin{bmatrix} \rho v \\ \rho uv \\ \rho v^2 + p \\ \rho vH \end{bmatrix} \quad (2.2)$$

The pressure p and enthalpy H can be written in terms of elements of the state vector U and the specific heat ratio γ , which is assumed to be constant.

$$p = (\gamma - 1)\rho \left(E - \frac{1}{2}(u^2 + v^2) \right) \quad (2.3)$$

$$H = E + \frac{p}{\rho} \quad (2.4)$$

The differential Equation (2.1) is written in conservation law form since the coefficients of the derivative terms are constant. Equations which can be written in conservation law form are well suited for finite volume numerical methods. The control volume used for these methods can be composed of quadrilaterals or triangles.

The flow variables are non-dimensionalized using the upstream stagnation density and stagnation speed of sound. This non-dimensionalization does not change the governing equations and gives the following stagnation quantities.

$$H = \frac{1}{\gamma - 1}, \quad \rho_o = 1, \quad p_o = \frac{1}{\gamma} \quad (2.5)$$

2.2 Quadrilateral Ni Scheme

Given values for the state vector U at time t for all nodes in the domain, it is desired to know what the state vector will be at time $t + \Delta t$. A second order Taylor series expansion for $U_{t+\Delta t}$ at a node is taken about the solution at time t where the subscripts represent the time where the function is evaluated.

$$\begin{aligned}
 U_{t+\Delta t} &= U_t + \Delta t \left(\frac{\partial U}{\partial t} \right)_t + \frac{1}{2} \Delta t^2 \left(\frac{\partial^2 U}{\partial t^2} \right)_t \\
 &= U_t - \Delta t \left(\frac{\partial F}{\partial x} + \frac{\partial G}{\partial y} \right)_t - \frac{1}{2} \Delta t \left(\frac{\partial}{\partial x} \Delta F + \frac{\partial}{\partial y} \Delta G \right)_t \\
 &= U_t - \frac{\Delta t}{A} \left(\oint (F dy - G dx) + \frac{1}{2} \oint (\Delta F dy - \Delta G dx) \right)_t \quad (2.6)
 \end{aligned}$$

where

$$\begin{aligned}
 \Delta F &= \Delta t \left(\frac{\partial F}{\partial t} \right)_t \\
 \Delta G &= \Delta t \left(\frac{\partial G}{\partial t} \right)_t \quad (2.7)
 \end{aligned}$$

Equation (2.6) is developed by substituting in the differential equation form of the governing Equation (2.1) and using Green's Theorem. The integral in Equation (2.6) is taken about a control volume around the node. This control volume is formed by connecting the centroids of the four cells surrounding the node to the midpoints of their shared faces. The control volume about node 1 is denoted by the dashed lines in Figure 2.1.

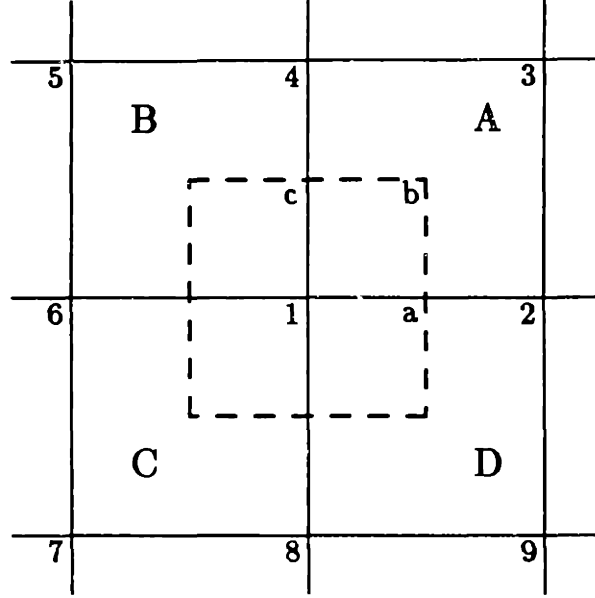


Figure 2.1: Quadrilateral cells surrounding node 1 with control volume for the Ni scheme

Equation (2.6) can be rearranged to define a change in the state vector U in time Δt .

$$\begin{aligned}\delta U &= U_{t+\Delta t} - U_t \\ &= \frac{\Delta t}{A} \left(\oint (F dy - G dx) + \frac{1}{2} \oint (\Delta F dy - \Delta G dx) \right)_t\end{aligned}\quad (2.8)$$

The first term in Equation (2.8) is the flux through the control volume and will be found by calculating the flux into each cell and distributing one quarter of this flux to each node which belongs to the cell. ΔF and ΔG will also be calculated for each cell. The second term in Equation (2.8) will be found by integrating around the control volume using the values of ΔF and ΔG for the cell which the boundary of the control volume passes through.

The change in the state vector at node 1 includes contributions from cells A, B, C and D given by

$$\delta U_1 = \delta U_{1A} + \delta U_{1B} + \delta U_{1C} + \delta U_{1D} \quad (2.9)$$

All four contributions are calculated in a similar manner.

δU_{1A} , the contribution of cell A to node 1, as shown in Figure 2.1 is given by

$$\begin{aligned} \delta U_{1A} &= (U_{t+\Delta t} - U_t)_{1A} \\ &= \left(\frac{\Delta t}{A} \right)_1 \left[-\frac{1}{4} \oint_{cell A} (F dy - G dx) - \frac{1}{2} \int_{abc} (\Delta F dy - \Delta G dx) \right]_t \\ &= \left(\frac{\Delta t}{A} \right)_1 \left[\frac{1}{4} \left(\frac{A}{\Delta t} \right)_A \Delta U_A - \frac{1}{4} (\Delta F(y_4 - y_2) - \Delta G(x_4 - x_2)) \right]_t \end{aligned} \quad (2.10)$$

where ΔU_A is found by simple trapezoidal integration around cell A.

$$\begin{aligned} \Delta U_A &= \frac{1}{2} \left(\frac{\Delta t}{A} \right)_A \left[-(F_2 + F_1)(y_2 - y_1) + (G_2 + G_1)(x_2 - x_1) \right. \\ &\quad -(F_3 + F_2)(y_3 - y_2) + (G_3 + G_2)(x_3 - x_2) \\ &\quad -(F_4 + F_3)(y_4 - y_3) + (G_4 + G_3)(x_4 - x_3) \\ &\quad \left. -(F_1 + F_4)(y_1 - y_4) + (G_1 + G_4)(x_1 - x_4) \right] \\ &= \frac{1}{2} \left(\frac{\Delta t}{A} \right)_A \left[(F_3 - F_1)(y_2 - y_4) - (G_3 - G_1)(x_2 - x_4) \right. \\ &\quad \left. + (F_4 - F_2)(y_3 - y_1) - (G_4 - G_2)(x_3 - x_1) \right] \end{aligned} \quad (2.11)$$

The terms $(\frac{\Delta t}{A})_A$ and $(\frac{\Delta t}{A})_1$ are the timestep divided by the area for cell A and node 1 respectively. The calculation of these terms will be described in Section 2.8.

Using the chain rule ΔF and ΔG evaluated at time t in Equation (2.7) can be rewritten as

$$\Delta F = \Delta t \left(\frac{\partial F}{\partial t} \right)_t = \left(\frac{\partial F}{\partial U} \right)_t \Delta U_t$$

$$\Delta G = \Delta t \left(\frac{\partial G}{\partial t} \right)_t = \left(\frac{\partial G}{\partial U} \right)_t \Delta U_t \quad (2.12)$$

where ΔU was defined in Equation (2.11). $(\frac{\partial F}{\partial U})_t$ and $(\frac{\partial G}{\partial U})_t$ are Jacobians of the flux vectors F and G evaluated at time t . For the Euler equations U , F and G are given in Equation (2.2), and for these state and flux vectors ΔF and ΔG are

$$\Delta F_A = \begin{bmatrix} (\Delta \rho u) \\ u(\Delta \rho u) + u(\rho \Delta u) + (\Delta p) \\ v(\Delta \rho u) + u(\rho \Delta v) \\ u \left((\Delta \rho E) + (\Delta p) \right) + H(\rho \Delta u) \end{bmatrix}_A \quad (2.13)$$

$$\Delta G_A = \begin{bmatrix} (\Delta \rho v) \\ u(\Delta \rho v) + v(\rho \Delta u) \\ v(\Delta \rho u) + v(\rho \Delta v) + (\Delta p) \\ v \left((\Delta \rho E) + (\Delta p) \right) + H(\rho \Delta v) \end{bmatrix}_A \quad (2.14)$$

where

$$U_A = \begin{bmatrix} \rho \\ \rho u \\ \rho v \\ \rho E \end{bmatrix}_A, \quad \Delta U_A = \begin{bmatrix} (\Delta \rho) \\ (\Delta \rho u) \\ (\Delta \rho v) \\ (\Delta \rho E) \end{bmatrix}_A \quad (2.15)$$

$$(\rho \Delta u) = (\Delta \rho u) - u(\Delta \rho)$$

$$(\rho \Delta v) = (\Delta \rho v) - v(\Delta \rho)$$

$$(\Delta p) = (\gamma - 1) \left((\Delta \rho E) - u(\Delta \rho u) - v(\Delta \rho v) + \frac{1}{2}(u^2 + v^2)(\Delta \rho) \right) \quad (2.16)$$

U_A is the value of the state vector at the cell which is the average of the state vector at the four nodes which belong the the cell. ΔU_A is found from Equation (2.11).

2.3 Triangular Ni Scheme

The triangular Ni scheme was developed here as another triangular scheme which can be used for comparison. The main difference between the quadrilateral and triangular Ni schemes is that now the control volume is formed by connecting the centroids of the triangular cells which surround the node to the midpoints of their shared faces. For discussion, let n be the number of cells surrounding a node. The control volume about node 1 where $n = 6$ is denoted by the dashed lines in Figure 2.2.

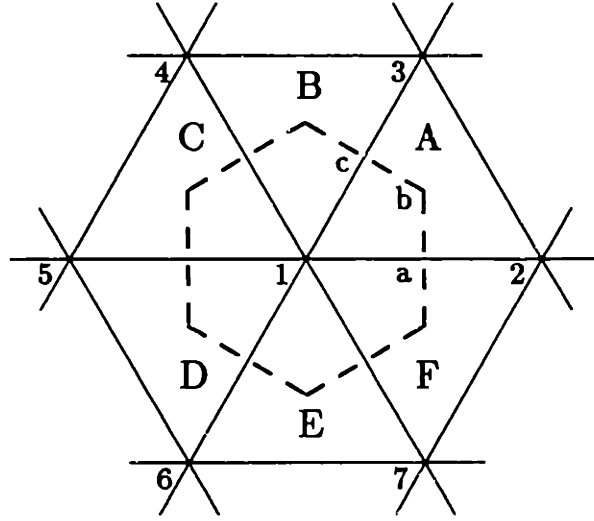


Figure 2.2: Triangular cells surrounding node 1 with control volume for the Ni scheme

The change in the state vector at node 1 includes contributions from all n cells surrounding node 1 and is given by

$$\delta U_1 = \delta U_{1A} + \delta U_{1B} + \delta U_{1C} + \cdots \text{ all } n \text{ cells} \quad (2.17)$$

All n contributions are calculated in a similar manner.

Equation (2.10) extended for the triangular scheme now becomes

$$\begin{aligned}
\delta U_{1A} &= (U_{t+\Delta t} - U_t)_{1A} \\
&= \left(\frac{\Delta t}{A} \right)_1 \left[-\frac{1}{3} \oint_{cell A} (F dy - G dx) - \frac{1}{2} \int_{abc} (\Delta F dy - \Delta G dx) \right]_t \\
&= \left(\frac{\Delta t}{A} \right)_1 \left[\frac{1}{3} \left(\frac{A}{\Delta t} \right)_A \Delta U_A - \frac{1}{4} (\Delta F(y_3 - y_2) - \Delta G(x_3 - x_2)) \right]_t \quad (2.18)
\end{aligned}$$

where ΔU_A is again found by simple trapezoidal integration around cell A.

$$\begin{aligned}
\Delta U_A &= \frac{1}{2} \left(\frac{\Delta t}{A} \right)_A \left[-(F_2 + F_1)(y_2 - y_1) + (G_2 + G_1)(x_2 - x_1) \right. \\
&\quad \left. -(F_3 + F_2)(y_3 - y_2) + (G_3 + G_2)(x_3 - x_2) \right. \\
&\quad \left. -(F_1 + F_3)(y_1 - y_3) + (G_1 + G_3)(x_1 - x_3) \right] \\
&= \frac{1}{2} \left(\frac{\Delta t}{A} \right)_A \left[-F_1(y_2 - y_3) + G_1(x_2 - x_3) \right. \\
&\quad \left. -F_2(y_3 - y_1) + G_2(x_3 - x_1) \right. \\
&\quad \left. -F_3(y_1 - y_2) + G_3(x_1 - x_2) \right] \quad (2.19)
\end{aligned}$$

The terms $(\frac{\Delta t}{A})_A$ and $(\frac{\Delta t}{A})_1$ are once again the timestep divided by the area for cell A and node 1 respectively. The calculation of these terms will be described in Section 2.8.

For the Euler equations ΔF and ΔG are found in the same manner as for the quadrilaterals from Equation (2.16), except U_A is the average of the state vector at the three nodes which belong the the cell and ΔU_A is found from Equation (2.19).

2.4 Quadrilateral Jameson Scheme

Like the Ni scheme, the value of the state vector, U , is known at time t for all nodes in the domain, and it is desired to know the value of this state vector at time $t + \Delta t$.

A multi-stage time stepping method is used for the Jameson scheme where a first order approximation is used at each step. In particular a four stage time stepping method is used in this thesis.

$$\begin{aligned}
U_0 &= U_t \\
U_1 &= U_0 + \alpha_1 \Delta t \left(\frac{\partial U}{\partial t} \right)_0 \\
&= U_0 - \alpha_1 \Delta t \left(\frac{\partial F}{\partial x} + \frac{\partial G}{\partial y} \right)_0 \\
&= U_0 - \alpha_1 \frac{\Delta t}{A} \oint (F dy - G dx)_0 \\
&= U_0 - \alpha_1 \frac{\Delta t}{A} (\text{flux})_0 \\
U_2 &= U_0 - \alpha_2 \frac{\Delta t}{A} (\text{flux})_1 \\
U_3 &= U_0 - \alpha_3 \frac{\Delta t}{A} (\text{flux})_2 \\
U_{t+\Delta t} &= U_0 - \alpha_4 \frac{\Delta t}{A} (\text{flux})_3
\end{aligned} \tag{2.20}$$

$$\alpha_1 = \frac{1}{4}, \quad \alpha_2 = \frac{1}{3}, \quad \alpha_3 = \frac{1}{2}, \quad \alpha_4 = 1 \tag{2.21}$$

where subscripts indicate which state vector is used to calculate the flux. The term $(\frac{\Delta t}{A})$ is the timestep divided by the area evaluated at the node to keep the scheme conservative. The calculation of this term will be described in Section 2.8.

Equation (2.20) is developed in a similar fashion as Equation (2.7) by substituting in the differential form of the governing Equation (2.1) and using Green's theorem. The first order terms in the Jameson scheme are the same as the first order terms in the Ni scheme, but are described differently by considering a different control volume. In fact, in the limit of small timestep the Ni scheme reduces to the Jameson scheme. The integral in Equation (2.20) is taken about a control volume around the node which is formed by all of the four cells around the node, therefore the control volumes in the

Jameson scheme are overlapping. This control volume is shown about node 1 by the dashed lines in Figure 2.3.

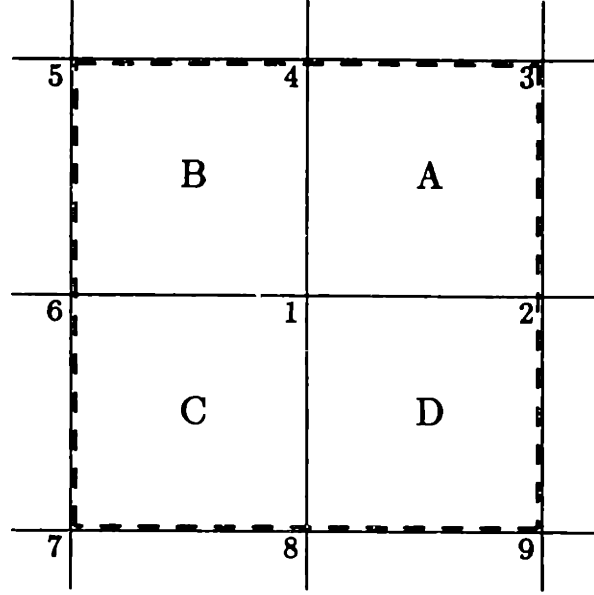


Figure 2.3: Quadrilateral cells surrounding node 1 with control volume for the Jameson scheme

The integral in Equation (2.20) is calculated by finding the contribution of each cell to the flux into the control volume for each node. The contribution from cell A to the flux at nodes 1, 2, 3 and 4 is the same.

$$\text{flux}_{1A} = \text{flux}_{2A} = \text{flux}_{3A} = \text{flux}_{4A} = \text{flux}_A \quad (2.22)$$

where

$$\begin{aligned} \text{flux}_A = \frac{1}{2} \bigg[& -(F_2 + F_1)(y_2 - y_1) + (G_2 + G_1)(x_2 - x_1) \\ & -(F_3 + F_2)(y_3 - y_2) + (G_3 + G_2)(x_3 - x_2) \\ & -(F_4 + F_3)(y_4 - y_3) + (G_4 + G_3)(x_4 - x_3) \\ & -(F_1 + F_4)(y_1 - y_4) + (G_1 + G_4)(x_1 - x_4) \bigg] \end{aligned}$$

$$= \frac{1}{2} \left[\begin{aligned} & (F_3 - F_1)(y_2 - y_4) - (G_3 - G_1)(x_2 - x_4) \\ & + (F_4 - F_2)(y_3 - y_1) - (G_4 - G_2)(x_3 - x_1) \end{aligned} \right] \quad (2.23)$$

The flux at node 1 is then the sum of the contributions from the four surrounding cells.

$$\text{flux}_1 = \text{flux}_{1A} + \text{flux}_{1B} + \text{flux}_{1C} + \text{flux}_{1D} \quad (2.24)$$

where flux_{1B} , flux_{1C} and flux_{1D} are calculated in a similar fashion as flux_{1A} .

2.5 Triangular Jameson Scheme

The triangular Jameson scheme is similar to the quadrilateral Jameson scheme except now the cells which surround a node are triangular. For discussion let n be the number of cells surrounding a node. The control volume about node 1 where $n = 6$ is denoted by the dashed lines in Figure 2.4.

The integral in Equation (2.20) is once again calculated by finding the contribution of each cell to the flux into the control volume for each node. The contribution from cell A to the flux at nodes 1, 2 and 3 is the same.

$$\text{flux}_{1A} = \text{flux}_{2A} = \text{flux}_{3A} = \text{flux}_A \quad (2.25)$$

where

$$\begin{aligned} \text{flux}_A = \frac{1}{2} \left[\begin{aligned} & -(F_2 + F_1)(y_2 - y_1) + (G_2 + G_1)(x_2 - x_1) \\ & -(F_3 + F_2)(y_3 - y_2) + (G_3 + G_2)(x_3 - x_2) \\ & -(F_1 + F_3)(y_1 - y_3) + (G_1 + G_3)(x_1 - x_3) \end{aligned} \right] \end{aligned}$$

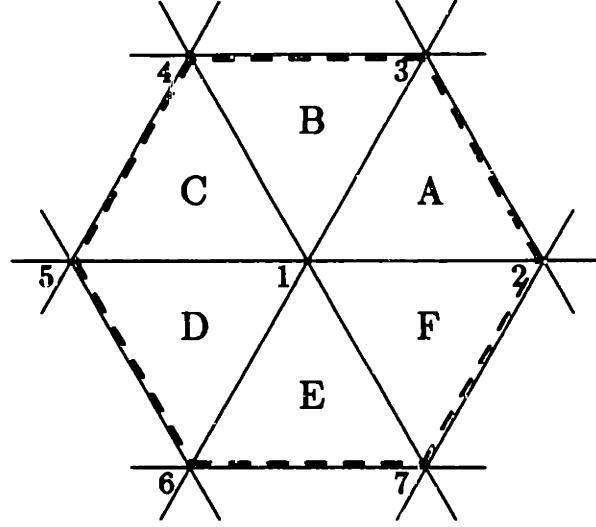


Figure 2.4: Triangular cells surrounding node 1 with control volume for the Jameson scheme

$$\begin{aligned}
 = \frac{1}{2} \bigg[& -F_1(y_2 - y_3) + G_1(x_2 - x_3) \\
 & -F_2(y_3 - y_1) + G_2(x_3 - x_1) \\
 & -F_3(y_1 - y_2) + G_3(x_1 - x_2) \bigg] \quad (2.26)
 \end{aligned}$$

The flux at node 1 is then the sum of the contributions from the n surrounding cells.

$$\text{flux}_1 = \text{flux}_{1A} + \text{flux}_{1B} + \dots \text{ all } n \text{ cells} \quad (2.27)$$

where flux_{1B} , flux_{1C} , \dots are calculated in a similar fashion as flux_{1A} .

2.6 Farfield Boundary Conditions

The farfield boundary conditions are applied at the inlet and outlet boundary nodes, and are used for both the Ni and Jameson schemes. The boundary conditions on hyperbolic equations must correctly close the system of equations. Linear characteristic

theory determines the direction of wave motion in and out of the domain, and thus where boundary conditions must be imposed. The following analysis is described by Giles in [7] and a general formulation is presented by Dannenhoffer in [3]. To simplify the process, primitive state vector variables U_p are used where

$$U_p = \begin{bmatrix} \rho \\ u \\ v \\ p \end{bmatrix} \quad (2.28)$$

The first step in the process is to linearize the governing equations where the spatial directions x and y are along the grid lines.

$$\frac{\partial U_p}{\partial t} + A \frac{\partial U_p}{\partial x} + B \frac{\partial U_p}{\partial y} = 0 \quad (2.29)$$

A and B are constant matrices evaluated at some reference state. The wave propagation normal to the boundary (in the x direction) is dominant, therefore variations parallel to the boundary (in the y direction) will be neglected. Equation (2.29) becomes

$$\frac{\partial U_p}{\partial t} + A \frac{\partial U_p}{\partial x} = 0 \quad (2.30)$$

The reference state for evaluation of the matrix A will be the flow on the boundary at the old timestep. To reduce computational effort, the average value of the state vector on the boundary will be used to evaluate A . This state will be denoted by the subscript $()_{old-ave}$. The constant matrix A is then

$$A = \begin{bmatrix} u & \rho & 0 & 0 \\ 0 & u & 0 & \frac{1}{\rho} \\ 0 & 0 & u & 0 \\ 0 & \rho a^2 & 0 & u \end{bmatrix}_{old-ave} \quad (2.31)$$

The matrix A can be diagonalized by a similarity transformation,

$$\mathbf{T}^{-1}\mathbf{A}\mathbf{T} = \begin{bmatrix} u & 0 & 0 & 0 \\ 0 & u & 0 & 0 \\ 0 & 0 & u+a & 0 \\ 0 & 0 & 0 & u-a \end{bmatrix}_{old-ave} = \mathbf{\Lambda} \quad (2.32)$$

where the matrix \mathbf{T} is the matrix of right eigenvectors of \mathbf{A} and the matrix \mathbf{T}^{-1} is the matrix of left eigenvectors of \mathbf{A} . Matrix $\mathbf{\Lambda}$ is a diagonal matrix whose elements are the eigenvalues of matrix \mathbf{A} .

$$\mathbf{T} = \begin{bmatrix} \frac{-1}{a^2} & 0 & \frac{1}{2a^2} & \frac{1}{2a^2} \\ 0 & 0 & \frac{1}{2\rho a} & \frac{-1}{2\rho a} \\ 0 & \frac{1}{\rho a} & 0 & 0 \\ 0 & 0 & \frac{1}{2} & \frac{1}{2} \end{bmatrix}_{old-ave}, \quad \mathbf{T}^{-1} = \begin{bmatrix} -a^2 & 0 & 0 & 1 \\ 0 & 0 & \rho a & 0 \\ 0 & \rho a & 0 & 1 \\ 0 & -\rho a & 0 & 1 \end{bmatrix}_{old-ave} \quad (2.33)$$

Multiplication of Equation (2.30) by \mathbf{T}^{-1} produces the equation

$$\frac{\partial \phi}{\partial t} + \mathbf{\Lambda} \frac{\partial \phi}{\partial x} = 0 \quad (2.34)$$

where $\phi = \mathbf{T}^{-1}U_p$. Variation from the values at the old timestep will be considered, therefore $\delta\phi = \mathbf{T}^{-1}\delta U_p$. The four equations in the system of equations (2.34) are now independent. The elements of ϕ are the linearized characteristic variables and the corresponding elements of $\mathbf{\Lambda}$ indicate the direction of the flow of information. For subsonic flow where $0 < u < a$ the first three characteristics give waves propagating downstream since the corresponding elements of $\mathbf{\Lambda}$ are positive and the fourth propagating upstream since the fourth element of $\mathbf{\Lambda}$ is negative. For supersonic flow where $u > a$ all waves propagate downstream since all elements of $\mathbf{\Lambda}$ are positive.

2.6.1 Subsonic Inlet Boundary

For subsonic flow three pieces of information must be specified at the inlet boundary; here they will be the total enthalpy, H , the entropy, s , and the flow angle, α .

$$H = \left(\frac{\gamma}{\gamma - 1} \right) \frac{p}{\rho} + \frac{1}{2}(u^2 + v^2) \quad (2.35)$$

$$s = \log(\gamma p) - \gamma \log(\rho) \quad (2.36)$$

$$\tan \alpha = \frac{v}{u} \quad (2.37)$$

A fourth piece of information comes from the interior of the flow field and will be taken from the values the numerical solver predicts, therefore denoted by the subscript $()_{pred}$. Let the subscript $()_{spec}$ stand for the value which is specified by the inlet flow conditions. The subscripts $()_{old}$ and $()_{new}$ will stand for the values at the old and new time steps. The amount needed to bring the old values of H , s and $\tan \alpha$ on the boundary to the specified values can be written in terms of a first order Taylor series in ϕ . The constant coefficient of the series can be changed by using the chain rule to contain elements of U_p .

$$\begin{aligned} (H)_{spec} &= (H)_{new} \\ &= (H)_{old} + \left(\frac{\partial H}{\partial \phi} \right)_{old-ave} \delta \phi \\ &= (H)_{old} + \left(\frac{\partial H}{\partial U_p} \right)_{old-ave} \mathbf{T} \delta \phi \\ &= (H)_{old} + \left[\frac{1}{(\gamma-1)\rho} \frac{v}{\rho a} \frac{a+u}{2\rho a} \frac{a-u}{2\rho a} \right]_{old-ave} \delta \phi \end{aligned} \quad (2.38)$$

$$(s)_{spec} = (s)_{old} + \left[\frac{1}{\rho} \quad \rho \quad 0 \quad 0 \right]_{old-ave} \delta \phi \quad (2.39)$$

$$(\tan \alpha)_{spec} = (\tan \alpha)_{old} + \left[0 \quad \frac{1}{\rho u a} \quad \frac{-v}{2\rho u^2 a} \quad \frac{v}{2\rho u^2 a} \right]_{old-ave} \delta \phi \quad (2.40)$$

The change in the fourth characteristic $\delta(\phi_4)$ is equal to the change that the flow field predicts, $\delta(\phi_4)_{pred}$. Since $\delta \phi = \mathbf{T}^{-1} \delta U_p$, the predicted change in the characteristic variable $\delta(\phi_4)_{pred}$ is found to be

$$\delta(\phi_4)_{pred} = (-\rho a)_{old} \delta u_{pred} + \delta p_{pred} \quad (2.41)$$

There are now four equations for the change in the characteristic variable ϕ which can be written in matrix form.

$$\begin{bmatrix} (H)_{spec} - (H)_{old} \\ s_{spec} - s_{old} \\ (\tan \alpha)_{spec} - (\tan \alpha)_{old} \\ \delta(\phi_4)_{pred} \end{bmatrix} = \begin{bmatrix} \frac{1}{(\gamma-1)\rho} & \frac{u}{\rho a} & \frac{a+u}{2\rho a} & \frac{a-u}{2\rho a} \\ \frac{1}{\rho} & \rho & 0 & 0 \\ 0 & \frac{1}{\rho u a} & \frac{-v}{2\rho u^2 a} & \frac{v}{2\rho u^2 a} \\ 0 & 0 & 0 & 1 \end{bmatrix}_{old-ave} \begin{bmatrix} \delta(\phi_1) \\ \delta(\phi_2) \\ \delta(\phi_3) \\ \delta(\phi_4) \end{bmatrix} \quad (2.42)$$

Using the relation $\delta U_p = T \delta \phi$, $\delta \phi$ in Equation (2.42) can now be changed back into primitive state vector values U_p .

$$\delta U_p = \begin{bmatrix} \delta(\rho) \\ \delta(u) \\ \delta(v) \\ \delta(p) \end{bmatrix}_{new} = \frac{1}{ua + u^2 + v^2} \begin{bmatrix} \frac{\rho u}{a} & \frac{-p}{a} \left(\frac{\gamma u}{\gamma-1} + \frac{u^2 + v^2}{a} \right) & \frac{-\rho v}{a} & \frac{u^2 + v^2}{a^2} \\ u & \frac{-up}{(\gamma-1)\rho} & -v & \frac{-u}{\rho} \\ v & \frac{-vp}{(\gamma-1)\rho} & a+u & \frac{-v}{\rho} \\ \rho a u & \frac{-a u p}{\gamma-1} & -\rho a v & u^2 + v^2 \end{bmatrix}_{old-ave} \begin{bmatrix} (H)_{spec} - (H)_{old} \\ s_{spec} - s_{old} \\ \tan \alpha_{spec} - \tan \alpha_{old} \\ \delta \phi_{4pred} \end{bmatrix} \quad (2.43)$$

To transform the change in the primitive state vector variables δU_p back into the change in the state vector δU the following transformation is performed

$$\delta U_{new} = \begin{bmatrix} \delta(\rho) \\ \delta(\rho u) \\ \delta(\rho v) \\ \delta(\rho E) \end{bmatrix}_{new} = \begin{bmatrix} 1 & 0 & 0 & 0 \\ u & \rho & 0 & 0 \\ v & \rho & 0 & 0 \\ \frac{1}{2}(u^2 + v^2) & \rho u & \rho v & \frac{1}{\gamma-1} \end{bmatrix}_{old} \begin{bmatrix} \delta(\rho) \\ \delta(u) \\ \delta(v) \\ \delta(p) \end{bmatrix}_{new} \quad (2.44)$$

So finally the new value of the state vector at the inlet nodes is

$$U_{new} = U_{old} + \delta U_{new} \quad (2.45)$$

2.6.2 Subsonic Outlet Boundary

For subsonic flow the outlet boundary is similar to the inlet boundary except now only one piece of information must be specified; here static pressure p will be used. Once again the amount needed to bring the old value of p to the specified value on the boundary can be written in terms of a first order Taylor series in ϕ .

$$\begin{aligned} p_{spec} &= p_{new} \\ &= p_{old} + \left(\frac{\partial p}{\partial \phi} \right)_{old-ave} \delta \phi \\ &= p_{old} + \left(\frac{\partial p}{\partial U_p} \right)_{old-ave} \left(\frac{\partial U_p}{\partial \phi} \right)_{old-ave} \delta \phi \\ &= p_{old} + \begin{bmatrix} 0 & 0 & \frac{1}{2} & \frac{1}{2} \end{bmatrix}_{old-ave} \delta \phi \end{aligned} \quad (2.46)$$

The change in the first, second and third characteristics is equal to the change the flow field predicts. Again, since $\delta \phi = T^{-1} \delta U_p$, the predicted change in the characteristic variables $\delta(\phi_1)_{pred}$, $\delta(\phi_2)_{pred}$ and $\delta(\phi_3)_{pred}$ are found to be

$$\delta(\phi_1)_{pred} = (-a^2)_{old} \delta \rho_{pred} + \delta p_{pred} \quad (2.47)$$

$$\delta(\phi_2)_{pred} = (\rho a)_{old} \delta v_{pred} \quad (2.48)$$

$$\delta(\phi_3)_{pred} = (\rho a)_{old} \delta u_{pred} + \delta p_{pred} \quad (2.49)$$

As in the subsonic inlet condition, there are now four equations for the change in the characteristic variable ϕ .

$$\begin{bmatrix} \delta(\phi_1)_{pred} \\ \delta(\phi_2)_{pred} \\ \delta(\phi_3)_{pred} \\ p_{spec} - p_{old} \end{bmatrix} = \begin{bmatrix} 1 & 0 & 0 & 0 \\ 0 & 1 & 0 & 0 \\ 0 & 0 & 1 & 0 \\ 0 & 0 & \frac{1}{2} & \frac{1}{2} \end{bmatrix} \begin{bmatrix} \delta(\phi_1) \\ \delta(\phi_2) \\ \delta(\phi_3) \\ \delta(\phi_4) \end{bmatrix} \quad (2.50)$$

Using the relation $\delta U_p = T \delta \phi$, $\delta \phi$ in Equation (2.50) can now be changed back into primitive state vector values U_p .

$$\begin{aligned} \delta U_p &= \begin{bmatrix} \delta(\rho) \\ \delta(u) \\ \delta(v) \\ \delta(p) \end{bmatrix}_{new} \\ &= \begin{bmatrix} \frac{-1}{a^2} & 0 & 0 & \frac{1}{a^2} \\ 0 & 0 & \frac{1}{\rho a} & \frac{-1}{\rho a} \\ 0 & \frac{1}{\rho a} & 0 & 0 \\ 0 & 0 & 0 & 1 \end{bmatrix}_{old-ave} \begin{bmatrix} \delta\phi_{1pred} \\ \delta\phi_{2pred} \\ \delta\phi_{3pred} \\ p_{spec} - p_{old} \end{bmatrix} \end{aligned} \quad (2.51)$$

The primitive state vector variables, δU_p , are transformed to the state vector, δU , by Equation (2.44) and the update is performed as in Equation (2.45).

2.6.3 Supersonic Inlet Boundary

Since for supersonic flow all waves flow downstream, to implement the inlet boundary condition it is simply necessary to prescribe the flow conditions. The inlet Mach number M_{spec} is specified as well as the flow angle α_{spec} , and the flow conditions are found from these variables.

$$\begin{aligned} \rho &= \left(1 + \frac{(\gamma-1)}{2} M_{spec}^2 \right)^{\frac{-1}{\gamma-1}} \\ a &= \left(1 + \frac{(\gamma-1)}{2} M_{spec}^2 \right)^{-\frac{1}{2}} \end{aligned}$$

$$w = \sqrt{u^2 + v^2} = M_{spec} a$$

$$p = \frac{\rho a^2}{\gamma}$$

The state vector is fixed at

$$U = \begin{bmatrix} \rho \\ \rho w \cos \alpha_{spec} \\ \rho w \sin \alpha_{spec} \\ \frac{p}{\gamma-1} + \frac{1}{2} w^2 \rho \end{bmatrix}_{new} \quad (2.52)$$

2.6.4 Supersonic Outlet Boundary

For the supersonic outlet all waves flow out of the boundary, so the change in the state vector predicted by the scheme is used here.

$$\delta U_{new} = \delta U_{pred} \quad (2.53)$$

2.7 Wall Boundary

Two conditions are applied at the solid wall boundaries; first that there is no flux through the wall faces and second that the flow is tangent to the wall at the wall nodes. It would seem that the first condition would be sufficient to satisfy the second condition without explicitly enforcing tangency, but this is not the case since the scheme is node based. The flux on the face is computed by averaging the flux at the nodes, therefore it is possible to have no flux through the walls while the nodal flux oscillates about zero at the wall nodes. By imposing tangency at the nodes, this oscillatory state cannot occur.

2.7.1 Ni Scheme

Since for the Ni scheme only half as many cells surround a boundary node as an interior node, the control volume is half the size of the control volumes in the interior of the flow field. Boundary cells are shown in Figure 2.5 with the boundaries of the control volume denoted by dashed lines. When computing ΔU for the wall boundary cells from Equation (2.11) for quadrilateral cells or Equation (2.19) for triangular cells, the flux through the wall face consists only of the pressure term. The contribution to ΔU from cells A, B (and C in the triangular case) to node 1 in Figure 2.5 is then found in the same manner as for the interior nodes. The second order contribution to node 1 is also the same as for the interior nodes, except that now the integral must also be taken along the wall boundary where ΔF and ΔG consist only of the pressure term. For cell A the second order contribution from ΔF and ΔG along the boundary become

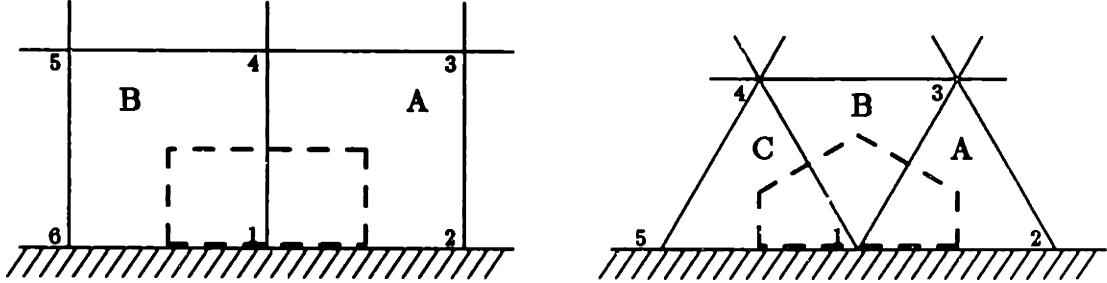


Figure 2.5: Boundary cells for Ni scheme with control volume

$$\Delta F_A = \begin{bmatrix} 0 \\ (\Delta p) \\ 0 \\ 0 \end{bmatrix}_A, \quad \Delta G_A = \begin{bmatrix} 0 \\ 0 \\ (\Delta p) \\ 0 \end{bmatrix}_A \quad (2.54)$$

where as before

$$(\Delta p) = (\gamma - 1) \left((\Delta \rho E) - u(\Delta \rho u) - v(\Delta \rho v) + \frac{1}{2}(u^2 + v^2)(\Delta \rho) \right) \quad (2.55)$$

2.7.2 Jameson Scheme

The control volume for the Jameson scheme boundary nodes is also half the size of the control volumes in the interior of the flow field. The boundary control volume is shown in Figure 2.6. The no flux condition is easier to implement for the Jameson scheme since there are only first order terms. When computing (flux) for the wall boundary cells from Equation (2.23) for quadrilateral cells or Equation (2.26) for triangular cells the flux through the wall face consists only of the pressure terms.

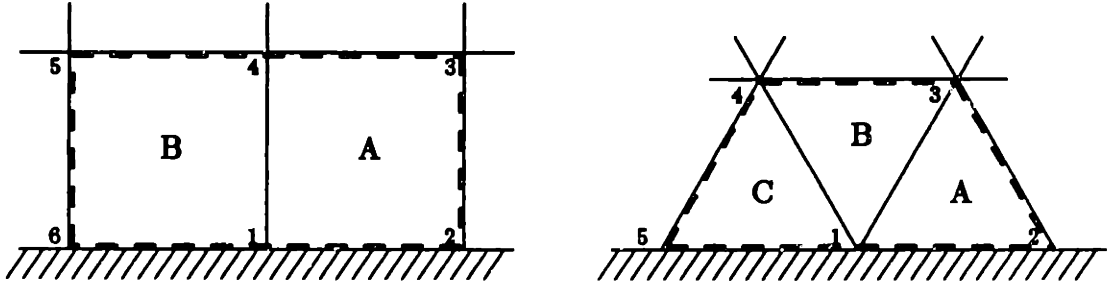


Figure 2.6: Boundary cells for Jameson scheme with control volume

2.7.3 Tangency Condition

The change in the state vector for both schemes is such that the new state vector will satisfy flow tangency on the boundary. At each wall node a flow angle, α , is prescribed.

For both schemes the second and third components of the state vector are changed from the value predicted by the solver, denoted by the subscript ()_{before}, so the new values become

$$\begin{aligned}(\rho u)_{new} &= (\rho u)_{before} + (\rho w)_n \sin \alpha \\(\rho v)_{new} &= (\rho v)_{before} - (\rho w)_n \cos \alpha\end{aligned}\tag{2.56}$$

where

$$(\rho w)_n = -(\rho u)_{before} \sin \alpha + (\rho v)_{before} \cos \alpha \quad (2.57)$$

and for the Ni scheme,

$$(\rho u)_{before} = (\rho u)_{old} + \delta(\rho u)_{old} \quad (2.58)$$

$$(\rho v)_{before} = (\rho v)_{old} + \delta(\rho v)_{old} \quad (2.59)$$

2.8 Timestep

In the description of the basic schemes the timestep divided by the area, $(\frac{\Delta t}{A})$, was referred to for a cell or a node. The term $(\frac{\Delta t}{A})$ is computed on a cell basis and the value at the node is found by summing $(\frac{\Delta t}{A})$ for the cells surrounding the node. A factor of $\frac{1}{4}$ is multiplied by the nodal $(\frac{\Delta t}{A})$ for the quadrilateral Ni scheme and $\frac{1}{3}$ for the triangular Ni scheme since the control volume at a node only includes this fraction of the cells surrounding that node.

For the quadrilateral schemes the maximum stable $(\frac{\Delta t}{A})$ on a cell A is defined by Usab in [22] from a Von Neumann stability analysis for the linearized 2-D Euler equations.

$$\left(\frac{\Delta t}{A}\right)_{max} = CFL \cdot \min \left\{ \frac{1}{|u\Delta y^l - v\Delta x^l| + a\Delta l}, \frac{1}{|u\Delta y^m - v\Delta x^m| + a\Delta m} \right\} \quad (2.60)$$

The flow variables u , v and a are the average of the nodal values for the cell. For cell A with nodes 1, 2, 3 and 4 running counterclockwise around the cell as shown in Figure 2.1,

$$\begin{aligned} \Delta x^l &= \frac{1}{2}(x_2 + x_3 - x_1 - x_4), & \Delta x^m &= \frac{1}{2}(x_1 + x_2 - x_4 - x_5) \\ \Delta y^l &= \frac{1}{2}(y_2 + y_3 - y_1 - y_4), & \Delta y^m &= \frac{1}{2}(y_1 + y_2 - y_4 - y_5) \\ \Delta l &= \sqrt{(\Delta x^l)^2 + (\Delta y^l)^2}, & \Delta m &= \sqrt{(\Delta x^m)^2 + (\Delta y^m)^2} \end{aligned} \quad (2.61)$$

For the triangular schemes the $(\frac{\Delta t}{A})$ is found by Giles in [6] from energy methods.

$$\left(\frac{\Delta t}{A}\right)_{\max} = \frac{CFL \cdot 2}{|u\Delta x^l - v\Delta y^l| + |u\Delta x^m - v\Delta y^m| + |u\Delta x^n - v\Delta y^n| + a(\Delta l + \Delta m + \Delta n)} \quad (2.62)$$

The flow variables u , v and a are again the average of the nodal values for the cell. For cell A with nodes 1, 2 and 3 running counterclockwise around the cell as shown in Figure 2.3

$$\begin{aligned} \Delta x^l &= \frac{1}{2}(x_2 - x_1), & \Delta y^l &= \frac{1}{2}(y_2 - y_1) \\ \Delta x^m &= \frac{1}{2}(x_3 - x_2), & \Delta y^m &= \frac{1}{2}(y_3 - y_2) \\ \Delta x^n &= \frac{1}{2}(x_1 - x_3), & \Delta y^n &= \frac{1}{2}(y_1 - y_3) \\ \Delta l &= \sqrt{(\Delta x^l)^2 + (\Delta y^l)^2} \\ \Delta m &= \sqrt{(\Delta x^m)^2 + (\Delta y^m)^2} \\ \Delta n &= \sqrt{(\Delta x^n)^2 + (\Delta y^n)^2} \end{aligned} \quad (2.63)$$

CFL stands for the Courant, Friedrichs, and Lewy number which gives the timestep limit for stability. In [22] Usab presents a Von Neumann stability analysis for the linearized 2-D Euler equations on the quadrilateral Ni scheme. The stability restriction from this analysis is $CFL \leq \frac{1}{\sqrt{2}}$. Usab then says that this restriction is too strict and that in practice $CFL \leq 1$ is used. This observation was confirmed in this work for both the quadrilateral and triangular Ni schemes. It is possible that the non-linearity in the Euler equations or numerical smoothing cause the increase in CFL limit. Using energy methods, Giles shows in [6] that for the four step method used here for the triangular Jameson scheme the stability limit gives $CFL \leq 2\sqrt{2}$. The method finds the CFL limit for which the energy associated with a solution is non-increasing. It has been shown that for the four step quadrilateral Jameson scheme the stability limit also gives $CFL \leq 2\sqrt{2}$. In practice it was found that this limit is not strict enough and $CFL \leq 2$ was used.

Chapter 3

Numerical Smoothing

Numerical smoothing is a dissipative operator which is added to numerical schemes to damp out oscillations in the solution and provide stability. The Jameson schemes do not have a dissipative term and are unstable without the addition of a dissipative smoothing operator. Numerical smoothing is also required for the Jameson schemes to eliminate steady state, spatially-oscillatory modes which are allowed by the scheme. Three modes are allowed for the quadrilateral Jameson scheme and are shown as modes a, b and c in Figure 3.1. Three modes are also allowed for the triangular Jameson scheme. One is shown in Figure 3.2 and the other two are similar modes shifted by one node. For the Ni schemes there is a dissipative term in the numerical operator and the scheme is stable for a smooth flow field. For the quadrilateral Ni scheme only one oscillatory mode is allowed which is shown as mode a in Figure 3.1. The triangular Ni scheme does not allow any of the oscillatory modes, but in the limit of very small timestep the Ni scheme reduces to the Jameson scheme which, as mentioned, allows three modes. To provide stability and to eliminate oscillatory modes a fourth difference *freestream* smoothing operator is used. For both schemes numerical smoothing is required to capture discontinuities such as shocks. The smoothing required to capture shocks will be referred to as *shock* smoothing.

First, the different second difference operators used here will be described. These operators are used to formulate the smoothing operators. Next the different methods of freestream and shock smoothing will be described.

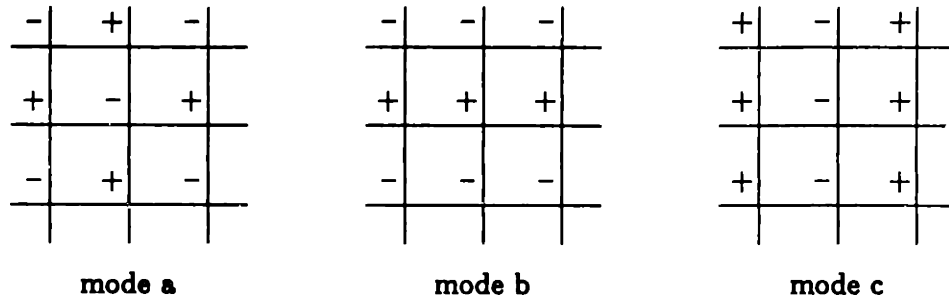


Figure 3.1: Possible modes for quadrilateral cells

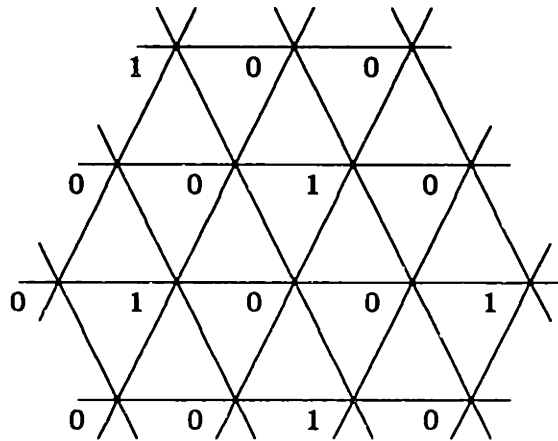


Figure 3.2: Possible mode for triangular cells

3.1 Second Difference Operators

To compute the fourth difference smoothing operator for freestream smoothing, a second difference of a second difference is computed. For both quadrilaterals and triangles two second difference operators are examined. The first is a relatively fast operator which gives a non-zero second difference for a linear function on a non-uniform grid. The second operator is slower but results in a zero second difference for a linear function. By examining the effect of the second difference operator on a linear function the accuracy of the operator is tested, since for second order or higher accuracy the contribution must be zero.

Typical triangular and quadrilateral cells are shown in Figure 3.3 with the corresponding nodes.

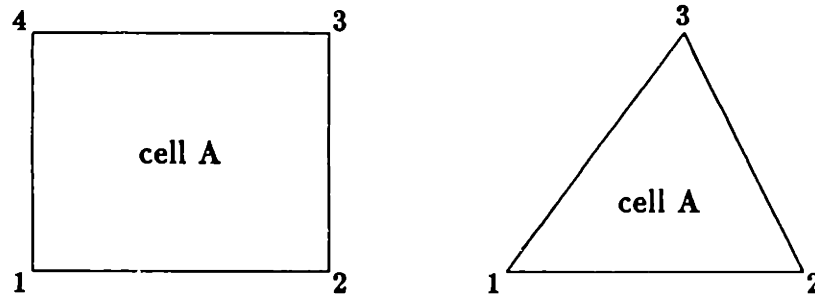


Figure 3.3: Typical triangular and quadrilateral cells

3.1.1 Low-Accuracy Second Difference

This operator is not dependent on the location of the nodes surrounding the node for which the second difference is computed, but merely on the function values at these nodes. For a triangular mesh the contribution from cell A to the second difference at node 1 is

$$(D^2S)_{1A} = (S_3 + S_2 - 2S_1) \quad (3.1)$$

where S is the variable for which the second difference is computed. For a quadrilateral mesh the contribution from cell A to the second difference at node 1 is

$$(D^2S)_{1A} = (S_4 + S_3 + S_2 - 3S_1) \quad (3.2)$$

This second difference is conservative for both triangular and quadrilateral meshes since the total contribution of each cell is zero.

3.1.2 High-Accuracy Second Difference

This operator consists of finding the first derivative for each cell and then combining the derivatives on the cells surrounding a node to form a second difference. Unlike the low-accuracy second difference operator, this operator is dependent on the grid geometry. The operator for a triangular cell mesh will be examined first since the operator for a quadrilateral cell mesh is essentially a triangular operator.

Referring to Figure 2.3 the first derivative is found with respect to x and y for cell A.

$$\begin{aligned} (S_x)_A &= \frac{1}{A_A} \iint_{\text{cell } A} \frac{\partial S}{\partial x} dx dy \\ &= \frac{1}{A_A} \int_{1-2-3} S dy \\ &= \frac{1}{A_A} \left(\frac{1}{2}(S_2 + S_1)(y_2 - y_1) + \frac{1}{2}(S_3 + S_2)(y_3 - y_2) + \frac{1}{2}(S_1 + S_3)(y_1 - y_3) \right) \\ &= \frac{1}{2A_A} \left(S_1(y_2 - y_3) + S_2(y_3 - y_1) + S_3(y_1 - y_2) \right) \end{aligned} \quad (3.3)$$

$$\begin{aligned} (S_y)_A &= \frac{1}{A_A} \iint_{\text{cell } A} \frac{\partial S}{\partial y} dx dy \\ &= \frac{-1}{A_A} \int_{1-2-3} S dx \end{aligned}$$

$$\begin{aligned}
&= \frac{-1}{A_A} \left(\frac{1}{2}(S_2 + S_1)(x_2 - x_1) + \frac{1}{2}(S_3 + S_2)(x_3 - x_2) + \frac{1}{2}(S_1 + S_3)(x_1 - x_3) \right) \\
&= \frac{-1}{2A_A} \left(S_1(x_2 - x_3) + S_2(x_3 - x_1) + S_3(x_1 - x_2) \right) \quad (3.4)
\end{aligned}$$

A similar process is performed to create a second difference. The integration is taken around all the triangles which surround the node for which the second difference is computed, using the derivative values calculated at the cells. To get a second difference instead of a second derivative, there is no division by the area of the integrated region. The contribution to the second difference at node 1 from cell A is

$$\begin{aligned}
(D^2 S)_{1A} &= \int_{2-3} (S_x)_A dy - (S_y)_A dx \\
&= \frac{1}{2} \left[(S_x)_A (y_3 - y_2) - (S_y)_A (x_3 - x_2) \right] \quad (3.5)
\end{aligned}$$

This second difference operator takes about twice as long to compute as the low-accuracy second difference operator. It is also conservative since again the total contribution of each cell is zero.

To formulate a second difference operator for a quadrilateral mesh a similar process is employed. It turns out that if the first derivative is found by integrating around the complete quadrilateral, the oscillatory modes are not damped out, and the primary purpose of the operator is not fulfilled. To prevent this problem, the quadrilateral is broken into triangles and the triangular operator is applied. The division of cell A is shown in Figure 3.4. For cell A the first derivatives for the triangle corresponding to node 1 is

$$\begin{aligned}
(S_x)_{A1} &= \frac{1}{A_{A1}} \iint_{\text{cell A}} \frac{\partial S}{\partial x} dx dy \\
&= \frac{1}{A_{A1}} \int_{1-2-4} S dy \\
&= \frac{1}{A_{A1}} \left(\frac{1}{2}(S_2 + S_1)(y_2 - y_1) + \frac{1}{2}(S_4 + S_2)(y_4 - y_2) + \frac{1}{2}(S_1 + S_4)(y_1 - y_4) \right)
\end{aligned}$$

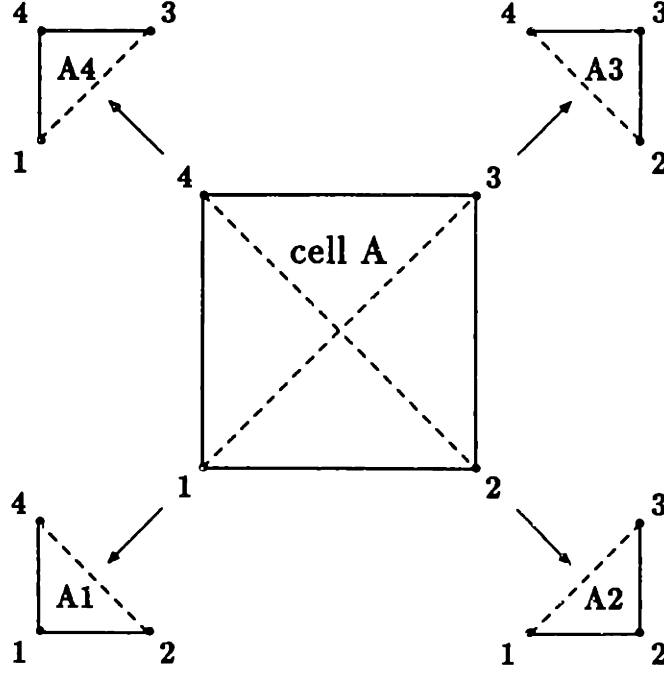


Figure 3.4: Division of quadrilateral cell for non-conservative, high-accuracy second difference

$$= \frac{1}{2A_{A1}} \left(S_1(y_2 - y_4) + S_2(y_4 - y_1) + S_4(y_1 - y_2) \right) \quad (3.6)$$

$$\begin{aligned} (S_y)_{A1} &= \frac{1}{A_{A1}} \iint_{\text{cell A}} \frac{\partial S}{\partial y} dx dy \\ &= \frac{-1}{A_{A1}} \int_{1-2-4} S dx \\ &= \frac{-1}{A_{A1}} \left(\frac{1}{2}(S_2 + S_1)(x_2 - x_1) + \frac{1}{2}(S_4 + S_2)(x_4 - x_2) + \frac{1}{2}(S_1 + S_4)(x_1 - x_4) \right) \\ &= \frac{-1}{2A_{A1}} \left(S_1(x_2 - x_4) + S_2(x_4 - x_1) + S_4(x_1 - x_2) \right) \end{aligned} \quad (3.7)$$

where A_{A1} refers to the area of the triangle corresponding to node 1 for cell A.

Next the integral is taken around the outside edge of the triangle to give the contribution of cell A1 to the second difference at node 1.

$$\begin{aligned}(D^2S)_{1A} &= \int_{2-4} (S_x)_{A1} dy - (S_y)_{A1} dx \\ &= \frac{1}{2} \left[(S_x)_{A1} (y_4 - y_2) - (S_y)_{A1} (x_4 - x_2) \right] \quad (3.8)\end{aligned}$$

Similar contributions to corresponding nodes are found for the other three triangles into which cell A is divided. As with the triangular version of this operator it takes about twice as long to compute this operator as the low-accuracy second difference operator. For the quadrilateral mesh this second difference is not conservative since the total contribution from each cell is not zero since the first derivative contribution is not constant in the cell.

3.2 Freestream Smoothing

To damp out oscillations and provide stability a fourth difference operator is added throughout the flow field. It is desirable to have a conservative operator so all terms in the flow field cancel and therefore no mass, momentum or energy production occurs due to the smoothing.

The first method of creating a fourth difference is to use the low-accuracy second difference twice by operating first on the state vector and then operating on this second difference. This fourth difference is conservative since the contribution of each cell to the numerical smoothing is zero, but is only second order accurate on a uniform mesh since the second difference operator used is only second order accurate on a uniform mesh. The second method is to compute a second difference of the state vector using the high-accuracy method and operate on this second difference with the low-accuracy second difference. This operator is second order accurate since the first operator is second order accurate and conservative since the second operator is conservative and again the contribution of each cell to the numerical smoothing is zero. The second

method is more expensive than the first, but the effect per iteration is an increase of only 5-10% which is a small increase for the gain in accuracy.

The fourth difference is computed as a contribution from each cell to the nodes which make up that cell. The freestream smoothing is multiplied by a coefficient $\epsilon^{(1)}$, between 0.0 and 0.05, to control the amount of smoothing added to the scheme. To make the numerical smoothing term consistent with the numerical scheme, the fourth difference added to the change in the state vector at each node must be multiplied by the ratio of $(\frac{\Delta t}{A})$ for the node to $(\frac{\Delta t}{A})$ for the cell as described by Roberts in [18]. For the quadrilateral Ni scheme in Equation (2.9) and for the triangular Ni scheme in Equation (2.17) the change to the state vector due to cell A at node 1 becomes

$$\delta U_{1A} = \delta U_{1A} + \epsilon^{(1)} \left(\frac{\Delta t}{A} \right)_1 \left(\frac{A}{\Delta t} \right)_A (D^2(D^2U))_{1A} \quad (3.9)$$

The value of the flux contribution for the quadrilateral Jameson scheme in Equation (2.24) and the triangular Jameson scheme in Equation (2.27) is changed such that the contribution from cell A to node 1 becomes

$$\text{flux}_{1A} = \text{flux}_{1A} + \epsilon^{(1)} \left(\frac{A}{\Delta t} \right)_A (D^2(D^2U))_{1A} \quad (3.10)$$

The flux terms are multiplied by $(\frac{\Delta t}{A})$ for each node when the change in the state vector is computed in Equation (2.20), so the smoothing term is consistent with the numerical scheme. The numerical smoothing term for the Jameson scheme is only computed after the first two of the four steps.

3.3 Shock Smoothing

In regions with strong discontinuities the fourth difference smoothing is not enough to cause any of the schemes mentioned here to be stable. When strong discontinuities are detected the freestream smoothing is turned off since it is destabilizing, and shock smoothing is turned on. For the Jameson schemes the shock smoothing consists of a

low-accuracy second difference which is turned on in regions of high pressure gradient. Another method is used for the Ni schemes which adds a bulk viscosity term in regions of high velocity flux.

3.3.1 Second Difference Smoothing

To determine when to turn on the second difference smoothing a pressure switch is used. This switch is found for each node and consists of the second difference of pressure computed using the low-accuracy operator as shown in Equation (3.1) for triangular meshes and Equation (3.2) for quadrilateral meshes and divided by the pressure at the node. Near shocks this switch will be of order 1 but in the freestream it will be of order Δx^2 . For node 1 this switch is

$$(s_p)_1 = \frac{D^2 p}{p_1} \quad (3.11)$$

Once the switch is found it is used when finding the second difference of the state vector at the nodes which is computed using an operator similar to the low-accuracy operator and multiplied by a coefficient, $\epsilon^{(2)}$, between 0.0 and 0.1.

For the triangular scheme the value of the flux contribution in Equation (2.27) is changed such that the contribution from cell A to node 1 becomes

$$\begin{aligned} \text{flux}_{1A} = \text{flux}_{1A} + \epsilon^{(2)} \frac{1}{2} \left[\left((s_p)_1 + (s_p)_2 \right) (U_2 - U_1) \right. \\ \left. + \left((s_p)_1 + (s_p)_3 \right) (U_3 - U_1) \right] \end{aligned} \quad (3.12)$$

Similarly for the quadrilateral scheme the flux contribution in Equation (2.24) is changed such the contribution from cell A to node 1 becomes

$$\begin{aligned}
\text{flux}_{1A} = \text{flux}_{1A} + \epsilon^{(2)} \frac{1}{2} \left[\right. & \left((s_p)_1 + (s_p)_2 \right) (U_2 - U_1) \\
& + \left((s_p)_1 + (s_p)_3 \right) (U_3 - U_1) \\
& \left. + \left((s_p)_1 + (s_p)_4 \right) (U_4 - U_1) \right] \quad (3.13)
\end{aligned}$$

3.3.2 Bulk Viscosity Smoothing

To capture shocks, viscosity is added near shocks since viscosity would capture a shock if it were accounted for in the governing equations. This method is similar to the method described by Richtmyer and Morton in [17]. To ensure that the shock width remains nearly the same, regardless of shock strength, terms quadratic in the strain rate are added to the differential equation. The volumetric dilatation, or the velocity divergence, is a measure of the strain rate. In shocks the volumetric dilatation is highly negative since shocks represent regions of extreme compression. The shock smoothing is turned on when the volumetric dilatation is negative and is limited such that it is never less than -0.1. The viscosity term which is added to the flux vectors is proportional to the volumetric dilatation squared and, to prevent excessive smoothing at stagnation points, the velocity squared. The change in the flux vectors, ΔF and ΔG now have the viscosity term added to their second and third elements respectively.

$$\begin{aligned}
(\Delta F_2)_A &= (\Delta F_2)_A + \frac{1}{2} \max(-0.1, \min(0., \text{div} \bar{w}_A)) \text{div} \bar{w}_A (u^2 + v^2) \\
(\Delta G_3)_A &= (\Delta G_3)_A + \frac{1}{2} \max(-0.1, \min(0., \text{div} \bar{w}_A)) \text{div} \bar{w}_A (u^2 + v^2)
\end{aligned} \quad (3.14)$$

For triangles the volumetric dilatation for cell A can be found by

$$\begin{aligned}
\text{div} \bar{w}_A &= \text{scaled volumetric dilatation of triangular cell A} \\
&= \sqrt{\frac{2}{A}} \iint_{\text{cell A}} \left(\frac{\partial u}{\partial x} + \frac{\partial v}{\partial y} \right) dx dy
\end{aligned}$$

$$\begin{aligned}
&= \sqrt{\frac{2}{A}} \oint_{1-2-3} (u dy - v dx) \\
&= \frac{1}{\sqrt{2A}} \left(u_1(y_2 - y_3) + u_2(y_3 - y_1) + u_3(y_1 - y_2) \right. \\
&\quad \left. - (v_1(x_2 - x_3) + v_2(x_3 - x_1) + v_3(x_1 - x_2)) \right) \tag{3.15}
\end{aligned}$$

and similarly for quadrilaterals

$$\begin{aligned}
\text{div} \vec{w}_A &= \text{scaled volumetric dilatation of quadrilateral cell A} \\
&= \sqrt{\frac{2}{A}} \iint_{\text{cell A}} \left(\frac{\partial u}{\partial x} + \frac{\partial v}{\partial y} \right) dx dy \\
&= \sqrt{\frac{2}{A}} \oint_{1-2-3-4} (u dy - v dx) \\
&= \frac{1}{\sqrt{2A}} \left((u_1 - u_3)(y_2 - y_4) + (u_2 - u_4)(y_3 - y_1) \right. \\
&\quad \left. - (v_1 - v_3)(x_2 - x_4) + (v_2 - v_4)(x_3 - x_1) \right) \tag{3.16}
\end{aligned}$$

Chapter 4

Mesh Generation and Pointer System

In this chapter the mesh generation and the system for storing the information in the mesh are described. An elliptic mesh generator was used to generate a quadrilateral mesh and, since the development of a triangular mesh generator was not a major focus of this work, the quadrilaterals in this mesh were divided into triangles to generate a triangular mesh.

For meshes consisting of a regular arrangement of quadrilateral cells, the the nodes of the mesh have traditionally been described in two dimensions by an (i, j) indexing. An alternative to this method of describing a mesh is to assign each node a number and describe the connection between the nodes by one dimension of an array of mesh elements. These elements can consist of cells, faces, edges or any other element which is important in the numerical scheme. This system of describing a mesh will be referred to as a pointer system. When a pointer system is used the flow solver is separated from the mesh generator. Due to the inherent irregularity of most triangular meshes, it is usually not possible to use the first method of describing a mesh and a pointer system must be used. All the meshes used for this study are described using a pointer system.

4.1 Elliptic Mesh Generator

Elliptic partial differential equations are used to generate a smooth mesh. The equations are defined in a computational plane with coordinates ξ and η whose nodes have a one to one mapping to the physical plane. The equations used are

$$\xi_{xx} + \xi_{yy} = P(\xi, \eta) \quad (4.1)$$

$$\eta_{xx} + \eta_{yy} = Q(\xi, \eta) \quad (4.2)$$

where P and Q are forcing terms defined by Steger and Sorenson [21]. By switching the independent and dependent variables these equations become

$$\alpha x_{\xi\xi} - 2\beta x_{\xi\eta} + \gamma x_{\eta\eta} = -J^2(Px_\xi + Qx_\eta) \quad (4.3)$$

$$\alpha y_{\xi\xi} - 2\beta y_{\xi\eta} + \gamma y_{\eta\eta} = -J^2(Py_\xi + Qy_\eta) \quad (4.4)$$

where

$$J = x_\xi y_\eta - x_\eta y_\xi$$

$$\alpha = x_\eta^2 + y_\eta^2$$

$$\beta = x_\xi x_\eta + y_\eta y_\xi$$

$$\gamma = x_\xi^2 + y_\xi^2$$

An H-mesh in a duct or blade cascade is created where the line $\xi = 0$ corresponds to the inlet boundary and the line $\xi = 1$ corresponds to the outlet boundary. The lines $\eta = 0$ and $\eta = 1$ correspond to the upper and lower surface of the domain.

4.2 Complete Pointer System

To assure that the constraints of the pointer system would not restrict the code development process, a very complete pointer system was put together. This system stores three to four times as much information as is required to program the numerical schemes described here. The minimum requirements are described in the following section.

The pointer system used here consists of arrays of nodes, cells, faces, edge faces and edge nodes. These arrays are interconnected in that elements in one array will point

to elements in another array. Nodes are the only elements which do not point to other elements. The interconnection of the arrays is shown in Figure 4.1 where the arrows indicate pointing from one array to another. In essence the complete pointer system overdefines the connection between the arrays, but this allowed more freedom in the code development process.

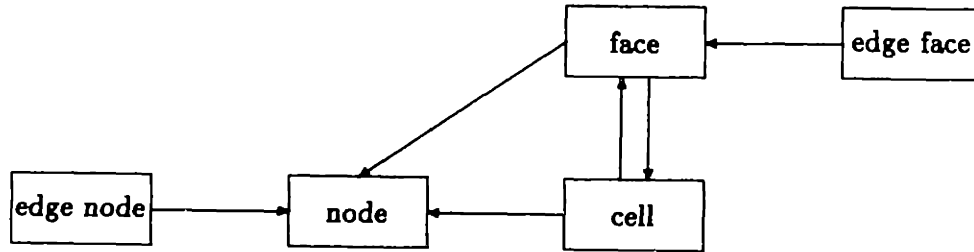


Figure 4.1: Interconnection of pointer arrays

4.2.1 Node Arrays

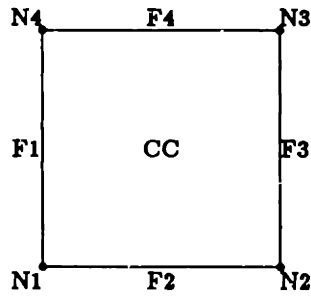
Each node is assigned a number. Arrays containing all the nodes hold information about the x and y coordinate as well as the four elements of the state vector U for each node. These variables were stored in

$$\begin{array}{ll}
 x(N) & N = 1, N_{max} \\
 y(N) & N = 1, N_{max} \\
 U(i, N) & N = 1, N_{max} \\
 & i = 1, 4
 \end{array}$$

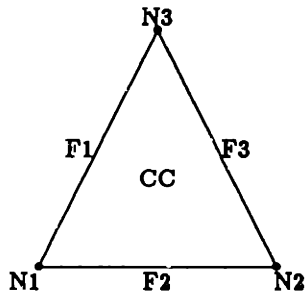
Other information is temporarily stored at each node, such as the flux vectors F and G and the change in the state vector δU .

4.2.2 Cell Arrays

The cell is the basic computational unit for all the schemes described here. The value of a flux, for example, is calculated for the cell and distributed to the nodes which make up the cell. To completely describe the cell and its surrounding elements, each cell points to the nodes and faces which make up the cell. The cell array for cell CC consists of



```
cell(1, CC) = F1
cell(2, CC) = F2
cell(3, CC) = F3
cell(4, CC) = F4
cell(5, CC) = N1
cell(6, CC) = N2
cell(7, CC) = N3
cell(8, CC) = N4
```

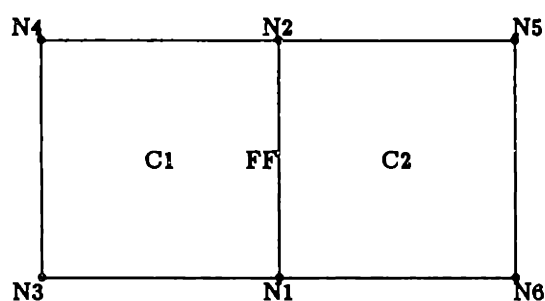


```
cell(1, CC) = F1
cell(2, CC) = F2
cell(3, CC) = F3
cell(4, CC) = N1
cell(5, CC) = N2
cell(6, CC) = N3
```

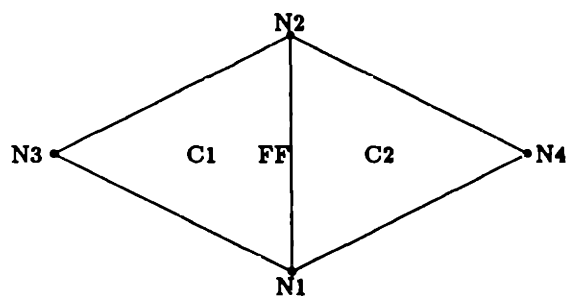
As with the nodal arrays, other information is temporarily stored at each cell, such as the area divided by the timestep ($\frac{A}{\Delta t}$) and the change in the state vector at the cell ΔU . This information can be accessed by merely knowing the cell number.

4.2.3 Face Arrays

At one time, using the face as the basic computational element was considered. This method is effective for the Jameson schemes mentioned here, but for consistency between the Ni and Jameson schemes was replaced by the cell based method. Currently the face elements are used to indirectly describe edge faces and plot the computational mesh. Each face points to the cells on either side of it and the nodes which belong to these cells. When a node or face lies outside the computational domain its value is set to zero. The face array for face FF consists of



$\text{face}(1, \text{FF}) = \text{C1}$
 $\text{face}(2, \text{FF}) = \text{C2}$
 $\text{face}(3, \text{FF}) = \text{N1}$
 $\text{face}(4, \text{FF}) = \text{N2}$
 $\text{face}(5, \text{FF}) = \text{N3}$
 $\text{face}(6, \text{FF}) = \text{N4}$
 $\text{face}(7, \text{FF}) = \text{N5}$
 $\text{face}(8, \text{FF}) = \text{N6}$



$\text{face}(1, \text{FF}) = \text{C1}$
 $\text{face}(2, \text{FF}) = \text{C2}$
 $\text{face}(3, \text{FF}) = \text{N1}$
 $\text{face}(4, \text{FF}) = \text{N2}$
 $\text{face}(5, \text{FF}) = \text{N3}$
 $\text{face}(6, \text{FF}) = \text{N4}$

4.2.4 Edge Face Arrays

To implement the boundary conditions it is necessary to know which faces or cells lie on the boundary and their orientation with respect to the boundary. The elements

of the edge face array point to the elements of the face array. Each face which is on the boundary is oriented such that C1 and N3 (and N4 for the quadrilaterals) lie inside the domain.

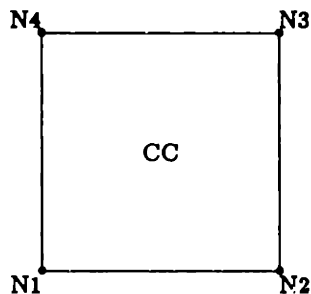
$$\text{edge}(\text{EF}) = \text{FF}$$

4.2.5 Edge Node Arrays

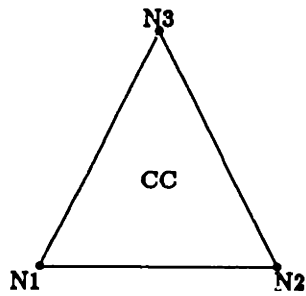
For convenience an array of edge nodes was also created. These nodes correspond to the solid wall boundary nodes. Information such as the flow angle α at a boundary node was stored in an array indexed by the edge nodes.

4.3 Required Pointer System

The pointer system described in the previous section contains more information than is required to implement the numerical schemes described here. The nodal arrays are the independent variables of the system and should remain as they were described. The rest of the pointer system can be replaced by a single array of cells. To specify which cells lie on the boundaries the edge cells can be placed at the beginning of the array with the first two nodes of these cells lying on the boundary. The cell array should be arranged such that the entry for cell CC would consist of



$$\begin{aligned} \text{cell}(1, \text{CC}) &= \text{N1} \\ \text{cell}(2, \text{CC}) &= \text{N2} \\ \text{cell}(3, \text{CC}) &= \text{N3} \\ \text{cell}(4, \text{CC}) &= \text{N4} \end{aligned}$$



$\text{cell}(1, \text{CC}) = \text{N1}$
 $\text{cell}(2, \text{CC}) = \text{N2}$
 $\text{cell}(3, \text{CC}) = \text{N3}$

4.4 Vectorization

To eliminate data dependencies which restrict vectorization and to create long vector lengths, a coloring system is used. This system assigns a color to each element in the array such that all elements of the same color do not contain data dependencies. The arrays are then sorted such that all elements of the same color occupy consecutive locations in the array. For example, data dependencies may occur in an operation on an array of cells when two different cells contain a node in common. For quadrilateral cells it is possible for simple cases to color the cell array with four colors such that the cells of one color do not have common nodes.

All arrays in the pointer system used here were colored so the code would run on an Alliant FX/3 computer which used vector/concurrent arithmetic.

Chapter 5

Computational Examples

To demonstrate the similarities and differences between the numerical methods described here, some examples will be shown. Little reference will be made to computational times since in practice multi-grid or adaptation will be used to accelerate the computational time. The convergence rate is also related to the numerical smoothing used for freestream and shock capturing.

A discussion on the accuracy of the numerical schemes discussed here is given in Chapter 6. In general the accuracy of a numerical method is reduced by using the low-accuracy smoothing as opposed to the high-accuracy smoothing.

The regular and irregular quadrilateral meshes used here are similar to the meshes shown in Figures 6.1 and 6.2. The irregular triangular meshes are similar to the mesh shown in Figure 6.3.

5.1 Supersonic Circular Arc Bump

The first problem is supersonic flow in a channel with a 4% circular arc bump on the lower surface. The inlet Mach number is 1.4. A 64×16 cell mesh is used. This problem illustrates the effect of the shock capturing methods used as well as interactions between shocks and solid wall boundaries, and shocks with other shocks.

The quadrilateral and triangular Jameson and Ni schemes with the high-accuracy numerical smoothing required essentially the same amount of computational time to converge from a uniform freestream initial condition. Slightly less time was required when the low-accuracy numerical smoothing was used since it is less expensive to compute

than the high-accuracy smoothing. Essentially twice as many iterations were required for the Ni schemes than the Jameson schemes since the CFL number for the Ni schemes is half as large as for the Jameson schemes.

Several Mach number contours with increment 0.05 are shown in Figures 5.1 to 5.7 for different numerical schemes and numerical smoothing techniques. For the most part the solutions look the same.

In Figure 5.1 the effect of the low-accuracy smoothing on an irregular mesh can be seen. Comparing the solution in Figure 5.1 to the solutions in Figure 5.2 with the low-accuracy smoothing on a regular mesh and Figure 5.3 with the high-accuracy smoothing on an irregular mesh, the contours are not as smooth. This effect exists because the low-accuracy smoothing includes no information about the location of the neighboring mesh points. For an irregular mesh this becomes important. In Figure 5.1 another effect of the low-accuracy smoothing can be seen. The contours on the lower surface do not intersect the solid wall smoothly. A slight turning of the contour lines can be seen. This effect exists because the inviscid solution has $\frac{\partial u}{\partial n} \neq 0$ on curved walls, whereas the low-accuracy smoothing has a one-sided bias which implicitly assumes that $\frac{\partial u}{\partial n} = 0$.

All the solutions pick up the normal shock on the upper surface of the duct in the reflection from the leading edge shock. This reflection interacts with the trailing edge shock behind the bump, and reflects off the lower surface of the duct to combine with the trailing edge shock. In general the quadrilateral schemes pick up this interaction between the shocks better than the triangular schemes. This is partly due to the resolution since as the mesh is refined, the shock is picked up by the triangular schemes as well as the quadrilateral schemes. The triangular schemes have a harder time keeping the leading edge shock and its first reflection straight. More work must be done with the shock capturing smoothing to correct this.

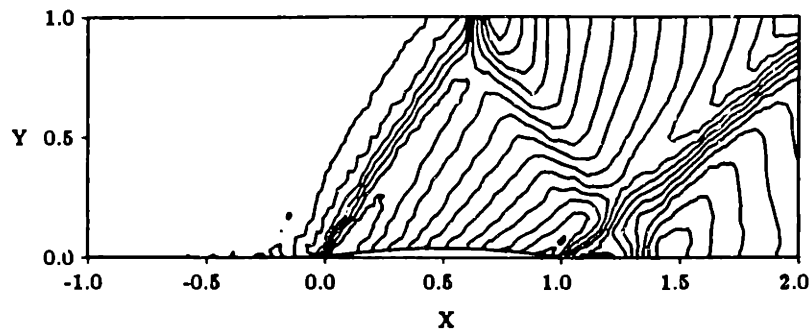


Figure 5.1: Supersonic case: quadrilateral Jameson scheme with low-accuracy smoothing on an irregular mesh

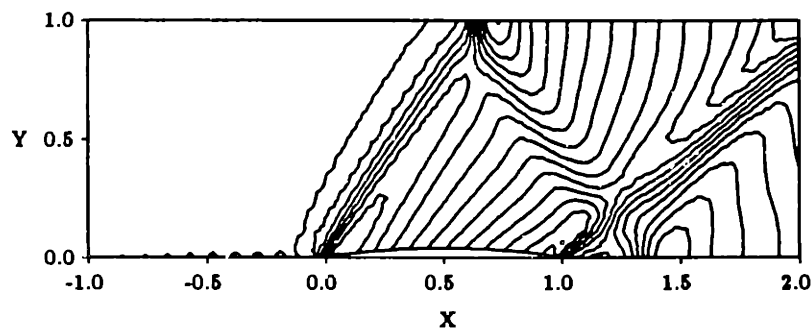


Figure 5.2: Supersonic case: quadrilateral Jameson scheme with low-accuracy smoothing on a regular mesh

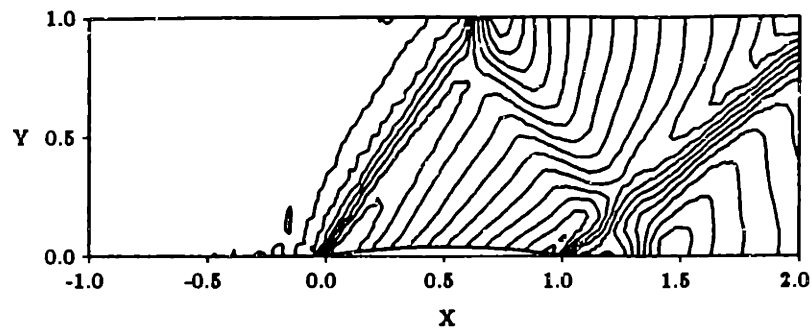


Figure 5.3: Supersonic case: quadrilateral Jameson scheme with high-accuracy smoothing on an irregular mesh

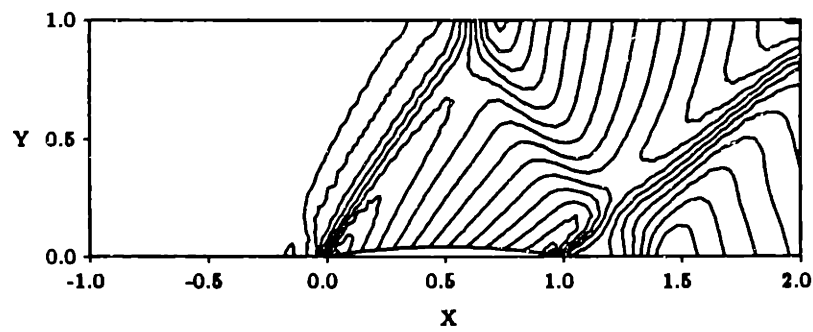


Figure 5.4: Supersonic case: quadrilateral Ni scheme with high-accuracy smoothing on a regular mesh

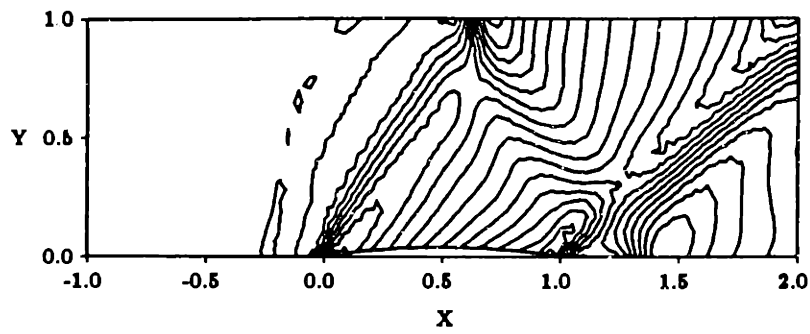


Figure 5.5: Supersonic case: triangular Jameson scheme with high-accuracy smoothing on an irregular mesh

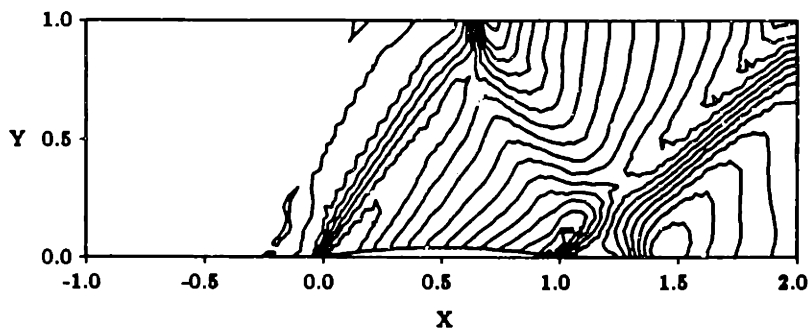


Figure 5.6: Supersonic case: triangular Jameson scheme with low-accuracy smoothing on an irregular mesh

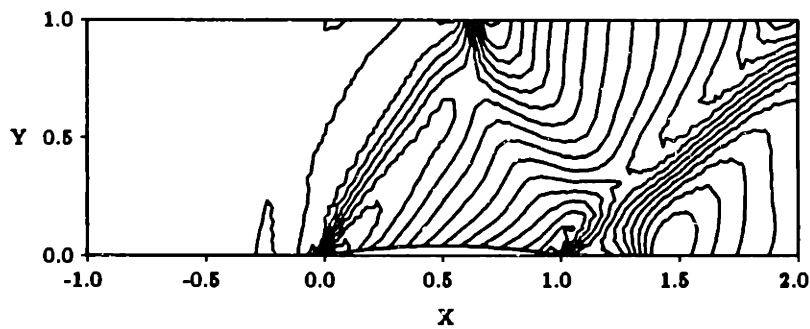


Figure 5.7: Supersonic case: triangular Ni scheme with high-accuracy smoothing on an irregular mesh

5.2 Transonic Circular Arc Bump

The second problem is transonic flow in a channel with a 10% circular arc bump on the lower surface. The inlet Mach number is 0.675. A 64×16 cell mesh is used. The inlet is subsonic, but the flow is accelerated over the top of the bump and a shock forms. This problem again illustrates the shock capturing methods as well as subsonic interactions with solid wall boundaries.

Several Mach number contours with increment 0.1 are shown in Figures 5.8 to 5.15 for different numerical schemes and numerical smoothing techniques. Again, for the most part the solutions look the same. Some plots show tighter shocks, but this is due to different shock capturing methods more than the numerical schemes themselves.

As with the supersonic test case, the solution found using the quadrilateral Jameson scheme with low-accuracy smoothing on an irregular mesh, as shown in Figure 5.8, is significantly less smooth than the same case on a regular mesh, as shown in Figure 5.9. The high-accuracy smoothing used with the quadrilateral Jameson scheme produces much smoother results with both regular and irregular meshes as shown in Figures 5.11 and 5.10. In fact the irregular mesh has a very smooth solution with the high-accuracy smoothing.

In Figure 5.13 an effect of the low-accuracy smoothing on the interaction with solid wall boundaries can be seen. The smoothing tends to keep the Mach contours from intersecting the solid wall smoothly. This can be seen by a turning of the contour lines near the wall. This effect can also be seen in for the supersonic case in Figure 5.1. Once again the lack of information about the location of the mesh points in the low-accuracy smoothing and the imposition of an unnatural boundary condition for the numerical smoothing causes a problem.

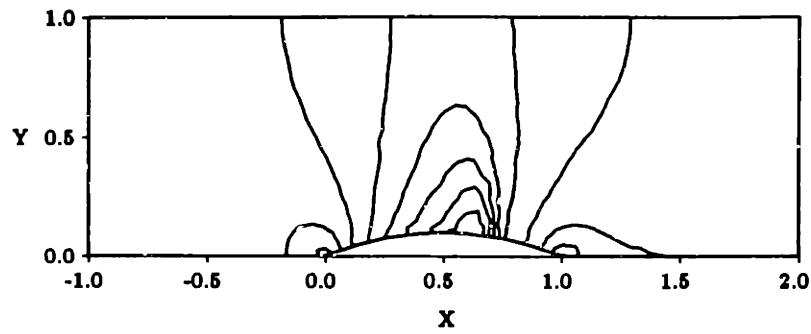


Figure 5.8: Transonic case: quadrilateral Jameson scheme with low-accuracy smoothing on an irregular mesh

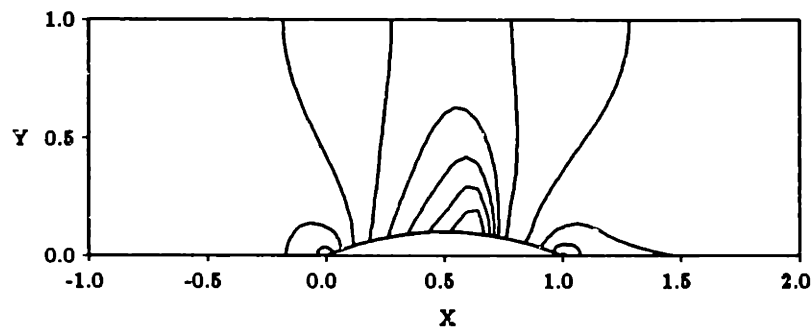


Figure 5.9: Transonic case: quadrilateral Jameson scheme with low-accuracy smoothing on a regular mesh

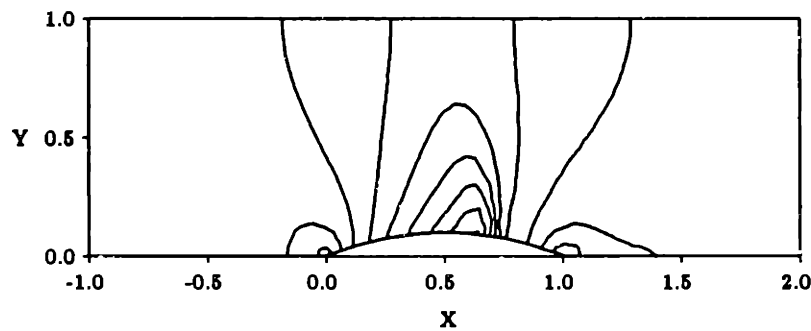


Figure 5.10: Transonic case: quadrilateral Jameson scheme with high-accuracy smoothing on an irregular mesh

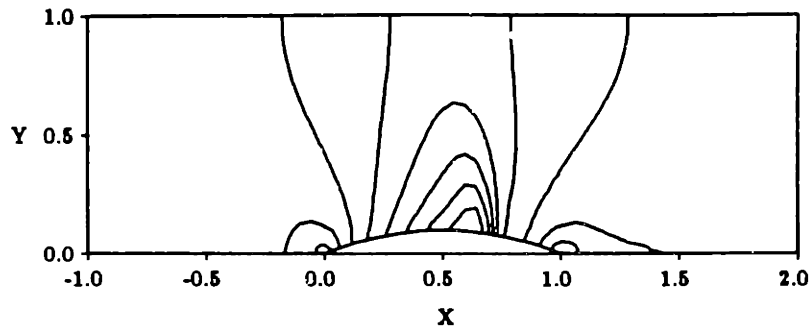


Figure 5.11: Transonic case: quadrilateral Jameson scheme with high-accuracy smoothing on a regular mesh

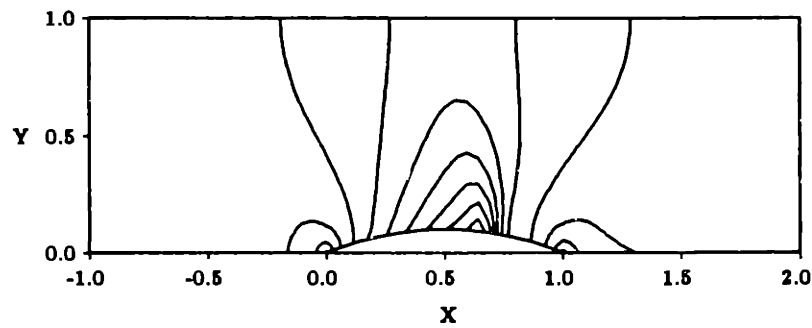


Figure 5.12: Transonic case: quadrilateral Ni scheme with high-accuracy smoothing on a regular mesh

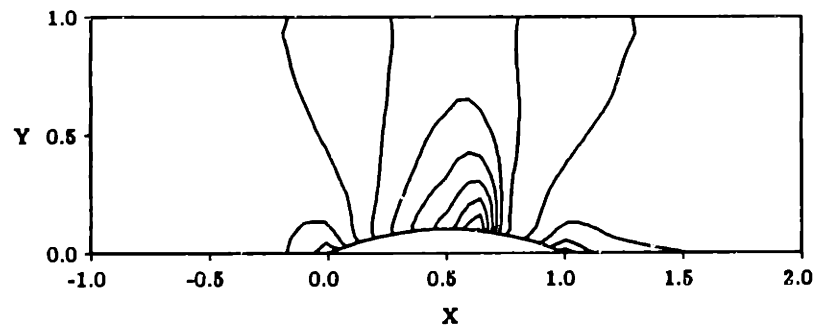


Figure 5.13: Transonic case: triangular Jameson scheme with low-accuracy smoothing on an irregular mesh

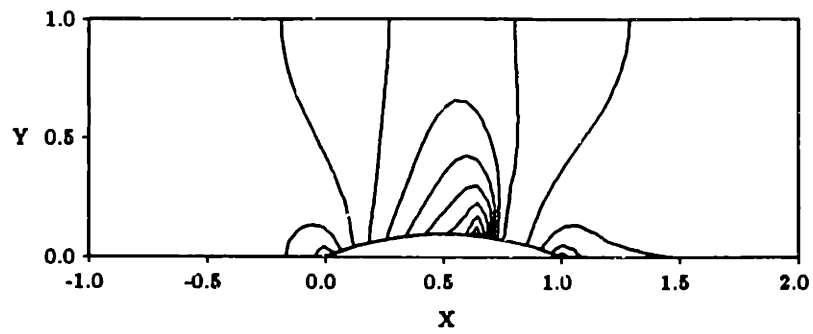


Figure 5.14: Transonic case: triangular Jameson scheme with high-accuracy smoothing on an irregular mesh

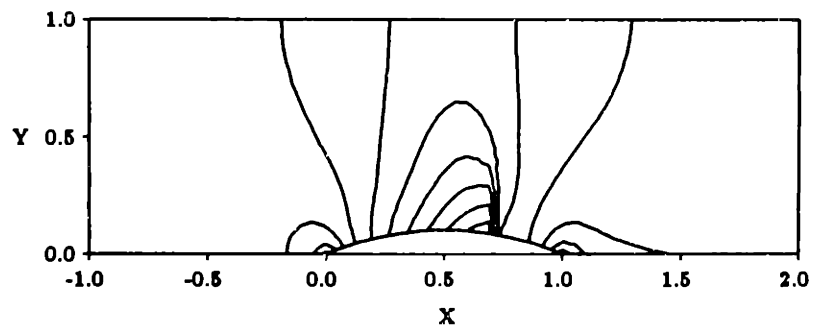


Figure 5.15: Transonic case: triangular Ni scheme with high-accuracy smoothing on an irregular mesh

Chapter 6

Accuracy Study

It is shown mathematically by Giles [5] that both the Ni scheme and the Jameson scheme are second order accurate for either quadrilateral or triangular meshes. The nature of the mathematical proof as well as a numerical accuracy study which confirms this result are examined in this chapter.

6.1 Mathematical Study

Giles shows in [5] that the global solution error is second order for steady state node based finite volume schemes on irregular meshes. The analysis states that the local truncation error is first order, but that this does not imply that the global error is also first order. It is assumed that the numerical finite volume scheme is conservative and h is some typical cell length. With this in mind, it is shown that the truncation error has a spectral content which can be split into two parts, a low-frequency component with an amplitude which is $O(h^2)$ and a high-frequency component with an amplitude which is $O(h)$. It is then shown that the hyperbolic differential operator

$$\mathcal{L}U = \frac{\partial F(U)}{\partial x} + \frac{\partial G(U)}{\partial y} \quad (6.1)$$

has a transfer function which is $O(1)$ at low frequencies and $O(1/h)$ at high frequencies. From this it can be concluded that the global error is $O(h^2)$ for both low and high frequencies, therefore the schemes are second order accurate. An important assumption in the analysis is that the numerical smoothing does not produce a truncation error which is worse than first order on irregular meshes.

6.2 Numerical Study

A numerical study was performed to determine the accuracy of the schemes described in Chapter 2. The purpose of this study was to confirm that these schemes are second order accurate.

For inviscid flow, there should be no total pressure loss for smooth, subsonic flow. Thus, for such flows any total pressure loss is purely numerical in nature. With this in mind, the total pressure loss is used to define the error in subsonic flow through a duct with a $\sin^2 x$ bump on the lower surface and a 0.50 inlet Mach number. The height of the lower surface is given by

$$y = 0.10 \sin^2(\pi x) \quad 0 \leq x \leq 1 \quad (6.2)$$

To describe the numerical error on a mesh a global error is defined as the L_2 -norm of this local error.

$$\eta = \frac{(p_o)_{upstream} - (p_o)_{local}}{(p_o)_{upstream}} \quad (6.3)$$

$$error = \sqrt{\frac{\sum_{i=1}^N \eta_i^2}{N}} \quad (6.4)$$

For meshes composed of quadrilateral cells it is easy to define an appropriate mesh for an accuracy study. To create a *regular quadrilateral mesh* a mesh is laid out in a rectangular duct with a $\sin^2 x$ bump on the lower surface which is 3 units long and 1 unit high where cells away from the bump are square. h , a typical cell length, is defined as the length of the cell faces. Four different meshes are used for the study with 8, 16, 32 and 64 faces per unit length. The second mesh in this series is shown in Figure 6.1. To create an *irregular quadrilateral mesh* the nodes of these meshes are perturbed by an amount determined by a sine function whose period has no relationship to the mesh. Four different meshes are again used for the study with 8, 16, 32 and 64 faces per unit length. The second mesh is shown in Figure 6.2.

Two different mesh types are used for triangular cells. The first type is an *irregular triangular mesh* and is identical to the meshes described for the quadrilateral cells where the cells are split along the shorter diagonal to create triangles. h is defined in the same manner as for the quadrilaterals since this defines a reference length for the cells. Again four different meshes are used with 8, 16, 32 and 64 faces per unit length. An example of the second mesh is shown in Figure 6.3. A second type of mesh was created which contains only triangles which are very nearly equilateral. This type of mesh is what could be referred to as a *regular triangular mesh*. The duct for this mesh is slightly wider than the previously described duct to facilitate the use of equilateral triangles. The dimensions of the domain are now $3 \times \frac{5\sqrt{3}}{8}$ units instead of 3×1 units. Three different meshes are used with 8, 16, and 32 faces per unit length. The second mesh is shown in Figure 6.4.

Mach contours are shown in Figure 6.5 for the quadrilateral mesh shown in Figure 6.1 which are computed using the quadrilateral Ni scheme. Figure 6.6 shows % total pressure loss contours for this flow. These are representative examples of solutions for this flow field regardless of the numerical scheme or mesh used.

The error is computed for each scheme on the previously defined sets of meshes. The order of accuracy for the scheme is found by plotting $\log(h)$ verses $\log(error)$ and finding the slope of the resulting line. Since the data will not exactly lie in a line, a least squares approximation was used to fit a line through the data points and compute the slope.

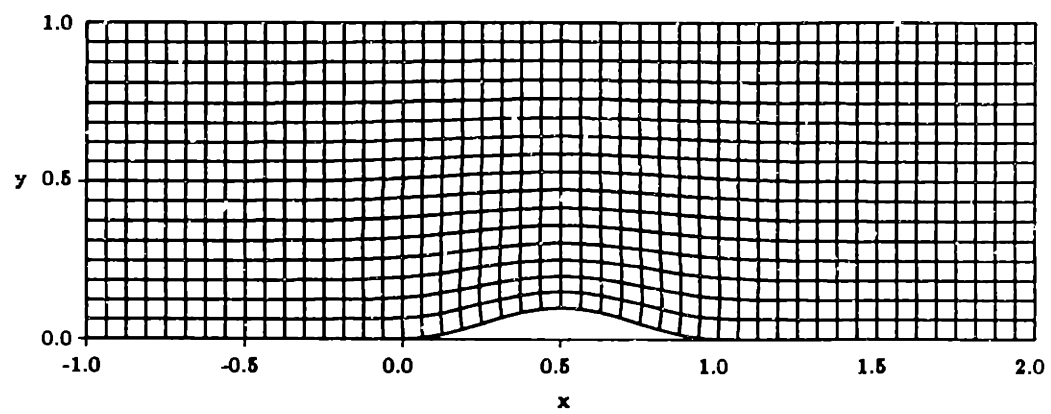


Figure 6.1: Quadrilateral cell mesh for $\sin^2 x$ duct

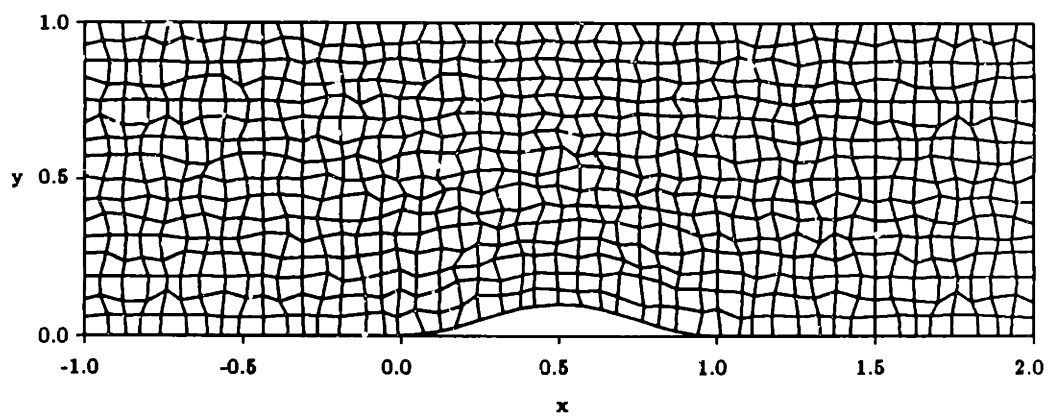


Figure 6.2: Irregular quadrilateral cell mesh for $\sin^2 x$ duct

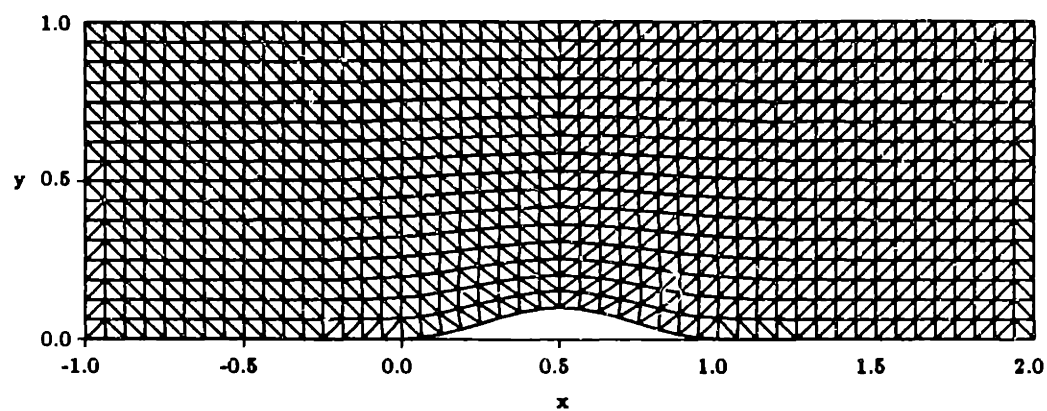


Figure 6.3: Split quadrilateral triangle cell mesh for $\sin^2 x$ duct

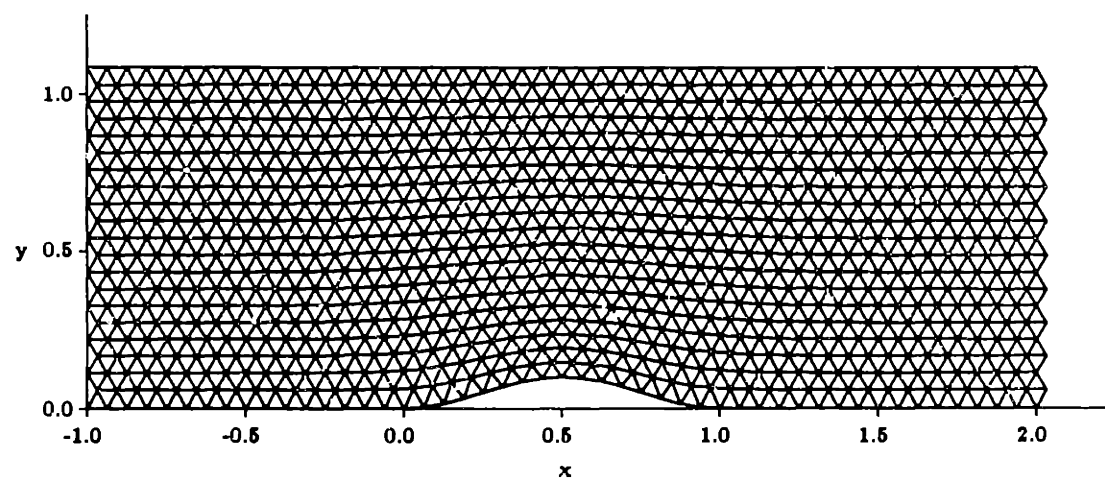


Figure 6.4: Equilateral triangle cell mesh for $\sin^2 x$ duct

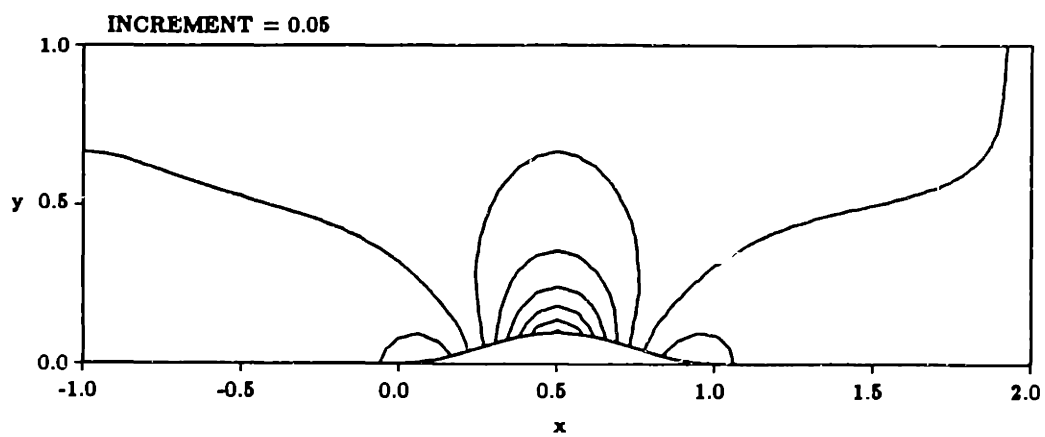


Figure 6.5: Mach contours for $\sin^2 x$ duct

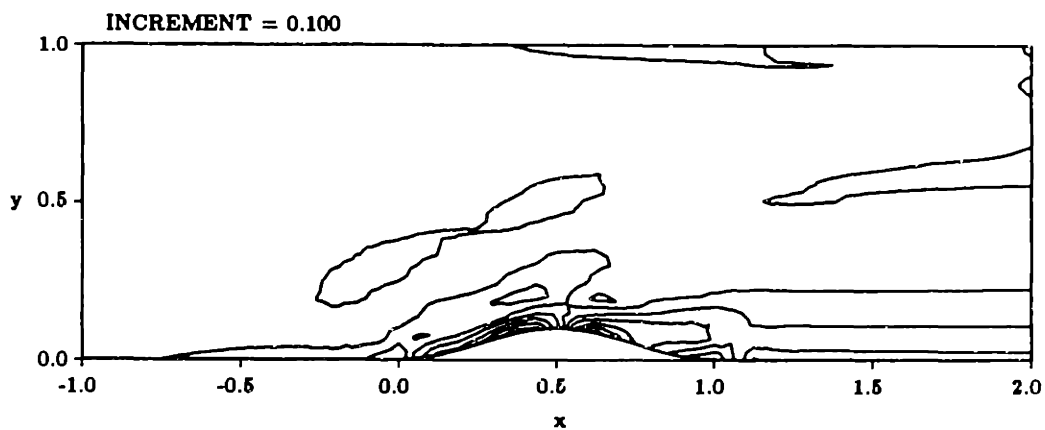


Figure 6.6: % total pressure loss contours for $\sin^2 x$ duct

6.3 Results of Numerical Study

For each of the numerical schemes described in Chapter 2 a numerical accuracy study was performed as described in the proceeding section. For the Jameson schemes two different numerical smoothing methods are used as described in Chapter 3. The schemes described here are all shown mathematically to be second order accurate. The order of accuracy of the numerical smoothing has an effect on the accuracy of the numerical scheme. This effect will be described in this section.

In the plots of the numerical accuracy data shown in Figures 6.7 to 6.14 it can be seen that not all points lie on a straight line. For all points the smoothing coefficient is the same, and the effect of this constant coefficient varies as h varies. In general, the points lie on a line and a least squares fit to the points is a good approximation to find the slope of this line. When fitting a line through the data there is some error due to a limited number of points which are used. Some sets of data consist of three points and others consist of four points, but the error is obviously small and depends on the particular case. It is also possible for the numerical accuracy to be more than 2, which is the numerical accuracy of the basic scheme, since the truncation errors are not composed of terms exactly proportional to h^2 , but will also contain higher order terms. This will be seen in the results presented here.

6.3.1 Effect of Numerical Smoothing on Accuracy

Two numerical fourth difference freestream smoothing methods were described in Chapter 3 for both the triangular and quadrilateral schemes. The first was a low-accuracy method and the second was a high-accuracy method, both of which are conservative. In the accuracy studies only freestream smoothing which is applied throughout the flow field was used. This is sufficient for stability since the problem used for the accuracy study was designed to have sufficiently smooth flow. For the Ni schemes only the high-accuracy smoothing was used. However, to illustrate the effect of smoothing on accuracy, both freestream methods were used for the Jameson schemes.

With the high-accuracy smoothing, the data for both the triangular and quadrilateral schemes show that these schemes are second order accurate on both regular and irregular meshes. This effect can be seen for the triangular Ni scheme by referring to Figures 6.7 and 6.8, and for the quadrilateral Ni scheme by referring to Figures 6.9 and 6.10. For the triangular Jameson scheme this can be seen by referring to Figures 6.11 and 6.12, and for the quadrilateral Jameson scheme by referring to Figures 6.13 and 6.14. Second order accuracy is also achieved for smoothing coefficient values larger than the values used here.

For the low-accuracy smoothing the story is different than when using the high-accuracy smoothing. The Jameson schemes require some smoothing to be stable. With a coefficient which is large enough to barely provide stability, the Jameson schemes may retain their second order accuracy. It is important to note that the flow field here is very smooth, and the value of the smoothing coefficient which barely provides stability here is smaller than it would be in most flow fields. With a smoothing coefficient a factor of five times larger the accuracy drops. In the first case the error due to the numerical scheme which is second order accurate dominates over the error due to the numerical smoothing. The order of accuracy which is calculated is effectively the order of accuracy of the numerical scheme without the smoothing. As the mesh resolution increases, eventually the smoothing error will dominate and the order of accuracy will deteriorate. Formally, order of accuracy is concerned with the limit $h \rightarrow 0$, and in this limit it will be less than second order. The beginning of this effect can be seen in Figure 6.22. As the smoothing coefficient is increased the accuracy is reduced and the error due to the numerical smoothing begins to play a role in the accuracy. The accuracy decreases for the regular as well as irregular meshes with the low-accuracy smoothing due to effects on the boundary. Dissipative errors occur on the boundary because the solid wall boundary condition which is imposed is not appropriate for the low-accuracy smoothing. The error can be estimated by noting that the smoothing attempts to enforce $\frac{\partial u}{\partial n} = 0$, and so it will create a numerical boundary layer at least one cell wide with an error that is of order $h(\frac{\partial u}{\partial n})_{\text{inviscid}} = O(h)$. The numerical boundary layer has $O(1/h)$ nodes compared to the total number of nodes which is $O(1/h^2)$, so a lower bound on the order of magnitude of the root mean square error is

$$error = O\left(\sqrt{\frac{1}{h}h^2/\frac{1}{h^2}}\right) = O(h^{3/2}) \quad (6.5)$$

This explains why the numerical error reduces to approximately $O(h^{3/2})$. The effect of increasing the smoothing coefficient can be seen for the triangular Jameson scheme on a regular mesh by referring to Figure 6.15, and Figure 6.16 for high and low smoothing coefficients respectively. The lowest smoothing coefficient which produces stability for an irregular triangular mesh would not give second order accuracy as shown in Figure 6.18. For the quadrilateral Jameson scheme this effect can be seen for both regular and irregular meshes by referring to Figures 6.19 and 6.20, and Figure 6.21 and 6.22 for high and low smoothing coefficients respectively.

The numerical smoothing formulation used is very important. The accuracy of the scheme will only be as good as the least accurate component, be that the basic numerical scheme or the numerical smoothing. If the numerical smoothing is only first order accurate, then the accuracy of the numerical scheme will be contaminated by the smoothing.

6.3.2 Conclusions

In Figure 6.23 the results of the numerical accuracy study are summarized. For the high-accuracy numerical smoothing, both the Ni and Jameson schemes for both quadrilateral and triangular meshes are second order accurate. This applies for both regular and irregular meshes. On relatively coarse meshes, when the smoothing coefficient for the low-accuracy smoothing is at the minimum required to keep a scheme stable the Jameson schemes for both quadrilateral and triangular meshes are approximately second order accurate. When this coefficient is above this minimum or when much finer meshes are used the accuracy of the basic numerical scheme is contaminated and the accuracy drops below second order.

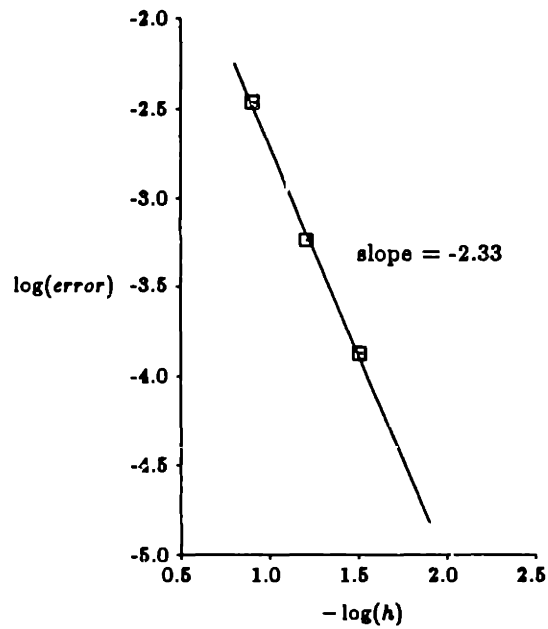


Figure 6.7: Order of accuracy for triangular Ni scheme with high-accuracy smoothing on a regular mesh

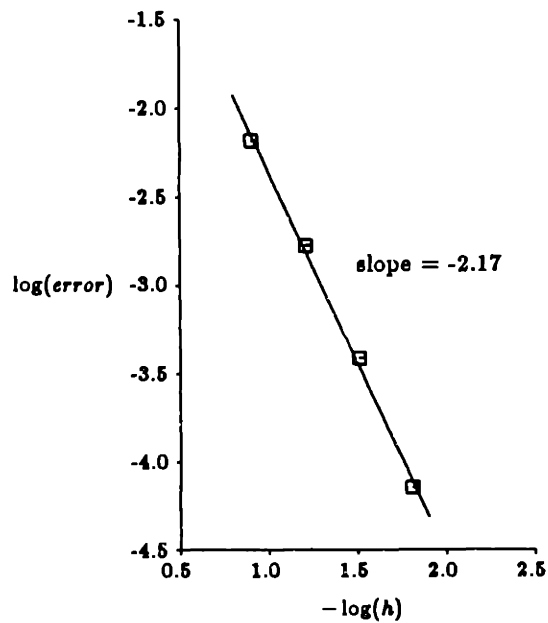


Figure 6.8: Order of accuracy for triangular Ni scheme with high-accuracy smoothing on an irregular mesh

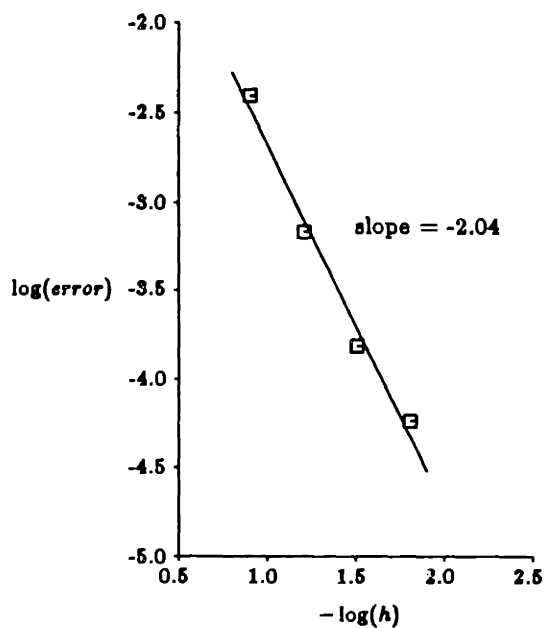


Figure 6.9: Order of accuracy for quadrilateral Ni scheme with high-accuracy smoothing on a regular mesh

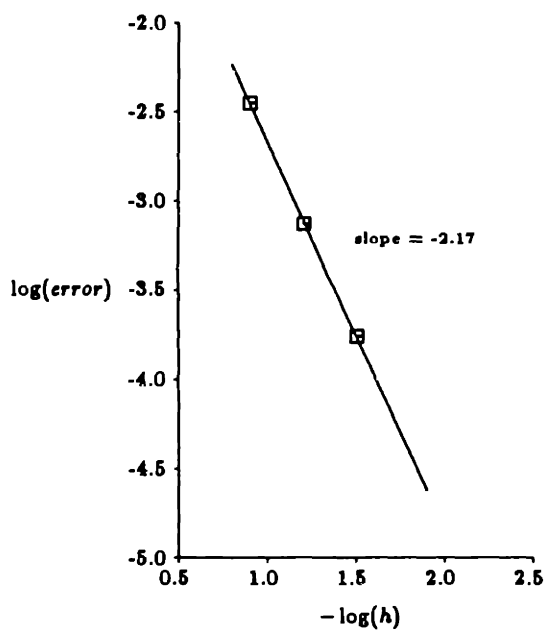


Figure 6.10: Order of accuracy for quadrilateral Ni scheme with high-accuracy smoothing on an irregular mesh

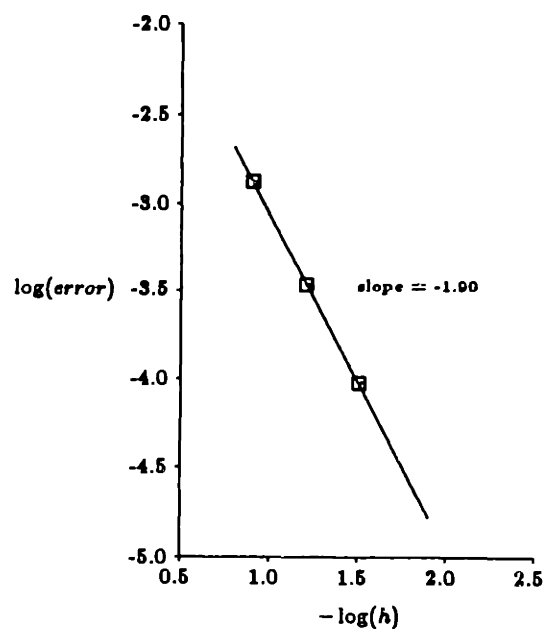


Figure 6.11: Order of accuracy for triangular Jameson scheme with high-accuracy smoothing on a regular mesh

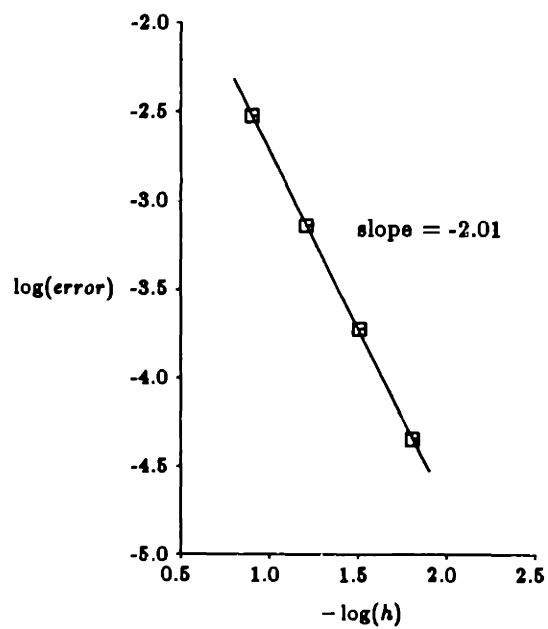


Figure 6.12: Order of accuracy for triangular Jameson scheme with high-accuracy smoothing on an irregular mesh

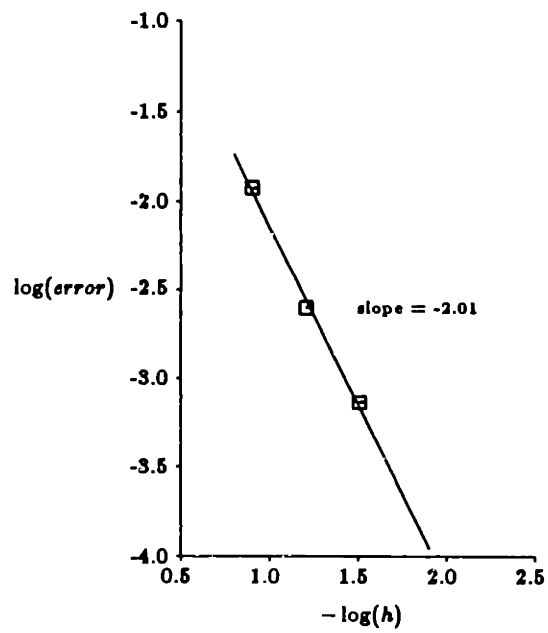


Figure 6.13: Order of accuracy for quadrilateral Jameson scheme with high-accuracy smoothing on a regular mesh

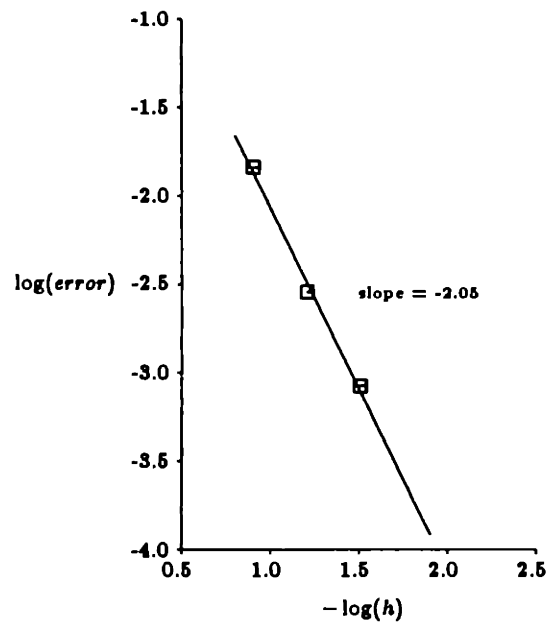


Figure 6.14: Order of accuracy for quadrilateral Jameson scheme with high-accuracy smoothing on an irregular mesh

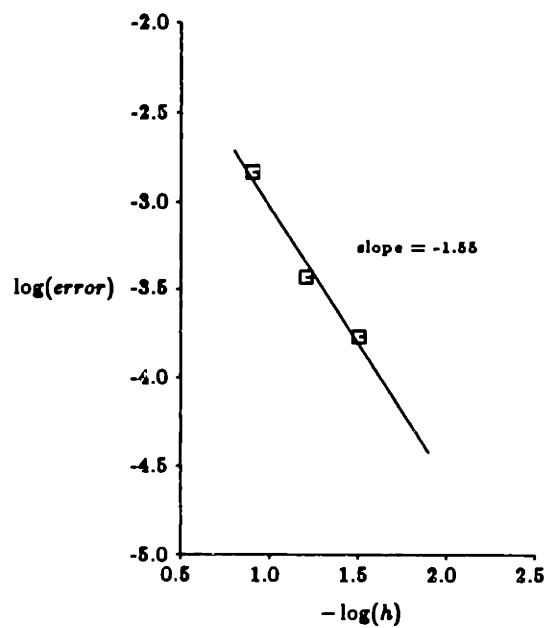


Figure 6.15: Order of accuracy for triangular Jameson scheme with low-accuracy smoothing on a regular mesh, smoothing coefficient $\epsilon = 0.0005$

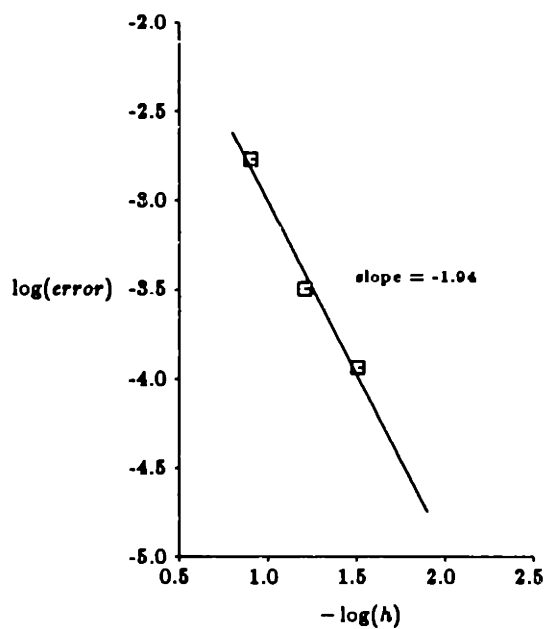


Figure 6.16: Order of accuracy for triangular Jameson scheme with low-accuracy smoothing on a regular mesh, smoothing coefficient $\epsilon = 0.0001$

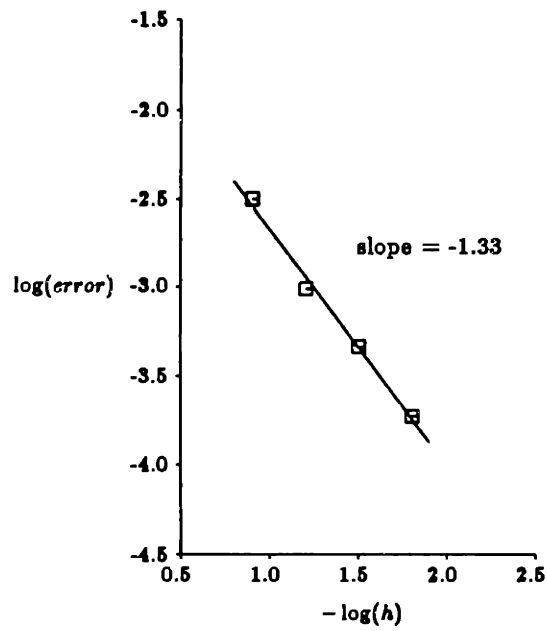


Figure 6.17: Order of accuracy for triangular Jameson scheme with low-accuracy smoothing on an irregular mesh, smoothing coefficient = 0.0005

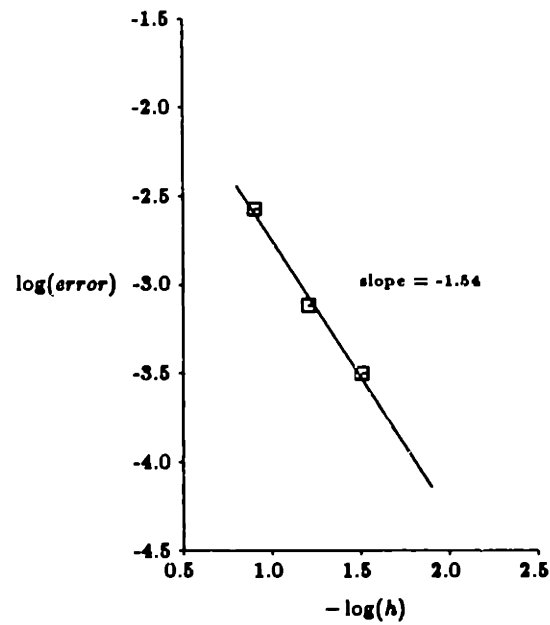


Figure 6.18: Order of accuracy for triangular Jameson scheme with low-accuracy smoothing on an irregular mesh, smoothing coefficient = 0.0001

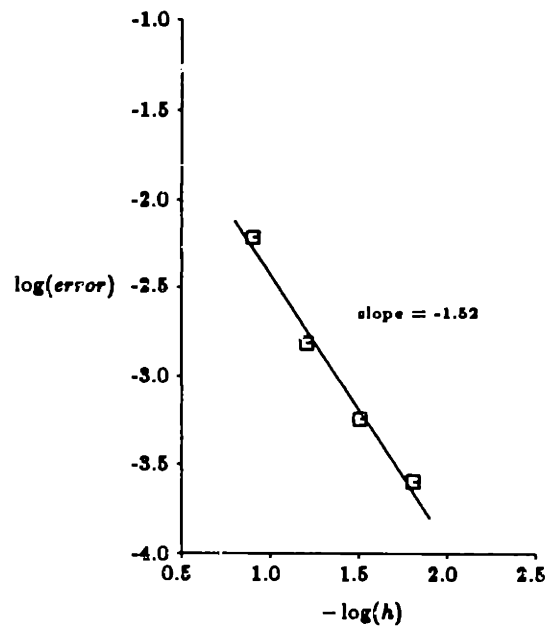


Figure 6.19: Order of accuracy for quadrilateral Jameson scheme with low-accuracy smoothing on a regular mesh, smoothing coefficient = 0.005

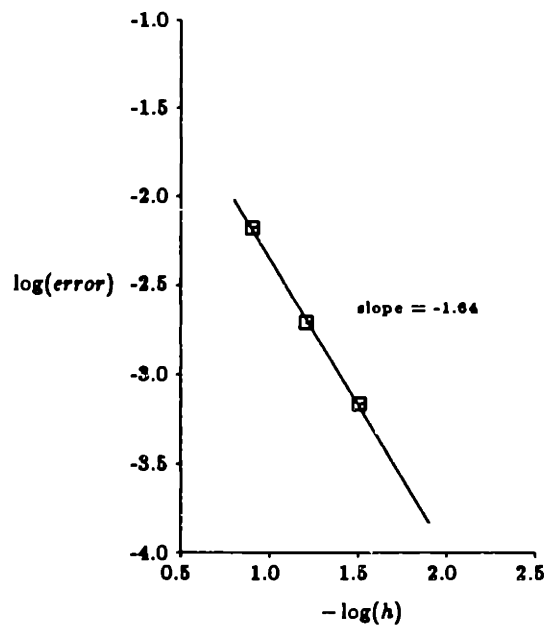


Figure 6.20: Order of accuracy for quadrilateral Jameson scheme with low-accuracy smoothing on an irregular mesh, smoothing coefficient = 0.005

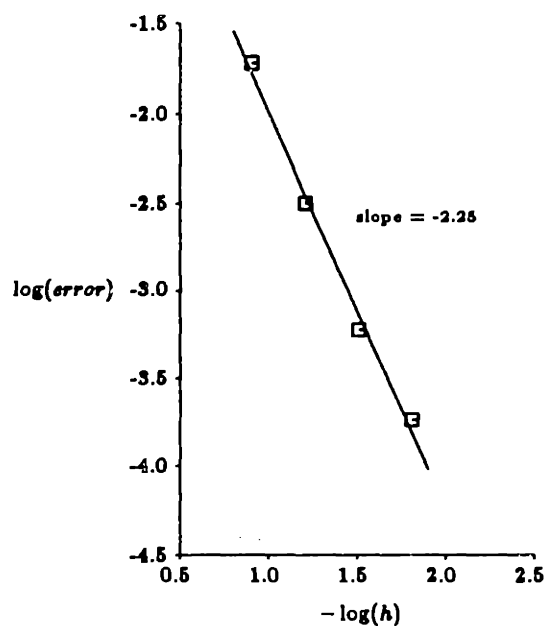


Figure 6.21: Order of accuracy for quadrilateral Jameson scheme with low-accuracy smoothing on a regular mesh, smoothing coefficient = 0.001

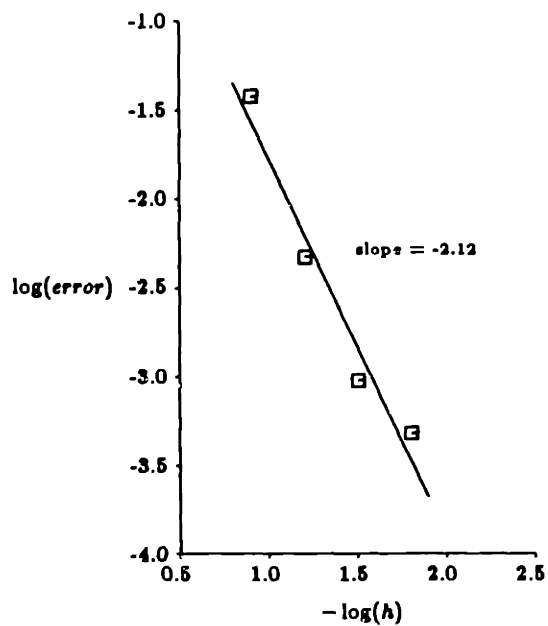


Figure 6.22: Order of accuracy for quadrilateral Jameson scheme with low-accuracy smoothing on an irregular mesh, smoothing coefficient = 0.001

	Numerical Accuracy	Smoothing Coef.
Triangular Ni scheme on regular mesh with high-accuracy smoothing	2.33	0.0025
Triangular Ni scheme on irregular mesh with high-accuracy smoothing	2.17	0.0025
Triangular Jameson scheme on regular mesh with low-accuracy smoothing	1.55	0.0005
	1.94	0.0001
Triangular Jameson scheme on irregular mesh with low-accuracy smoothing	1.33	0.0005
	1.54	0.0001
Triangular Jameson scheme on regular mesh with high-accuracy smoothing	1.90	0.0025
Triangular Jameson scheme on irregular mesh with high-accuracy smoothing	2.01	0.0025
Quadrilateral Ni scheme on regular mesh with high-accuracy smoothing	2.04	0.005
Quadrilateral Ni scheme on irregular mesh with high-accuracy smoothing	2.17	0.005
Quadrilateral Jameson scheme on regular mesh with low-accuracy smoothing	1.52	0.005
	2.25	0.001
Quadrilateral Jameson scheme on irregular mesh with low-accuracy smoothing	1.64	0.005
	2.12	0.001
Quadrilateral Jameson scheme on regular mesh with high-accuracy smoothing	2.01	0.005
Quadrilateral Jameson scheme on irregular mesh with high-accuracy smoothing	2.05	0.005

Figure 6.23: Numerical order of accuracy of numerical schemes

Chapter 7

Adaptation

In order to reduce the computational time required to find the solution to a problem, spatial adaptation can be used. Spatial adaptation is a method which places small cells where the physical characteristic length is small, such as in shocks, and large cells where the physical characteristic length is large. Since the solution is not known before hand, one method for creating an adapted mesh is to start with a course mesh and to divide cells when the physical characteristic length becomes larger than some prescribed value.

7.1 Adaptation Criteria

The features which will be resolved by the adaptation procedure depend on the parameter used to determine when to adapt. Dannenhoffer [3] discusses the merit of different parameters, and concludes that the first difference of density provides a parameter which will resolve both shocks and slip lines in isoenergetic flow, and is inexpensive to calculate. If the division will take place on a cell basis, the first difference of density is found for all the cells, and if the division will take place on a face basis, it is found for all the faces. All cells or faces whose first difference of density is greater than some reference value are divided. The mean and standard deviation of the first difference of density are found over all the cells or faces, and the reference value will be the average plus a factor between 0 and 1 of the standard deviation. It was found that a factor of 0.4 produces good results.

7.2 Quadrilateral Adaptation

For quadrilateral cells the division process typically is made on a cell basis. Several cells are chosen for adaptation and are divided into four smaller cells by adding a node in the center of each chosen course cell and in the center of each of its faces. This process produces an interface between a region of course cells and a region of fine cells. This interface is a result of the regularity in a quadrilateral mesh. An example of such an interface is shown in Figure 7.1. Methods have been developed for dealing with the problem of interfaces [3], but will not be discussed in detail here. Part of the problem is the need to store data on the adaptation history and interface location and to deal with interfaces with special methods in the numerical solver. Results produced by Shapiro [20] for a quadrilateral mesh are shown in Figure 7.2. The computations by Shapiro were done using a cell vertex finite element scheme which is essentially the same as the quadrilateral Jameson scheme discussed here. The three meshes shown have 192, 609 and 1647 cells.

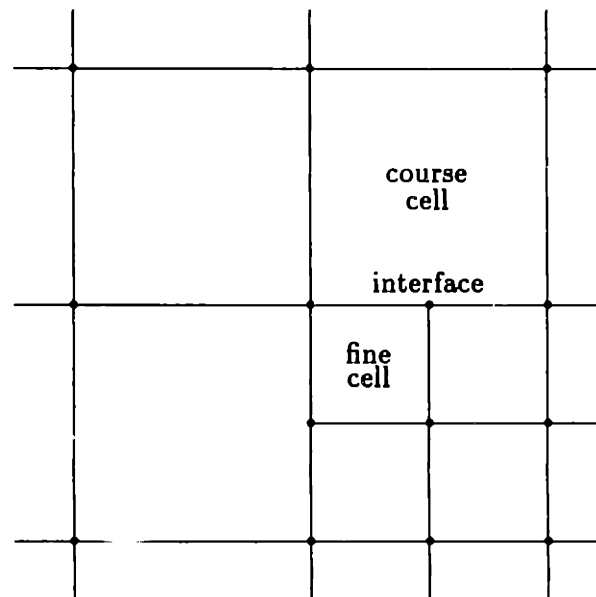


Figure 7.1: Manner of quadrilateral cell division with interfaces

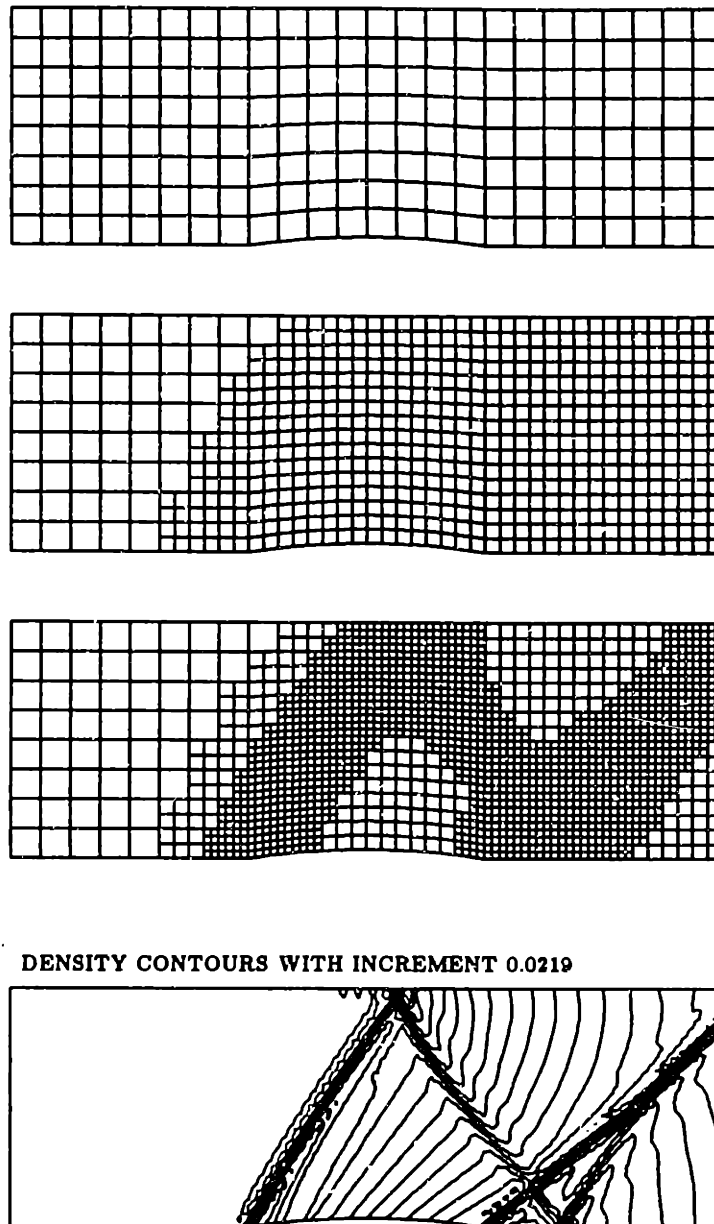


Figure 7.2: Levels of adaptation with a quadrilateral mesh

7.3 Triangular Adaptation

For triangular cells the irregularity in the mesh allows a cell to be divided into smaller cells without creating interfaces. This means that once a cell is divided, no added information must be stored and no special cases are required in the solver. The order of accuracy of the numerical scheme as well as consistency are also preserved. To illustrate this, a simple method for dividing triangular cells was developed.

The division process used here for triangular cells is face based. It is possible for a cell to have one, two or three faces selected for division. To create a smooth mesh, when two sides of a triangle are chosen for division the third is also divided. The manner of cell division is shown in Figure 7.3. Other processes of cell division have been devised which are based on cell division or Delaunay triangulation. These methods would probably create a better adapted mesh, but were not investigated here since the sole objective was to demonstrate the relative ease with which triangular adaptation can be implemented. A simple triangular mesh was used as a base mesh and the meshes after each level of adaptation are shown in Figure 7.4. The triangular Jameson scheme was used to compute the solution. The three meshes shown have 384, 1001 and 2228 cells.

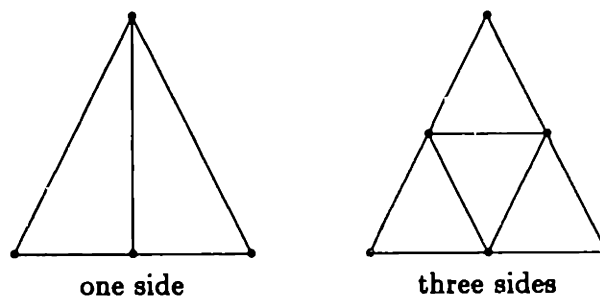
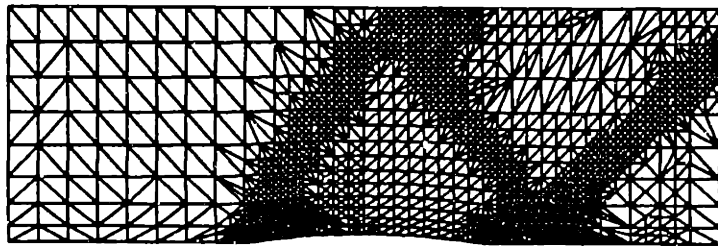
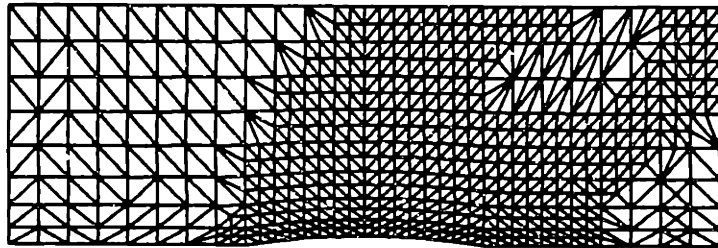
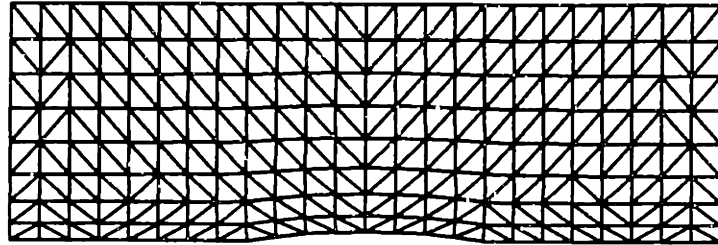


Figure 7.3: Manner of triangular cell division



DENSITY CONTOURS WITH INCREMENT 0.02

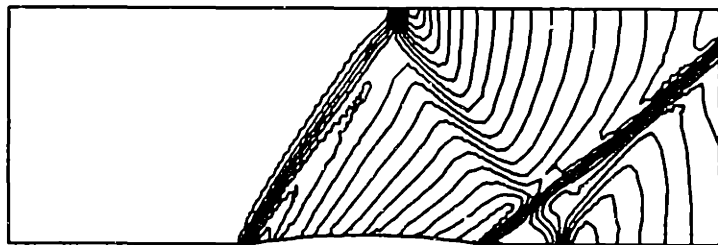


Figure 7.4: Levels of adaptation with a triangular mesh

Chapter 8

Conclusions

This work sheds some light on the controversy in the CFD community over the advantages of quadrilateral and triangular cell schemes. Two different node based finite volume schemes for the Euler equations were studied: a Jameson scheme and a Ni scheme.

A triangular Jameson scheme flow solver was programmed as described by Mavriplis [13] and a similar quadrilateral Jameson scheme was also programmed. A quadrilateral Ni scheme flow solver as described by Ni [15] and Giles [7] was also programmed and a triangular extension of this scheme was developed. Both schemes on triangular and quadrilateral meshes produce similar results. In fact, the quality of the results is more dependent on the numerical smoothing used than the flow solver.

Two different freestream smoothing techniques are examined and implemented with the numerical schemes. Numerical smoothing is required for stability of the Jameson schemes and to reduce undesirable modes in all the schemes. The freestream smoothing consists of a fourth difference of the state vector variables which is formed by taking the second difference of a second difference. Two different methods for finding a second difference are examined, one of which has low-accuracy and the other which has high-accuracy but requires more time to compute. The result is that two conservative fourth differences for numerical smoothing can be created one of which has low-accuracy and another which has high-accuracy, but is more expensive. The low-accuracy smoothing contains no information about the locations of the nodes in the mesh, and as a result produces less smooth solutions on irregular meshes and poorer solutions at solid walls for triangular meshes than the high-accuracy smoothing. The problems with the solid wall boundaries are caused by imposing boundary conditions on the flow field which

cause the low-accuracy numerical smoothing to be convective.

A mathematical study shows that these node based numerical schemes for both quadrilateral and triangular meshes are second order accurate. This is verified here by a numerical accuracy study. The effect of numerical smoothing on the accuracy of the scheme can be large. A numerical smoothing method which is first order accurate can reduce this second order accuracy to first order, since the scheme is only as accurate as the lowest order accurate component. This effect is demonstrated here. When using the low-accuracy smoothing, the accuracy drops below second order due to the lower accuracy of the smoothing. The high-accuracy smoothing allows the schemes to retain their second order accuracy. Therefore both the quadrilateral and triangular schemes are second order accurate if the proper numerical smoothing is used.

Spatial adaptation reduces the computational time required to compute a solution by placing small cells where the physical characteristic length of the flow is small and larger cells where the physical characteristic length of the flow is large. One of the advantages of triangles is the ability to divided cells such that no special interfaces exist. Quadrilateral spatial adaptation, however, requires interfaces between regions of fine and coarse cells. While with triangular adaptation both second order accuracy and conservation can be retained in the adapted region, the quadrilateral adaptation can retain only one of these qualities at the interfaces. In all, spatial adaptation is better suited for triangular meshes, but methods have been developed for quadrilateral meshes which are quite satisfactory.

Triangular meshes by nature of their irregularity allow for mesh generation about complex geometries. The mesh generation process may be quite complicated, but can be made to handle arbitrary several types of geometries and is extendable to three dimensions. The ability to be quite general can be advantageous when compared with the complexities involved in block meshes or overlapping meshes which require user input when generated which is harder to visualize in three dimensions.

A method to define an irregular mesh which is referred to as a pointer system must be used for triangular meshes. Quadrilateral meshes require pointer systems only

when adaptation or other special mesh operations are performed. A pointer system describes the interconnection between elements of a mesh and can easily deal with any irregularities or changes in the mesh structure. In general, pointer systems are beginning to be used for both triangular and quadrilateral meshes, and don't require much more information than the traditional (i, j) method of defining a mesh and allow for longer vector lengths and more concurrency for a parallel processing computer.

This study has focussed on the basic development, performance and accuracy of quadrilateral and triangular mesh schemes. The triangular schemes perform as well as the quadrilateral schemes in both quality of results and accuracy provided an appropriate numerical smoothing technique is used. This result justifies the continued research in the use of triangular mesh schemes. More work must be done in the field of triangular mesh generation and adaptation to bring these computational methods for triangular meshes to a similar level of understanding as on quadrilateral meshes. In three dimensions the use of tetrahedra as opposed to hexahedra significantly increases the ability to produce meshes around complex three dimensional geometries and for this reason more than any other justifies the need to continue research in this area.

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Appendix A

Computer Code

This appendix contains the sources for the flow solvers for the quadrilateral Ni scheme, the quadrilateral Jameson scheme, the triangular Ni scheme and the triangular Jameson scheme. Also included is the code for the mesh generator and the plotting program. The subroutines from the graphics package GRAFIC are not included.

A.1 Triangular Schemes

A.1.1 Common Files

This is file TRI.INC which includes many declarations and common block statements and is included in all subroutines for the triangular schemes.

```

parameter Maxnodes = 5000
parameter Maxfaces = 15141      ! = 3*Maxnodes + 2*sqrt(Maxnodes)
parameter Maxcells = 10000     ! = 2*Maxnodes
parameter Maxedges = 424       ! = 6*sqrt(Maxnodes)

integer Nmax                    !number of nodes
integer Fmax                    !number of faces (including edges)
integer Cmax                    !number of cells (including edge cells)
integer Imax, Omax              !number of inlet and outlet nodes
integer Vmax                    !number of wall faces
integer Emax                    !number of edge faces
integer Pmax                    !number of periodic nodes
integer ENmax                   !number of edge nodes
integer tnode, bnode, inode    !location of diff. types of edge nodes

real pitch                      !blade pitch
real pout                       !outlet pressure for blades
real Sinl                       !tan of inlet flow angle
real x(Maxnodes)                !x values of nodes
real y(Maxnodes)                !y values of nodes
integer eface(Maxedges)         !array of edge faces
integer enode(Maxedges)         !array of edge nodes
real senode(Maxedges)           !sin and cos at edge nodes
real cenode(Maxedges)
integer face(Maxfaces,6)        !array of faces
integer cell(Maxcells,6)        !array of cells
integer innode(Maxedges)        !inlet nodes

```

```

integer outnode(Maxedges)      !outlet nodes
integer pnode(Maxedges,2)      !periodic nodes
integer NCcolor(50)            !number of cells colored each color
integer Ccolormax               !max number of colors used
integer NFcolor(50)            !number of faces colored each color
integer Fcolormax               !max number of colors used
integer NEcolor(10)            !number of edges colored each color
integer Eicolormax, E2colormax !max number of colors used

common /tri/ Imax, Omax, Wmax, Pmax
common /tri/ Nmax, Fmax, Cmax, Emax
common /tri/ ENmax, tnode, bnode, inode
common /tri/ pitch, pout, Sini
common /tri/ x, y, eface, enode, senode, cenode, face, cell
common /tri/ innode, outnode, pnode
common /color/ NCcolor, Ccolormax, NFcolor, Fcolormax
common /color/ NEcolor, Eicolormax, E2colormax

real Minl                      !inlet Mach number
integer Maxiter                 !max number of iterations
real CFL                       !CFL number
real epsicoef, eps2            !smoothing coef
real sigE, sigV                !smoothing coef
real vol(Maxcells)             !for vorticity smoothing
real durms                     !rms difference in state vector
real U(4,0:Maxnodes)           !state vector
real dU(4,Maxnodes)            !change in state vector at nodes
real Uc(4,Maxcells)            !state vector at cells
real dUc(4,0:Maxcells)         !change in state vector at cells
real F(4,Maxnodes)             !flux vectors at nodes
real G(4,Maxnodes)
real areaN(0:Maxnodes)         !control area around each node
real areaC(0:Maxcells)         !area of each cell
real deltN(0:Maxnodes)         !time step at each node (Jameson)
                                ! or cell (Ni)
real deltC(0:Maxcells)         !time step at each cell
real dis(4,Maxnodes)           !dissipation at each node
real flux(4,0:Maxnodes)        !flux at each node

common /flo/ Minl, durms, Maxiter, CFL
common /flo/ epsicoef, eps2, sigE, sigV, vol
common /flo/ U, dU, Uc, dUc
common /flo/ F, G
common /flo/ deltC, deltN, areaC, areaN, dis, flux

parameter gam = 1.4
parameter gam1 = 0.4

```

This file contains the subroutines for input and output and is linked with all the triangular schemes.

```

c*****
c*
c*   write or read grid data from file
c*
c*****

subroutine gridio(process)
implicit none

include 'TRI.INC'
integer process                !read or write?
integer i, NN, FF, CC, EF, EN, col !pointers
integer IN, ON, WN, PN

if (process.eq.1) then

```

```

c** read in data from file
    open(unit=30, status='unknown', form='unformatted')

    read(30) Nmax, (x(NN),NN=1,Nmax), (y(NN),NN=1,Nmax)
    read(30) Fmax, ((face(FF,i),i=1,6),FF=1,Fmax)
    read(30) Wmax, Emax, (eface(EF),EF=1,Emax)
    read(30) tnode, bnode, inode, ENmax, (enode(EN),EN=1,ENmax)
    read(30) (senode(EN),EN=1,ENmax), (cenode(EN),EN=1,ENmax)
    read(30) lmax, (innode(IN),IN=1,lmax)
    read(30) Omax, (outnode(ON),ON=1,Omax)
    read(30) Pmax, ((pnode(PN,i),i=1,2),PN=1,Pmax)
    read(30) Cmax, ((cell(CC,i),i=1,6),CC=1,Cmax)
    read(30) Ccolormax, (NCcolor(col),col=1,Ccolormax)
    read(30) Fcolormax, (NFcolor(col),col=1,Fcolormax)
    read(30) Ecolormax, E2colormax
    read(30) (NEcolor(col),col=1,E2colormax)
    read(30) pitch

    close(unit=30)

    else if (process.eq.0) then

c* write data to file
    open(unit=30, status='unknown', form='unformatted')

    write(30) Nmax, (x(NN),NN=1,Nmax), (y(NN),NN=1,Nmax)
    write(30) Fmax, ((face(FF,i),i=1,6),FF=1,Fmax)
    write(30) Wmax, Emax, (eface(EF),EF=1,Emax)
    write(30) tnode, bnode, inode, ENmax, (enode(EN),EN=1,ENmax)
    write(30) (senode(EN),EN=1,ENmax), (cenode(EN),EN=1,ENmax)
    write(30) lmax, (innode(IN),IN=1,lmax)
    write(30) Omax, (outnode(ON),ON=1,Omax)
    write(30) Pmax, ((pnode(PN,i),i=1,2),PN=1,Pmax)
    write(30) Cmax, ((cell(CC,i),i=1,6),CC=1,Cmax)
    write(30) Ccolormax, (NCcolor(col),col=1,Ccolormax)
    write(30) Fcolormax, (NFcolor(col),col=1,Fcolormax)
    write(30) Ecolormax, E2colormax
    write(30) (NEcolor(col),col=1,E2colormax)
    write(30) pitch

    close(unit=30)

endif

return
end

```

```

c*****
c*
c* write or read flow data from file
c*
c*****

```

```

    subroutine flowio(process)
    implicit none

    include 'TRI.INC'
    integer process          !read or write?
    integer i, NN            !pointers

    if (process.eq.1) then

c** read in data from file
        open(unit=25, status='unknown', form='unformatted')

        read(25) Minl
        read(25) Nmax, ((U(i,NN),i=1,4),NN=1,Nmax)

```

```

        close(unit=25)

    else if (process.eq.0) then
c* write data to file
        open(unit=25, status='unknown', form='unformatted')

        write(25) Minl
        write(25) Hmax, ((U(1,NN),i=1,4),NN=1,Nmax)

        close(unit=25)

    endif

    return
end

c*****
c*
c* read input data from file
c*
c*****

subroutine input
implicit none

include 'TRI.INC'
real pi !the one and only

pi = 3.14159

open(unit=20, status='old')

read(20,*) Maxiter, CFL
read(20,*) Minl, Sinl, pout
read(20,*) sigE, sigV, epsicoef, eps2
close (unit=20)

pout = pout/gm
Sinl = tan(pi+Sinl/180.0)

return
end

```

A.1.2 Mesh Generator

This is file GRID.INC which includes many declarations and common block statements for the mesh generator.

```

parameter Maxdim = 200

real xx(0:Maxdim,0:Maxdim) !x coordinate of grid points
real yy(0:Maxdim,0:Maxdim) !y coordinate of grid points
integer Nx !number cells on x axis
integer Ny ! " " " y "
integer ILE, ITE !leading and trailing edge of blade

```

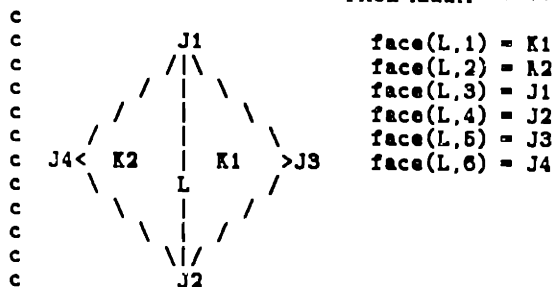
```
common /grid/ xx, yy, Nx, Ny
common /grid/ ILE, ITE
```

```
parameter IBX=251
```

```
real GSINL, GSOUT, CHINL, CHOUT
integer II, JJ
real XB(IBX), XPB(IBX), YB(IBX), YPB(IBX), SB(IBX)
integer IIB, IBLE
real SBLE, SBLOLD, SS, SP
real XMIN, XMAX, YMIN, XLE, YLE, XPOS(Maxdim), YPOS(Maxdim)
integer NINL, NOUT, NBLD
```

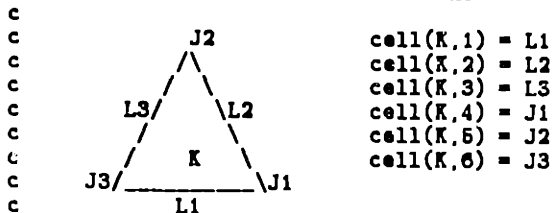
```
common /c01/ GSINL, GSOUT, CHINL, CHOUT, II, JJ,
*      XB, XPB, YB, YPB, SB,
*      IIB, IBLE, SBLE, SBLOLD, SS, SP,
*      XMIN, XMAX, YMIN, XLE, YLE, XPOS, YPOS,
*      NINL, NOUT, NBLD
```

```
C***** FACE ARRAY *****
```



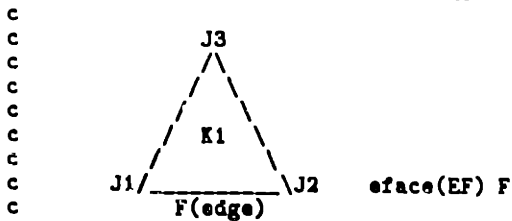
```
face(L,1) = K1
face(L,2) = K2
face(L,3) = J1
face(L,4) = J2
face(L,5) = J3
face(L,6) = J4
```

```
C***** CELL ARRAY *****
```



```
cell(K,1) = L1
cell(K,2) = L2
cell(K,3) = L3
cell(K,4) = J1
cell(K,5) = J2
cell(K,6) = J3
```

```
C***** EDGE ARRAY *****
```



```
eface(EF) F
```

```
C*****
```

```
program gridgen
implicit none
```

```
include 'TRI.INC'
include 'GRID.INC'
integer gtype !type of grid
integer EN, EF, FF, NN, CC, col !pointers
integer ON, IN, PN
integer i, j !pointers
```



```

        integer JJJ

c** determine geometry
        type*, 'Type of grid '
        type*, ' 1) N1 bump '
        type*, ' 2) blade'
        type 100
100    format(0, ' selection = ')
        accept 110, gtype
110    format(I)

        write(6,*) ' '
        write(6,*) 'Generating grid ...'
        write(6,*) ' '

        if (gtype.eq.1) then
c** create rectangular mesh
            call rectangle
            pitch = 0.

            else if (gtype.eq.2) then
c** Read in, normalize and spline blade data
                call readin

c** Initialize grid
                call grinit

c** Fix up grid
                call ellip(Maxdim,Maxdim,II,JJ,JJJ,XX,YY,YPOS,XPOS)
                call improv

c** change pointers
                do i = 1, ii
                    do j = 1, jj
                        xx(i-1,j-1) = xx(i,j)
                        yy(i-1,j-1) = yy(i,j)
                    enddo
                enddo
                Nx = ii-1
                Ny = jj-1
            endif

c** change rectangular mesh to triangular mesh
            call pointers
            call bpointers

c** rearrange node numbers on edges
            call edgenumber

c** color cells and faces
            call cellcolor
            call facecolor
            call edgecolor

c** write out data to file
            call gridio(0)

            stop
            end

subroutine rectangle
implicit none

include 'GRID.INC'

real*4 yymax, xxmax, xxmin      !coordinates of grid boundaries

```

```

real*4 delta_x           !grid spacing in x direction
real*4 delta_y           !grid spacing in y direction
integer n, i, j, kk, m  !counters
real*4 num
real*4 delta_xbot
real*4 omega             !relaxation constant for
                           !interior point SLOR
real*4 pi                !the one and only
real*4 tau               !height of bump
real*4 rr               !radius of bump
real*4 yc               !rr-tau
real*4 ang, delta_ang   !angle of buap, angle between nodes
real*4 angplus
real*4 delx, dely       !help define initial conditions
real*4 alpha(Maxdim), beta(Maxdim) !coef. in modified equation
real*4 gamma(Maxdim)
real*4 AA(Maxdim), BB(Maxdim) !values for system of equations
real*4 CC(Maxdim), DD(Maxdim)
real*4 EPS              !allowable error
integer Niter           !number of iterations so far
real*4 error           !largest error for an iteration
real*4 x_xi(Maxdim), x_eta(Maxdim) !derivatives on boundary
real*4 x_xi_xi(Maxdim), x_eta_eta(Maxdim)
real*4 x_eta_xi(Maxdim)
real*4 y_xi(Maxdim), y_eta(Maxdim)
real*4 y_xi_xi(Maxdim), y_eta_eta(Maxdim)
real*4 y_eta_xi(Maxdim)
real*4 theta, AR        !angle and aspect ratio of edge cells
real*4 R1, R2           !part of source term
real*4 Q1(Maxdim), P1(Maxdim) !part of source term, function of xi
real*4 a1(Maxdim), b1(Maxdim) !alpha, beta, gamma on lower boundary
real*4 c1(Maxdim)
real*4 omega_P, omega_Q !relaxation conts. for source terms
real*4 Jacobi1(Maxdim)  !Jacobian on the lower boundary
real*4 Jacobi2(Maxdim)  !Jacobian squared in region
real*4 xxi, yxi, xeta, yeta !derivatives in region
real*4 a, b             !exponents in source terms

pi = 3.14159

c** number of nodes on x and y axes
xxmax = 2.
xxmin = -1.
yymin = 1.

call Irequest('      Nx',Nx)
call Irequest('      Ny',Ny)
call Rrequest('      tau',tau)
c  call Rrequest('      omega',omega)
c  call Rrequest('      omega_P',omega_P)
c  call Rrequest('      omega_Q',omega_Q)
c  call Rrequest('      a',a)
c  call Rrequest('      b',b)
c  call Rrequest('      AR',AR)
c  call Rrequest('      EPS',EPS)

omega = 1.
a = 0.8
b = 0.8
AR = 0.5
EPS = 0.0005

if (tau.eq.0.) then
  omega_P = 0.
  omega_Q = 0.
else if (tau.lt.0.) then
  AR = 1.
  omega_P = 0.02
  omega_Q = 0.02
else
  omega_P = 0.02
  omega_Q = 0.02

```

```

endif

theta = .5*pi
delta_x = (xxmax-xxmin)/real(Nx)

c** set initial and boundary conditions for x and y
if (tau.gt.0.) then
  yc = 0.5*(-tau**2 + 0.25)/tau
  rr = sqrt(0.25 + yc**2)
  ang = asin(.5/rr)
  if (mod(Nx,3).ne.0) then
    num = real(Nx - mod(Nx,3))
    delta_xbot = (xxmax-xxmin)/num
    angplus = 2.*ang/(num/3. + real(mod(Nx,3)))
  else
    angplus = 2.*ang*delta_x
    delta_xbot = delta_x
  endif
  delta_ang = 0.
  do i = 0, Nx
    yy(1,Ny) = yymax
    xx(1,Ny) = xxmin + i*delta_x
    if (xx(1,Ny).le.0.) then
      xx(1,0) = xxmin + i*delta_xbot
      yy(1,0) = 0.
    else if (xx(1,Ny).gt.0. .and. xx(1,Ny).lt.1.) then
      delta_ang = delta_ang + angplus
      xx(1,0) = 0.5 - rr*sin(ang-delta_ang)
      yy(1,0) = rr*cos(ang-delta_ang) - yc
    else if (xx(1,Ny).ge.1.) then
      xx(1,0) = xxmin + (i-mod(Nx,3))*delta_xbot
      yy(1,0) = 0.
    endif
    delx = (xx(1,Ny) - xx(1,0))/real(Ny)
    dely = (yy(1,Ny) - yy(1,0))/real(Ny)
    do j = 1, Ny-1
      xx(1,j) = delx*j + xx(1,0)
      yy(1,j) = dely*j + yy(1,0)
    enddo
  enddo
else if (tau.lt.0.) then
  delta_xbot = delta_x
  do i = 0, Nx
    yy(1,Ny) = yymax
    xx(1,Ny) = xxmin + i*delta_x
    if (xx(1,Ny).le.0.) then
      xx(1,0) = xxmin + i*delta_xbot
      yy(1,0) = 0.
    else if (xx(1,Ny).gt.0. .and. xx(1,Ny).lt.1.) then
      xx(1,0) = xxmin + i*delta_xbot
      yy(1,0) = -tau*(sin(pi*xx(1,0)))**2
    else if (xx(1,Ny).ge.1.) then
      xx(1,0) = xxmin + i*delta_xbot
      yy(1,0) = 0.
    endif
    delx = (xx(1,Ny) - xx(1,0))/real(Ny)
    dely = (yy(1,Ny) - yy(1,0))/real(Ny)
    do j = 1, Ny-1
      xx(1,j) = delx*j + xx(1,0)
      yy(1,j) = dely*j + yy(1,0)
    enddo
  enddo
else if (tau.eq.0) then
  delta_y = yymax/real(Ny)
  do i = 0, Nx
    do j = 0, Ny
      xx(1,j) = i*delta_x + xxmin
      yy(1,j) = j*delta_y
    enddo
  enddo
endif

```

```

c* Solve for source terms
do n = 1, Nx-1
    P1(n) = 0.
    Q1(n) = 0.
    x_xi(n) = .5*(xx(n+1,0) - xx(n-1,0))
    y_xi(n) = .5*(yy(n+1,0) - yy(n-1,0))

    x_xi_xi(n) = (xx(n+1,0) - 2.*xx(n,0) + xx(n-1,0))
    y_xi_xi(n) = (yy(n+1,0) - 2.*yy(n,0) + yy(n-1,0))

    if (xx(n,0).eq.0. .or. xx(n,0).eq.1.) then
        x_eta(n) = max(5.*tau,1.)*AR*(-x_xi(n)*cos(theta)
        & - y_xi(n)*sin(theta))
        y_eta(n) = max(5.*tau,1.)*AR*(-y_xi(n)*cos(theta)
        & + x_xi(n)*sin(theta))
    else
        x_eta(n) = AR*(-x_xi(n)*cos(theta) - y_xi(n)*sin(theta))
        y_eta(n) = AR*(-y_xi(n)*cos(theta) + x_xi(n)*sin(theta))
    endif

    x_eta_xi(n) = 0.5*(x_eta(n+1)-x_eta(n-1))
    y_eta_xi(n) = 0.5*(y_eta(n+1)-y_eta(n-1))

    Jacobi(n) = x_xi(n)*y_eta(n) - x_eta(n)*y_xi(n)

    a1(n) = x_eta(n)**2 + y_eta(n)**2
    b1(n) = x_xi(n)*x_eta(n) + y_xi(n)*y_eta(n)
    c1(n) = x_xi(n)**2 + y_xi(n)**2
enddo

c* SOR by lines
Niter = 0
error = 9999.
do while (error.gt.EPS)
    error = 0.
    Niter = Niter + 1

c** solve for source terms
    if (tau.ne.0.) then
        do n = 1, Nx-1
            x_eta_eta(n) = 0.5*(-7.*xx(n,0) + 8.*xx(n,1) - xx(n,2))
            & - 3.*x_eta(n)
            y_eta_eta(n) = 0.5*(-7.*yy(n,0) + 8.*yy(n,1) - yy(n,2))
            & - 3.*y_eta(n)
            R1 = (-a1(n)*x_xi_xi(n) + 2.*b1(n)*x_eta_xi(n)
            & - c1(n)*x_eta_eta(n))/Jacobi(n)**2
            R2 = (-a1(n)*y_xi_xi(n) + 2.*b1(n)*y_eta_xi(n)
            & - c1(n)*y_eta_eta(n))/Jacobi(n)**2
            P1(n) = P1(n) + omega_P*((y_eta(n)*R1 - x_eta(n)*R2)
            & /Jacobi(n) - P1(n))
            Q1(n) = Q1(n) + omega_Q*((-y_xi(n)*R1 + x_xi(n)*R2)
            & /Jacobi(n) - Q1(n))
        enddo
    endif

c* solve for each line
    do kk = 1, Ny-1

c* evaluate alpha, beta, and gamma
        do i = 1, Nx-1
            alpha(i) = .25*(xx(i,kk+1)-xx(i,kk-1))**2
            & + .25*(yy(i,kk+1)-yy(i,kk-1))**2
            beta(i) = .25*(xx(i+1,kk)-xx(i-1,kk))*(xx(i,kk+1)
            & -xx(i,kk-1)) + .25*(yy(i+1,kk)-yy(i-1,kk))
            & *(yy(i,kk+1)-yy(i,kk-1))
            gamma(i) = .25*(xx(i+1,kk)-xx(i-1,kk))**2
            & + .25*(yy(i+1,kk)-yy(i-1,kk))**2
            Jacobi2(i) = (.25*(xx(i+1,kk)-xx(i-1,kk))*
            & (yy(i,kk+1)-yy(i,kk-1))
            & - .25*(yy(i+1,kk)-yy(i-1,kk))*
            & (xx(i,kk+1)-xx(i,kk-1)))**2
        enddo
    enddo

```

```

c* Solve for x
c* set up matrix for tridiagonal system of equations
do i = 1, Nx-1
    AA(i) = omega*alpha(i)
    DD(i) = -2.*(gamma(i) + alpha(i))
    BB(i) = omega*alpha(i)
enddo

c* set up vector of constants for tridiagonal system of equations
do i = 1, Nx-1
    xxi = xx(i+1,kk) - xx(i-1,kk)
    xeta = xx(i,kk+1) - xx(i,kk-1)

    CC(i) = -omega*gamma(i)*(xx(i,kk+1) + xx(i,kk-1))
    & + omega*0.5*beta(i)*(xx(i+1,kk+1)
    & - xx(i+1,kk-1) - xx(i-1,kk+1) + xx(i-1,kk-1))
    & + 2.*(omega-1.)*(alpha(i) + gamma(i))*xx(i,kk)
    & - Jacobi2(i)*(Pi(i)*xxi*exp(-a*kk) +
    & Q1(i)*xeta*exp(-b*kk))
enddo
CC(1) = CC(1) - BB(1)*xx(0,kk)
CC(Nx-1) = CC(Nx-1) - AA(Nx-1)*xx(Nx,kk)

c* solve tridiagonal system of equations
call tridiag(1, Nx-1, BB, DD, AA, CC)
do m = 1, Nx-1
    if (abs(CC(m)-xx(m,kk)).gt.error) then
        error = abs(CC(m) - xx(m,kk))
    endif
    xx(m,kk) = CC(m)
enddo

c* Solve for y
c* set up matrix for tridiagonal system of equations
do i = 1, Nx-1
    AA(i) = omega*alpha(i)
    DD(i) = -2.*(gamma(i) + alpha(i))
    BB(i) = omega*alpha(i)
enddo

c* set up vector of constants for tridiagonal system of equations
do i = 1, Nx-1
    yxi = yy(i+1,kk) - yy(i-1,kk)
    yeta = yy(i,kk+1) - yy(i,kk-1)

    CC(i) = -omega*gamma(i)*(yy(i,kk+1) + yy(i,kk-1))
    & + 0.5*omega*beta(i)*(yy(i+1,kk+1)
    & - yy(i+1,kk-1) - yy(i-1,kk+1) + yy(i-1,kk-1))
    & + 2.*(omega-1.)*(alpha(i)+gamma(i))*yy(i,kk)
    & - Jacobi2(i)*(Pi(i)*yxi*exp(-a*kk) +
    & Q1(i)*yeta*exp(-b*kk))
enddo
CC(1) = CC(1) - BB(1)*yy(0,kk)
CC(Nx-1) = CC(Nx-1) - AA(Nx-1)*yy(Nx,kk)

c* solve tridiagonal system of equations
call tridiag(1, Nx-1, BB, DD, AA, CC)
do m = 1, Nx-1
    if (abs(CC(m)-yy(m,kk)).gt.error) then
        error = abs(CC(m) - yy(m,kk))
    endif
    yy(m,kk) = CC(m)
enddo

enddo

c* set x=xxmax and x=xxmin boundary conditions to next interior point
do kk = 1, Ny-1
    yy(0,kk) = yy(1,kk)
    yy(Nx,kk) = yy(Nx-1,kk)
enddo

```

```

        type*, 'iteration number = ',Niter,' error = ',error
    enddo

    return
end

c*****
c*
c*      Subroutine to solve a tridiagonal system of equations.
c*      Taken from "Computational Fluid Mechanics and Heat Transfer"
c*      by Anderson, Tannehill and Pletcher.
c*
c*****

SUBROUTINE TRIDIAG(IL,IU,BB,DD,AA,RR)
    IMPLICIT NONE

    include 'GRID.INC'

    INTEGER IL                      !SUBSCRIPT OF FIRST EQUATION
    INTEGER IU                      !SUBSCRIPT OF LAST EQUATION
    REAL*4 BB(Naxdim)              !COEFFICIENT BEHIND DIAGONAL
    REAL*4 DD(Naxdim)              !COEFFICIENT ON DIAGONAL
    REAL*4 AA(Naxdim)              !COEFFICIENT AHEAD OF DIAGONAL
    REAL*4 RR(Naxdim)              !ELEMENT OF CONSTANT VECTOR
    INTEGER LP
    INTEGER I, J                    !POINTERS
    REAL*4 R

C**      ESTABLISH UPPER TRIANGULAR MATRIX

    LP = IL + 1
    DO I = LP, IU
        R = BB(I)/DD(I-1)
        DD(I) = DD(I)-R*AA(I-1)
        RR(I) = RR(I)-R*RR(I-1)
    ENDDO

C**      BACK SUBSTITUTION

    RR(IU) = RR(IU)/DD(IU)
    DO I = LP, IU
        J = IU - I + 1
        RR(J) = (RR(J)-AA(J)*RR(J+1))/DD(J)
    ENDDO

C**      SOLUTION STORED IN RR

    RETURN
END

c*****
c*
c*      this subroutine requests the user to input the value of a
c*      real variable
c*
c*****

subroutine Rrequest (name,var)
    character*10 name
    real*4 var

    write(6,1) name
1   format(9,' ',A,' = ')
    accept 11, var

```

```

11      format(F)

      return
      end

C*****
C*
C*      this subroutine requests the user to input the value of a      *
C*      integer variable      *
C*      *
C*****

      subroutine Irequest (name,var)
      character*10 name
      integer var

      write(6,1) name
1      format(1,' ',A,' = ')
      accept 11, var
11     format(I)

      return
      end

SUBROUTINE READIN

C---- Read in, normalize and spline blade data

      INCLUDE 'TRI.INC'
      INCLUDE 'GRID.INC'

      CHARACTER*32 NAMEXT
      CHARACTER*80 NAME

C---- Read in blade data
      OPEN(UNIT=3,STATUS='OLD')
1000  FORMAT(A32)
      READ(3,1000) NAME
      READ(3,*) GSINL, GSOUT, CHINL, CHOUT, PITCH

      WRITE(6,1001) NAME
1001  FORMAT(/,' Blade name: ',A60)

      READ(3,*) XB(1), YB(1)
      XMIN = XB(1)
      XMAX = XB(1)
      YMIN = YB(1)
      DO 1 IB = 2, 12345
         READ(3,*,END=11) XB(IB),YB(IB)
         XMAX = AMAX1(XMAX,XB(IB))
         IF(XMIN.GT.XB(IB)) THEN
            XMIN = XB(IB)
            YMIN = YB(IB)
         ENDIF
1      CONTINUE
11     IIB = IB - 1
      CLOSE(UNIT=3)

      IF(IIB.GT.IBX) STOP 'Array overflow: IBX too small'

C---- Normalize blade and calculate surface arc length array
      PITCH = PITCH/(XMAX-XMIN)
      DO 2 IB = 1, IIB
         XB(IB) = (XB(IB)-XMIN) / (XMAX-XMIN)
         YB(IB) = (YB(IB)-YMIN) / (XMAX-XMIN)
2      CONTINUE

```

```

C---- close t.e. if open
IF( XB(1).NE.XB(IIB) .OR. YB(1).NE.YB(IIB) ) THEN
  ASOUT = ATAN( (YB(2) -YB(1)) / (XB(2) -XB(1)) )
  APOUT = ATAN( (YB(IIB)-YB(IIB-1)) / (XB(IIB)-XB(IIB-1)) )
  DSTE = SQRT( (XB(1)-XB(IIB))**2 + (YB(1)-YB(IIB))**2 )
  AOUT = 0.5*(APOUT+ASOUT)
  XOUT = 0.5*(XB(1)+XB(IIB)) + 3.0*DSTE*COS(AOUT)
  YOUT = 0.5*(YB(1)+YB(IIB)) + 3.0*DSTE*SIN(AOUT)
  XS1 = (XE(1)-XOUT)*COS(AOUT) + (YB(1)-YOUT)*SIN(AOUT)
  YS1 = -(XB(1)-XOUT)*SIN(AOUT) + (YB(1)-YOUT)*COS(AOUT)
  YPS1 = TAN(ASOUT-AOUT) * XS1
  XP1 = (XB(IIB)-XOUT)*COS(AOUT) + (YB(IIB)-YOUT)*SIN(AOUT)
  YP1 = -(XB(IIB)-XOUT)*SIN(AOUT) + (YB(IIB)-YOUT)*COS(AOUT)
  YPP1 = TAN(APOUT-AOUT) * XP1

  WRITE(6,1002)
1002 FORMAT(/,' Input flap deflection angle (degrees): ',%)
  READ(5,*) AFLAP
  YPFLAP = TAN(3.14159*AFLAP/180.0)

  WRITE(NAMEXT,1003) AFLAP
1003 FORMAT(' (flap deflection angle =',F4.1,')')
  LENSTART = INDEX(NAME,' ')
  NAME(LENSTART:LENSTART+31) = NAMEXT

  IIB = IIB+20
  IF(IIB.GT.IBX) STOP 'Array overflow: IBX too small'

  DO 3 IB = IIB-20, 1, -1
    XB(IB+10) = XB(IB)
    YB(IB+10) = YB(IB)
3    CONTINUE

  DO 4 IB = 1, 10
    ETA = 0.1*(IB-1)
    XXS = XS1*ETA
    YYS = YS1*ETA*ETA*(3.0-2.0*ETA) + YPS1*ETA*ETA*(ETA-1.0)
    * - YPFLAP*XS1*0.5*(ETA-1.0)**2
    XB(IB) = XOUT + XXS*COS(AOUT) - YYS*SIN(AOUT)
    YB(IB) = YOUT + XXS*SIN(AOUT) + YYS*COS(AOUT)
4    CONTINUE

  DO 5 IB = IIB-9, IIB
    ETA = 0.1*(IIB-IB)
    XXP = XP1*ETA
    YYP = YP1*ETA*ETA*(3.0-2.0*ETA) + YPP1*ETA*ETA*(ETA-1.0)
    * - YPFLAP*XP1*0.5*(ETA-1.0)**2
    XB(IB) = XOUT + XXP*COS(AOUT) - YYP*SIN(AOUT)
    YB(IB) = YOUT + XXP*SIN(AOUT) + YYP*COS(AOUT)
5    CONTINUE

  ENDIF

C---- Spline blade surface(s) and find leading edge position
SB(1) = 0.
DO 6 IB = 2, IIB
  ALF = FLOAT( MIN(IB-1,IIB-IB) ) / FLOAT(IIB/2)
  SB(IB) = SB(IB-1) +
  * SQRT( (XB(IB)-XB(IB-1))**2 + (ALF*(YB(IB)-YB(IB-1)))**2 )
6  CONTINUE

  CALL SPLINE(XB,XPB,SB,IIB)
  CALL SPLINE(YB,YPB,SB,IIB)

  DO 7 IB=2, IIB
    DP1 = XPB(IB-1) + GSINL*YPB(IB-1)
    DP2 = XPB(IB) + GSINL*YPB(IB)
    IF(DP1.LT.0.0 .AND. DP2.GE.0.0) GO TO 71
7  CONTINUE

  STOP 'Leading edge not found'
71 DSB = SB(IB) - SB(IB-1)

```



```

SBLE = SB(IB-1) + DSB*DP1/(DP1-DP2)
XLE = SEVAL(SBLE,XB,XPB,SB,IIB)
YLE = SEVAL(SBLE,YB,YPB,SB,IIB)

```

```

RETURN
END

```

SUBROUTINE GRINIT

C---- Fix grid points on boundary of domain, and initialize interior

```

INCLUDE 'TRI.INC'
INCLUDE 'GRID.INC'

```

C---- Input and check grid size

```

WRITE(6,1000)
1000 FORMAT(/,' Input II, JJ: ',%)
READ(6,*) II,JJ
IF(II.GT.Maxdim) STOP 'Array overflow: Maxdim too small'
IF(JJ.GT.Maxdim) STOP 'Array overflow: Maxdim too small'

```

C---- Set various parameters

```

SLEN = CHINL + 0.6*SB(IIB) + CHOUT
NINL = INT( FLOAT(II)*CHINL/SLEN )
NOUT = INT( FLOAT(II)*CHOUT/SLEN )
NBLD = II - NOUT - NINL + 2
ILE = NINL
ITE = II - NOUT + 1

```

C---- Set inlet stagnation streamline

```

DO 1 K=1, NINL
  XX(K,1) = XLE + CHINL * FLOAT(K-NINL) / FLOAT(NINL-1)
  YY(K,1) = YLE + (XX(K,1)-XLE) * GSINL
  XX(K,JJ) = XX(K,1)
  YY(K,JJ) = YY(K,1) + PITCH
1 CONTINUE

```

C---- Set outlet stagnation streamline

```

XTE = XB(1)
YTE = YB(1)

DO 2 K= 1, NOUT
  I = II-NOUT+K
  XX(I,1) = XTE + CHOUT * FLOAT(K-1) / FLOAT(NOUT-1)
  YY(I,1) = YTE + (XX(I,1)-XTE) * GSOUT
  XX(I,JJ) = XX(I,1)
  YY(I,JJ) = YY(I,1) + PITCH
2 CONTINUE

```

C---- Set points on blade suction surface

```

DO 3 K=1, NBLD
  I = NINL + K - 1
  S = SBLE - SBLE*FLOAT(K-1)/FLOAT(NBLD-1)
  XX(I,1) = SEVAL(S,XB,XPB,SB,IIB)
  YY(I,1) = SEVAL(S,YB,YPB,SB,IIB)
3 CONTINUE

```

C---- Set points on blade pressure surface

```

DO 4 K=1, NBLD
  I = NINL + K - 1
  S = SBLE + (SB(IIB)-SBLE)*FLOAT(K-1)/FLOAT(NBLD-1)
  XX(I,JJ) = SEVAL(S,XB,XPB,SB,IIB)
  YY(I,JJ) = SEVAL(S,YB,YPB,SB,IIB) + PITCH
4 CONTINUE

```

C---- set up metrics

```

DO 5 I=1, II
  XPOS(I) = FLOAT(I-1)/FLOAT(II-1)

```

```

6    CONTINUE

      DO 6 J = 1, JJ
        RJ = FLOAT(J-1)/FLOAT(JJ-1)
        YPOS(J) = RJ - 1.8 * ( (RJ-0.5) * ((RJ-0.5)**2-0.25) )
6    CONTINUE

C---- Initialize interior grid
      DO 7 I = 1, II
        DO 71 J = 2, JJ-1
          XX(I,J) = XX(I,1) + YPOS(J)*(XX(I,JJ)-XX(I,1))
          YY(I,J) = YY(I,1) + YPOS(J)*(YY(I,JJ)-YY(I,1))
71   CONTINUE
7    CONTINUE

      RETURN
      END

```

SUBROUTINE IMPROV

C---- Improves grid after elliptic grid generation

```

      INCLUDE 'TRI.INC'
      INCLUDE 'GRID.INC'

      DIMENSION SUM(Maxdim), XT1(Maxdim), YT1(Maxdim)
      DIMENSION XT2(Maxdim), YT2(Maxdim)

      DO 1 I = 1, II
        IM = I-1
        IP = I+1
        IF(I.EQ.1) IM=1
        IF(I.EQ.II) IP=II

        SUM(1) = 0.
        DO 11 J = 1, JJ-1
          JP = J+1
          XS = XX(IP,J)+XX(IP,JP) - XX(IM,J)-XX(IM,JP)
          YS = YY(IP,J)+YY(IP,JP) - YY(IM,J)-YY(IM,JP)
          SS = SQRT(XS*XS + YS*YS)
          XS = XS/SS
          YS = YS/SS
          SUM(JP) = SUM(J) + ABS( (XX(I,J)-XX(I,JP))*YS
                                & - (YY(I,J)-YY(I,JP))*XS )
11   CONTINUE

        J = 1
        DO 12 JO = 2, JJ-1
          SUMJ = FLOAT(JO-1)/FLOAT(JJ-1) * SUM(JJ)
121   IF(SUMJ.GT.SUM(J+1)) THEN
            J = J+1
            GOTO 121
          ENDIF
          ALPHA = (SUMJ-SUM(J)) / (SUM(J+1)-SUM(J))
          XT2(JO) = XX(I,J) + ALPHA*(XX(I,J+1)-XX(I,J))
          YT2(JO) = YY(I,J) + ALPHA*(YY(I,J+1)-YY(I,J))
12   CONTINUE

        DO 13 J = 2, JJ-1
          IF(I.NE.1) THEN
            XX(IM,J) = XT1(J)
            YY(IM,J) = YT1(J)
          ENDIF
          XT1(J) = XT2(J)
          YT1(J) = YT2(J)
          IF(I.EQ.II) THEN

```

```

        XX(I,J) = XT1(J)
        YY(I,J) = YT1(J)
      ENDIF
13      CONTINUE

1      CONTINUE
      RETURN
      END

```

```

C*****C
C
C   ISES - an Integrated Steamtube Euler Solver
C
C   Written by M. Giles and M.Drela
C
C   Copyright M.I.T. (1985)
C*****C
C
C   SUBROUTINE ELLIP(IMAX,JMAX,II,JJ,III,X,Y,YPOS,XPOS)
C   DIMENSION X(0:IMAX,0:JMAX), Y(0:IMAX,0:JMAX)
C   DIMENSION YPOS(JMAX), XPOS(IMAX)
C   CHARACTER*1 ANS
C
C   DIMENSION C(400),D(2,400)
C   IF(II.GT.400) STOP 'ELLIP dimensions must be increased'
C
C   ITMAX = 50
C
C   DSET1 = 1.0E-1
C   DSET2 = 5.0E-3
C   DSET3 = 2.0E-4
C
C   RLX1 = 1.30      !           DMAX > DSET1
C   RLX2 = 1.5C      !   DSET1 > DMAX > DSET2
C   RLX3 = 1.60      !   DSET2 > DMAX > DSET3
CCC  STOP           !   DSET3 > DMAX
C
C   RLX = RLX1
C
C   DO 1 ITER = 1, ITMAX
C
C     DMAX = 0.
C     DO 5 JO=2, JJ-1
C       JM = JO-1
C       JP = JO+1
C
C       IF(JO.EQ.JJJ) THEN
C         DO 2 IO=2, II-1
C           X(IO,JO) = X(IO,JM)
C         CONTINUE
C         GO TO 5
C       ELSE IF(JO.EQ.JJJ+1) THEN
C         DO 3 IO=2, II-1
C           X(IO,JO) = X(IO,JP)
C         CONTINUE
C         GO TO 5
C       ENDIF
C
C     DO 6 IO=2, II-1
C       IM = IO-1
C       IP = IO+1
C
C       XIM = X(IM,JM)
C       XOM = X(IO,JM)
C       XPM = X(IP,JM)
C       XMO = X(IM,JO)
C       XOO = X(IO,JO)

```

```

      XPO = X(IP,JO)
      XMP = X(IM,JP)
      XOP = X(IO,JP)
      XPP = X(IP,JP)
      YMM = Y(IM,JM)
      YOM = Y(IO,JM)
      YPM = Y(IP,JM)
      YMO = Y(IM,JO)
      YOO = Y(IO,JO)
      YPO = Y(IP,JO)
      YMP = Y(IM,JP)
      YOP = Y(IO,JP)
      YPP = Y(IP,JP)
C
      DXIM = XPOS(IO)-XPOS(IM)
      DXIP = XPOS(IP)-XPOS(IO)
      DXIAV = 0.5*(DXIM+DXIP)
C
      DETM = YPOS(JO)-YPOS(JM)
      DETP = YPOS(JP)-YPOS(JO)
      DETAV = 0.5*(DETM+DETP)
C
      DXDET = ( XOP - XOM ) / DETAV
      DYDET = ( YOP - YOM ) / DETAV
      DXDXI = ( XPO - XMO ) / DXIAV
      DYDXI = ( YPO - YMO ) / DXIAV
C
      ALF = DXDET**2 + DYDET**2
      BET = DXDET*DXDXI + DYDET*DYDXI
      GAM = DXDXI**2 + DYDXI**2
C
      CXIM = 1.0 / (DXIM+DXIAV)
      CXIP = 1.0 / (DXIP+DXIAV)
      CETM = 1.0 / (DETM+DETA)
      CETP = 1.0 / (DETP+DETA)
C
      B = -ALF*CXIM
      A = ALF*(CXIM+CXIP) + GAM*(CETM+CETP)
      C(IO) = -ALF*CXIP
      IF(IO.EQ.2) B = 0
C
      D(1,IO) = ALF*((XMO-XOO)*CXIM + (XPO-XOC)*CXIP)
      & - 2.0*BET*(XPP-XMP-XPM+XMM) / (4.0*DXIAV*DETA)
      & + GAM*((XOM-XOO)*CETM + (XOP-XOC)*CETP)
C
      D(2,IO) = ALF*((YMO-YOO)*CXIM + (YPO-YOO)*CXIP)
      & - 2.0*BET*(YPP-YMP-YPM+YMM) / (4.0*DXIAV*DETA)
      & + GAM*((YOM-YOO)*CETM + (YOP-YOC)*CETP)
C
      AINV = 1.0/(A - B+C(IM))
      C(IO) = C(IO) * AINV
      D(1,IO) = ( D(1,IO) - B*D(1,IM) ) * AINV
      D(2,IO) = ( D(2,IO) - B*D(2,IM) ) * AINV
C
6      CONTINUE
C
      D(1,II) = 0.
      D(2,II) = 0.
C
      IFIN = II-1
      DO 8 IBACK=2, IFIN
      IO = II-IBACK+1
      IP = IO+1
      D( 1,IO) = D( 1,IO) - C(IO)*D(1,IP)
      D( 2,IO) = D( 2,IO) - C(IO)*D(2,IP)
      X(IO,JO) = X(IO,JO) + RLX*D(1,IO)
      Y(IO,JO) = Y(IO,JO) + RLX*D(2,IO)
      AD1 = ABS(D(1,IO))
      AD2 = ABS(D(2,IO))
      DMAX = AMAX1(DMAX,AD1,AD2)
8      CONTINUE
C

```

```

6    CONTINUE
C
C    WRITE(6,*) 'Dmax = ', DMAX, RLX
C
C    RLX = RLX1
C    IF(DMAX.LT.DSET1) RLX = RLX2
C    IF(DMAX.LT.DSET2) RLX = RLX3
C    IF(DMAX.LT.DSET3) RETURN
C
1    CONTINUE
C
C    RETURN
C    END ! ELLIP

SUBROUTINE SPLINE(X,XP,S,II)
C
C    DIMENSION X(1),XP(1),S(1)
C    DIMENSION A(480),B(480),C(480)
C
C    IF(II.GT.480) STOP 'SPLINE: Array overflow'
C    DO 1 I = 1, II
C
C----- Beginning points
C    IF(I.EQ.1 .OR. S(I).EQ.S(I-1)) THEN
C        DSMI = 0.
C        DXM = 0.
C        DSPI = 1.0 / (S(I+1)-S(I))
C        DXP = X(I+1) - X(I)
C
C----- End points
C    ELSE IF(I.EQ.II .OR. S(I).EQ.S(I+1)) THEN
C        DSMI = 1.0 / (S(I) - S(I-1))
C        DXM = X(I) - X(I-1)
C        DSPI = 0.
C        DXP = 0.
C
C----- Interior points
C    ELSE
C        DSMI = 1.0 / (S(I) - S(I-1))
C        DXM = X(I) - X(I-1)
C        DSPI = 1.0 / (S(I+1) - S(I))
C        DXP = X(I+1) - X(I)
C
C    ENDIF
C
C    B(I) = DSMI
C    A(I) = 2.0 * (DSMI + DSPI)
C    C(I) = DSPI
C    XP(I) = 3.0 * (DXP*DSPI**2 + DXM*DSMI**2)
C
1    CONTINUE
C
C    CALL TRISOL(A,B,C,XP,II)
C
C    RETURN
C    END ! SPLINE

SUBROUTINE TRISOL(A,B,C,D,KK)
C
C    DIMENSION A(1),B(1),C(1),D(1)
C
C    DO 1 K = 2, KK
C        KM = K - 1

```

```

      C(KM) = C(KM) / A(KM)
      D(KM) = D(KM) / A(KM)
      A(K) = A(K) - B(K) * C(KM)
      D(K) = D(K) - B(K) * D(KM)
1    CONTINUE
C
      D(KK) = D(KK) / A(KK)
C
      DO 2 K = KK-1, 1, -1
        D(K) = D(K) - C(K) * D(K+1)
2    CONTINUE
C
      RETURN
END ! TRISOL

```

```

      FUNCTION SEVAL(SS,X,XP,S,N)
      REAL X(1), XP(1), S(1)
C
      ILOW = 1
      I = N
C
10  IF(I-ILOW .LE. 1) GO TO 11
C
      IMID = (I+ILOW)/2
      IF(SS .LT. S(IMID)) THEN
        I = IMID
      ELSE
        ILOW = IMID
      ENDIF
      GO TO 10
C
11  DS = S(I) - S(I-1)
      T = (SS-S(I-1)) / DS
      CX1 = DS*XP(I-1) - X(I) + X(I-1)
      CX2 = DS*XP(I) - X(I) + X(I-1)
      SEVAL = T*X(I) + (1.0-T)*X(I-1) + (T-T*T)*((1.0-T)*CX1 - T*CX2)
      RETURN
END ! SEVAL

```

```

      FUNCTION DEVAL(SS,X,XP,S,N)
      REAL X(1), XP(1), S(1)
C
      ILOW = 1
      I = N
C
10  IF(I-ILOW .LE. 1) GO TO 11
C
      IMID = (I+ILOW)/2
      IF(SS .LT. S(IMID)) THEN
        I = IMID
      ELSE
        ILOW = IMID
      ENDIF
      GO TO 10
C
11  DS = S(I) - S(I-1)
      T = (SS-S(I-1)) / DS
      CX1 = DS*XP(I-1) - X(I) + X(I-1)
      CX2 = DS*XP(I) - X(I) + X(I-1)
      DEVAL = (X(I)-X(I-1) + (1.-4.*T+3.*T*T)*CX1 + T*(3.*T-2.)*CX2)/DS
      RETURN
END ! DEVAL

```

```

subroutine pointers
implicit none

include 'GRID.INC'
include 'TRI.INC'
integer i, j
real*4 D1, D2
integer NM, FF, CC
!pointers
!diagonals of rectangle
!pointers

NM = 0
CC = 0
FF = 0

c** assign values to node arrays
do j = 0, Ny
do i = 0, Nx
NM = NM + 1
x(NM) = xx(i,j)
y(NM) = yy(i,j)
enddo
enddo
Nmax = NM

c** assign horizontal and vertical faces and cells
do j = 0, Ny-1
do i = 1, Nx
FF = FF + 1
face(FF,1) = 2*Nx*j + 1
face(FF,2) = Nx*(2*j-1) + 1
face(FF,3) = (Nx+1)*j + 1
face(FF,4) = (Nx+1)*j + 1 + 1
CC = CC + 1
cell(CC,1) = FF
cell(CC,6) = face(FF,3)
cell(CC,4) = face(FF,4)
enddo
do i = 1, Nx
FF = FF + 1
face(FF,3) = (Nx+1)*(j+1) + 1
face(FF,4) = (Nx+1)*j + 1
CC = CC + 1
cell(CC,1) = FF + Nx + 1
cell(CC,4) = (Nx+1)*(j+1) + 1
cell(CC,6) = (Nx+1)*(j+1) + 1 + 1
enddo
FF = FF + 1
face(FF,3) = (Nx+1)*(j+2)
face(FF,4) = (Nx+1)*(j+1)
enddo

Cmax = CC

c** assign top boundary faces
do i = 1, Nx
FF = FF + 1
face(FF,1) = 0
face(FF,2) = Nx*(2*Ny-1) + 1
face(FF,3) = (Nx+1)*Ny + 1
face(FF,4) = (Nx+1)*Ny + 1 + 1
face(FF,5) = 0
enddo

c** set bottom edge face array
do i = 1, Nx
face(i,2) = 0
face(i,6) = 0
enddo

```

```

c** set inlet edge face array
do j = 1, Ny
  face((2*Nx+1)*j-Nx,2) = 0
  face((2*Nx+1)*j-Nx,6) = 0
enddo

c** set outlet edge face array
do j = 1, Ny
  face((2*Nx+1)*j,1) = 0
  face((2*Nx+1)*j,5) = 0
enddo

c** split rectangular cells into two triangular cells in grid
do j = 1, Ny
  do i = 1, Nx
    FF = FF + 1
    D1 = (xx(1,j)-xx(1-1,j-1))**2 + (yy(1,j)-yy(1-1,j-1))**2
    D2 = (xx(1-1,j)-xx(1,j-1))**2 + (yy(1-1,j)-yy(1,j-1))**2
    if (D1 .gt. D2) then
c
      if (mod(i+j,2).eq.0) then
        face((2*Nx+1)*(j-1)+Nx+1,1) = 2*(j-1)*Nx + 1
        face((2*Nx+1)*j-Nx+1,2) = (2*j-1)*Nx + 1
        face((2*Nx+1)*(j-1)+Nx+1,5) = (Nx+1)*(j-1) + 1 + 1
        face((2*Nx+1)*j-Nx+1,6) = (Nx+1)*j + 1
        face((2*Nx+1)*(j-1)+1,5) = (Nx+1)*j + 1
        face((2*Nx+1)*j+1,6) = (Nx+1)*(j-1) + 1 + 1
        face(FF,3) = (Nx+1)*j + 1
        face(FF,4) = (Nx+1)*(j-1) + 1 + 1
        face(FF,5) = (Nx+1)*j + 1 + 1
        face(FF,6) = (Nx+1)*(j-1) + 1
        cell(2*Nx*(j-1)+1,2) = FF
        cell(2*Nx*(j-1)+1,3) = (2*Nx+1)*(j-1) + Nx + 1
        cell(2*Nx*(j-1)+1,5) = (Nx+1)*j + 1
        cell(2*Nx*(j-1)+Nx+1,2) = FF
        cell(2*Nx*(j-1)+Nx+1,3) = (2*Nx+1)*j - Nx + 1
        cell(2*Nx*(j-1)+Nx+1,5) = (Nx+1)*(j-1) + 1 + 1
      else
        face((2*Nx+1)*(j-1)+Nx+1,1) = (2*j-1)*Nx + 1
        face((2*Nx+1)*j-Nx+1,2) = 2*(j-1)*Nx + 1
        face((2*Nx+1)*(j-1)+Nx+1,5) = (Nx+1)*j + 1 + 1
        face((2*Nx+1)*j-Nx+1,6) = (Nx+1)*(j-1) + 1
        face((2*Nx+1)*(j-1)+1,5) = (Nx+1)*j + 1 + 1
        face((2*Nx+1)*j+1,6) = (Nx+1)*(j-1) + 1
        face(FF,3) = (Nx+1)*(j-1) + 1
        face(FF,4) = (Nx+1)*j + 1 + 1
        face(FF,5) = (Nx+1)*j + 1
        face(FF,6) = (Nx+1)*(j-1) + 1 + 1
        cell(2*Nx*(j-1)+1,2) = (2*Nx+1)*j - Nx + 1
        cell(2*Nx*(j-1)+1,3) = FF
        cell(2*Nx*(j-1)+1,5) = (Nx+1)*j + 1 + 1
        cell(2*Nx*(j-1)+Nx+1,2) = (2*Nx+1)*(j-1) + Nx + 1
        cell(2*Nx*(j-1)+Nx+1,3) = FF
        cell(2*Nx*(j-1)+Nx+1,5) = (Nx+1)*(j-1) + 1
      endif
      face(FF,1) = (2*j-1)*Nx + 1
      face(FF,2) = 2*(j-1)*Nx + 1
    enddo
  enddo

  Fmax = FF

  return
end

subroutine bpointers
implicit none

```



```

include 'GRID.INC'
include 'TRI.INC'
integer i, j, n
integer FF, EH, EF, IN, ON, WN, PN
integer CC
integer F1, F2
real dxM, dyM, dxP, dyP, dsM, dsP, dx, dy
integer node

!pointers
!pointers
!periodic faces
!change in x & y near node
!current node processed

EN = 0
EF = 0
IN = 0
ON = 0
WN = 0
PN = 0

c** wall boundaries
if (pitch.eq.0.) then
  do i = 1, Nx
    EF = EF + 1
    eface(EF) = Ny*(2*Nx+1) + 1
    EN = EN + 1
    node = (Nx+1)*Ny + 1
    enode(EN) = node
    dxP = x(node+1) - x(node)
    dyP = y(node+1) - y(node)
    if (i .eq. 1) then
      dxM = -dxP
      dyM = -dyP
    else
      dxM = x(node-1) - x(node)
      dyM = y(node-1) - y(node)
    endif
    if ((dxM*dxP + dyM*dyP) .gt. 0.) then
      dsM = sqrt(dxM**2 + dyM**2)
      dsP = sqrt(dxP**2 + dyP**2)
      dx = dxM/dsM + dxP/dsP
      dy = dyM/dsM + dyP/dsP
    else
      dx = dxP - dxM
      dy = dyP - dyM
    endif
    senode(EN) = dy/sqrt(dx**2 + dy**2)
    cenode(EN) = dx/sqrt(dx**2 + dy**2)
  enddo

  EN = EN + 1
  node = (Nx+1)*(Ny+1)
  enode(EN) = node
  dxM = x(node-1) - x(node)
  dyM = y(node-1) - y(node)
  dxP = -dxM
  dyP = -dyM
  dx = dxP - dxM
  dy = dyP - dyM
  senode(EN) = dy/sqrt(dx**2 + dy**2)
  cenode(EN) = dx/sqrt(dx**2 + dy**2)

  tnode = EN

  do i = 1, Nx
    EF = EF + 1
    eface(EF) = 1
    EN = EN + 1
    node = 1
    enode(EN) = node
    dxP = x(node+1) - x(node)
    dyP = y(node+1) - y(node)
    if (i .eq. 1) then
      dxM = -dxP
      dyM = -dyP

```

```

        else
            dxM = x(node-1) - x(node)
            dyM = y(node-1) - y(node)
        endif
        if ((dxM*dxP + dyM*dyP) .gt. 0.) then
            dsM = sqrt(dxM**2 + dyM**2)
            dsP = sqrt(dxP**2 + dyP**2)
            dx = dxM/dsM + dxP/dsP
            dy = dyM/dsM + dyP/dsP
        else
            dx = dxP - dxM
            dy = dyP - dyM
        endif
        senode(EN) = dy/sqrt(dx**2 + dy**2)
        cenode(EN) = dx/sqrt(dx**2 + dy**2)
    enddo

    EN = EN + 1
    node = Nx + 1
    enode(EN) = node
    dxM = x(node-1) - x(node)
    dyM = y(node-1) - y(node)
    dxP = -dxM
    dyP = -dyM
    dx = dxP - dxM
    dy = dyP - dyM
    senode(EN) = dy/sqrt(dx**2 + dy**2)
    cenode(EN) = dx/sqrt(dx**2 + dy**2)

    bnode = EN
    Wmax = 2*Nx
    else if (pitch.ne.0.) then
c** periodic nodes
        do i = 1, ILE
            PN = PN + 1
            pnode(PN,1) = 1
            pnode(PN,2) = (Nx+1)*Ny + 1
        enddo

        do i = ITE, Nx+1
            PN = PN + 1
            pnode(PN,1) = 1
            pnode(PN,2) = (Nx+1)*Ny + 1
        enddo
        Pmax = ILE + Nx + 2 - ITE

        do i = 1, ILE-1
            F1 = 1
            F2 = Ny*(2*Nx+1) + 1
            face(F1,2) = face(F2,2)
            face(F1,6) = face(F2,6)
            face(F2,1) = face(F1,1)
            face(F2,5) = face(F1,5)
        enddo

        do i = ITE, Nx
            F1 = 1
            F2 = Ny*(2*Nx+1) + 1
            face(F1,2) = face(F2,2)
            face(F1,6) = face(F2,6)
            face(F2,1) = face(F1,1)
            face(F2,5) = face(F1,5)
        enddo

c** the rest of the edge nodes and faces

c** lower edge of airfoil
        do i = ILE, ITE-1
            EN = EN + 1
            node = (Nx+1)*Ny + 1
            enode(EN) = node
            dxP = x(node+1) - x(node)

```

```

dyP = y(node+1) - y(node)
if (i .eq. ILE) then
  dxM = x(ILE+1) - x(node)
  dyM = y(ILE+1) + pitch - y(node)
else
  dxM = x(node-1) - x(node)
  dyM = y(node-1) - y(node)
endif
if ((dxM*dxP + dyM*dyP) .gt. 0.) then
  dsM = sqrt(dxM**2 + dyM**2)
  dsP = sqrt(dxP**2 + dyP**2)
  dx = dxM/dsM + dxP/dsP
  dy = dyM/dsM + dyP/dsP
else
  dx = dxP - dxM
  dy = dyP - dyM
endif
enode(EN) = dy/sqrt(dx**2 + dy**2)
cenode(EN) = dx/sqrt(dx**2 + dy**2)
FF = Ny*(2*Nx+1) + 1
do n = 1, 6
  face(FF-ILE+1,n) = face(FF,n)
enddo
EF = EF + 1
eface(EF) = FF - ILE + 1
enddo

do FF = Ny*(2*Nx+1)+Nx+1, Fmax
  CC = face(FF,1)
  cell(CC,1) = FF-Nx+ITE-ILE
  do n = 1, 6
    face(FF-Nx+ITE-ILE,n) = face(FF,n)
  enddo
enddo

Fmax = Fmax - Nx + ITE - ILE

EN = EN + 1
node = (Nx+1)*Ny + ITE
enode(EN) = node
dxM = x(node-1) - x(node)
dyM = y(node-1) - y(node)
dxP = x(ITE-1) - x(node)
dyP = y(ITE-1) + pitch - y(node)
if ((dxM*dxP + dyM*dyP) .gt. 0.) then
  dsM = sqrt(dxM**2 + dyM**2)
  dsP = sqrt(dxP**2 + dyP**2)
  dx = dxM/dsM + dxP/dsP
  dy = dyM/dsM + dyP/dsP
else
  dx = dxP - dxM
  dy = dyP - dyM
endif
enode(EN) = dy/sqrt(dx**2 + dy**2)
cenode(EN) = dx/sqrt(dx**2 + dy**2)

tnode = EN

c** upper edge of airfoil
do i = ILE, ITE-1
  EN = EN + 1
  EF = EF + 1
  eface(EF) = 1
  node = 1
  enode(EN) = node
  dxP = x(node+1) - x(node)
  dyP = y(node+1) - y(node)
  if (i .eq. ILE) then
    dxM = x((Nx+1)*Ny+ILE+1) - x(node)
    dyM = y((Nx+1)*Ny+ILE+1) - pitch - y(node)
  else
    dxM = x(node-1) - x(node)

```

```

        dyM = y(node-1) - y(node)
    endif
    if ((dxM*dxP + dyM*dyP) .gt. 0.) then
        dsM = sqrt(dxM**2 + dyM**2)
        dsP = sqrt(dxP**2 + dyP**2)
        dx = dxM/dsM + dxP/dsP
        dy = dyM/dsM + dyP/dsP
    else
        dx = dxP - dxM
        dy = dyP - dyM
    endif
    senode(EN) = dy/sqrt(dx**2 + dy**2)
    cenode(EN) = dx/sqrt(dx**2 + dy**2)
enddo

EN = EN + 1
node = ITE
enode(EN) = node
dxM = x(node-1) - x(node)
dyM = y(node-1) - y(node)
dxP = x((Nx+1)*Ny+ITE-1) - x(node)
dyP = y((Nx+1)*Ny+ITE-1) - pitch - y(node)
if ((dxM*dxP + dyM*dyP) .gt. 0.) then
    dsM = sqrt(dxM**2 + dyM**2)
    dsP = sqrt(dxP**2 + dyP**2)
    dx = dxM/dsM + dxP/dsP
    dy = dyM/dsM + dyP/dsP
else
    dx = dxP - dxM
    dy = dyP - dyM
endif
senode(EN) = dy/sqrt(dx**2 + dy**2)
cenode(EN) = dx/sqrt(dx**2 + dy**2)

bnode = EN
Wmax = 2*(ITE-ILE)

endif

c** inlet and outlet boundaries
do j = 1, Ny
    EN = EN + 1
    enode(EN) = (j-1)*(Nx+1) + 1
    IN = IN + 1
    innode(IN) = (j-1)*(Nx+1) + 1
    EF = EF + 1
    eface(EF) = (2*Nx+1)*j - Nx
enddo

EN = EN + 1
enode(EN) = Ny*(Nx+1) + 1
IN = IN + 1
innode(IN) = Ny*(Nx+1) + 1

inode = bnode + Ny + 1
imax = IN

do j = 1, Ny
    EN = EN + 1
    enode(EN) = (Nx+1)*j
    ON = ON + 1
    outnode(ON) = (Nx+1)*j
    EF = EF + 1
    eface(EF) = (2*Nx+1)*j
enddo

EN = EN + 1
enode(EN) = (Nx+1)*(Ny+1)
ON = ON + 1
outnode(ON) = (Nx+1)*(Ny+1)

ENmax = EN

```

```

Emax = EF
Omax = ON

```

```

return
end

```

```

c*****
c*
c*   renumber edges so nodes are consecutive along edges
c*   and NS is the interior node
c*
c*****

```

```

subroutine edgenumber
implicit none

```

```

include 'TRI.INC'
integer EF, FF, i           !pointers
integer save                !save value in face

```

```

do EF = 1, Emax
  FF = eface(EF)
  if (face(FF,1).eq.0) then
    do i = 1, 5, 2
      save = face(FF,i)
      face(FF,i) = face(FF,i+1)
      face(FF,i+1) = save
    enddo
  endif
enddo

```

```

return
end

```

```

subroutine cellcolor
implicit none

```

```

include 'TRI.INC'

```

```

integer CC, NM, FF           !pointers
integer Ncells               !number of cells colored
integer CC2(0:Maxcells)      !new number of cell
integer Ncolor               !current color
integer cell2(Maxcells,6)    !new cell number
logical hascolor(Maxcells)   !has node been colored current color?

```

```

c** Initialize everything
  Ncells = 0
  do CC = 1, Cmax
    CC2(CC) = 0
  enddo

```

```

c** Loop over colors
  do Ncolor = 1, 50

```

```

c** Initialize node color array
  NCcolor(Ncolor) = 0
  do NM = 1, Nmax
    hascolor(NM) = .FALSE.
  enddo

```

```

c** Loop over cells
  do CC = 1, Cmax

```

```

c** Check if cell is already colored with old color

```

```

        if(CC2(CC).NE.0) GOTO 22
c** Check if cell nodes are already colored with new color
        if( hascolor(cell(CC,4)) .OR.
&          hascolor(cell(CC,5)) .OR.
&          hascolor(cell(CC,6)) ) GOTO 22

c** Set color markers
        Ncells = Ncells + 1
        CC2(CC) = Ncells
        NCcolor(Ncolor) = NCcolor(Ncolor) + 1
        hascolor(cell(CC,4)) = .TRUE.
        hascolor(cell(CC,5)) = .TRUE.
        hascolor(cell(CC,6)) = .TRUE.

22      enddo

        if(Ncells.EQ.Cmax) GOTO 23

        enddo

        STOP 'COLOR; more than 50 colors required for cells'

23      Ccolormax= Ncolor

c** Redo pointers
        do CC = 1, Cmax
            cell12(CC2(CC),1) = cell(CC,1)
            cell12(CC2(CC),2) = cell(CC,2)
            cell12(CC2(CC),3) = cell(CC,3)
            cell12(CC2(CC),4) = cell(CC,4)
            cell12(CC2(CC),5) = cell(CC,5)
            cell12(CC2(CC),6) = cell(CC,6)
        enddo

        do CC = 1, Cmax
            cell(CC,1) = cell12(CC,1)
            cell(CC,2) = cell12(CC,2)
            cell(CC,3) = cell12(CC,3)
            cell(CC,4) = cell12(CC,4)
            cell(CC,5) = cell12(CC,5)
            cell(CC,6) = cell12(CC,6)
        enddo

        CC2(0) = 0

        do FF = 1, Fmax
            face(FF,1) = CC2(face(FF,1))
            face(FF,2) = CC2(face(FF,2))
        enddo

        return
end

subroutine facecolor
implicit none

include 'TRI.INC'
integer FF, NN, CC, EF
integer Nfaces
integer FF2(Maxfaces)
integer Ncolor
integer face2(Maxfaces,6)
logical hascolor(0:Maxfaces)

!pointers
!number of faces colored
!new number of face
!current color
!new face number
!has node been colored current color?

c** Initialize everything
        Nfaces = 0

```

```

        do FF = 1, Fmax
            FF2(FF) = 0
        enddo

c** Loop over colors
        do Ncolor = 1, 50

c** Initialize node color array
            NFcolor(Ncolor) = 0
            do NN = 1, Nmax
                hascolor(NN) = .FALSE.
            enddo

c** Loop over faces
            do FF = 1, Fmax

                hascolor(0) = .FALSE.

c** Check if face is already colored with old color
                if(FF2(FF).NE.0) GOTO 22

c** Check if face nodes are already colored with new color
                if( hascolor(face(FF,3)) .OR.
&                 hascolor(face(FF,4)) .OR.
&                 hascolor(face(FF,5)) .OR.
&                 hascolor(face(FF,6)) ) GOTO 22

c** Set color markers
                    Nfaces = Nfaces + 1
                    FF2(FF) = Nfaces
                    NFcolor(Ncolor) = NFcolor(Ncolor) + 1
                    hascolor(face(FF,3)) = .TRUE.
                    hascolor(face(FF,4)) = .TRUE.
                    hascolor(face(FF,5)) = .TRUE.
                    hascolor(face(FF,6)) = .TRUE.

22                enddo

                    if(Nfaces.EQ.Fmax) GOTO 23

                    enddo

                    STOP 'COLOR; more than 50 colors required for faces'

23                Fcolormax= Ncolor

c** Redo pointers
                do FF = 1, Fmax
                    face2(FF2(FF),1) = face(FF,1)
                    face2(FF2(FF),2) = face(FF,2)
                    face2(FF2(FF),3) = face(FF,3)
                    face2(FF2(FF),4) = face(FF,4)
                    face2(FF2(FF),5) = face(FF,5)
                    face2(FF2(FF),6) = face(FF,6)
                enddo

                do FF = 1, Fmax
                    face(FF,1) = face2(FF,1)
                    face(FF,2) = face2(FF,2)
                    face(FF,3) = face2(FF,3)
                    face(FF,4) = face2(FF,4)
                    face(FF,5) = face2(FF,5)
                    face(FF,6) = face2(FF,6)
                enddo

                do CC = 1, Cmax
                    cell(CC,1) = FF2(cell(CC,1))
                    cell(CC,2) = FF2(cell(CC,2))
                    cell(CC,3) = FF2(cell(CC,3))
                enddo

                do EF = 1, Emax

```

```

        eface(EF) = FF2(eface(EF))
    enddo

    return
end

subroutine edgecolor
implicit none

include 'TRI.INC'
integer FF, NN, EF          !pointers
integer Nedges              !number of faces colored
integer EF2(Maxedges)       !new number of face
integer Ncolor              !current color
integer eface2(Maxedges)    !new face number
logical hascolor(Maxnodes)  !has node been colored current color?

c** Initialize everything
    Nedges = 0
    do EF = 1, Emax
        EF2(EF) = 0
    enddo

c** Loop over colors
    do Ncolor = 1, 5

c** Initialize node color array
        NEcolor(Ncolor) = 0
        do NN = 1, Nmax
            hascolor(NN) = .FALSE.
        enddo

c** Loop over faces
        do EF = 1, Wmax
            FF = eface(EF)

c** Check if face is already colored with old color
            if(EF2(EF).NE.0) GOTO 22

c** Check if face nodes are already colored with new color
            if( hascolor(face(FF,3)) .OR.
&             hascolor(face(FF,4)) .OR.
&             hascolor(face(FF,5)) ) GOTO 22

c** Set color markers
                Nedges = Nedges + 1
                EF2(EF) = Nedges
                NEcolor(Ncolor) = NEcolor(Ncolor) + 1
                hascolor(face(FF,3)) = .TRUE.
                hascolor(face(FF,4)) = .TRUE.
                hascolor(face(FF,5)) = .TRUE.

22            enddo

                if(Nedges.EQ.Wmax) GOTO 23

        enddo

        STOP 'COLOR; more than 5 colors required for wall edges'

23        Eicolormax = Ncolor

c** Loop over colors
        do Ncolor = Eicolormax+1, 10

c** Initialize node color array
            NEcolor(Ncolor) = 0
            do NN = 1, Nmax
                hascolor(NN) = .FALSE.
            enddo

```



```

        enddo

c** Loop over faces
      do EF = Wmax+1, Emax
        FF = eface(EF)

c** Check if face is already colored with old color
        if(EF2(EF).NE.0) GOTO 32

c** Check if face nodes are already colored with new color
        if( hascolor(face(FF,3)) .OR.
           &   hascolor(face(FF,4)) .OR.
           &   hascolor(face(FF,5)) ) GOTO 32

c** Set color markers
          Nedges = Nedges + 1
          EF2(EF) = Nedges
          NEcolor(Ncolor) = NEcolor(Ncolor) + 1
          hascolor(face(FF,3)) = .TRUE.
          hascolor(face(FF,4)) = .TRUE.
          hascolor(face(FF,5)) = .TRUE.

32      enddo

          if(Nedges.EQ.Emax) GOTO 33

        enddo

        STOP 'COLOR; more than 10 colors required for all edges'

33      E2colormax= Ncolor

c** Redo pointers
      do EF = 1, Emax
        eface2(EF2(EF)) = eface(EF)
      enddo

      do EF = 1, Emax
        eface(EF) = eface2(EF)
      enddo

      return
      end

```

A.1.3 Ni Scheme

```

c*****
c*
c*      main program for triangular Ni scheme
c*
c*****

      program triangle
      implicit none

      include 'TRI.INC'
      integer Niter           !number of iterations
      integer NN, CC, 1       !pointer
      integer N1, N2, N3      !nodes at corners of cell
      real maxchange          !max change in state vector
      integer maxnode, maxeqn !where max change occurs

```

```

        real onethird          !one divided by three

c* read in data from file
    call gridio(1)
    call flowio(1)
    call input

c* calculate control area around each cell
    call calcarea

    Niter = 0
    durms = 999.
    onethird = 1./3.

c* start history file from the top
    open(unit=35,status='unknown',form='formatted')
    write(35,2) Min1
    close(unit=35)
    2    format(' inlet Mach number = ',f5.3)

c* loop until converged
    do while ((Niter.lt.Maxiter) .and. (durms.gt.2.e-7))
        Niter = Niter + 1

c* set cell values of state vector
        do CC = 1, Cmax
            N1 = cell(CC,4)
            N2 = cell(CC,5)
            N3 = cell(CC,6)
            Uc(1,CC) = (U(1,N1) + U(1,N2) + U(1,N3))*onethird
            Uc(2,CC) = (U(2,N1) + U(2,N2) + U(2,N3))*onethird
            Uc(3,CC) = (U(3,N1) + U(3,N2) + U(3,N3))*onethird
            Uc(4,CC) = (U(4,N1) + U(4,N2) + U(4,N3))*onethird
        enddo

c* calculate time step for each cell
        call timestep

c* calculate flux at each node
        call nodeflux

c* calculate change in state vector and fluxes at each cell
        call delcell

c* calculate change in state vector at each node
        call delstate

c* add smoothing term
        call smooth

c** account for periodic nodes
        call bperiodic

c* set inlet and outlet boundary conditions
        call binlet
        call boutlet

c* change momentum change to make flow tangent at walls
        call tangent

c* update state vector value
        do NN = 1, Nmax
            U(1,NN) = U(1,NN) + dU(1,NN)
            U(2,NN) = U(2,NN) + dU(2,NN)
            U(3,NN) = U(3,NN) + dU(3,NN)
            U(4,NN) = U(4,NN) + dU(4,NN)
        enddo

c* find root mean square difference in state vector
        if (mod(Niter,10).eq.0 .or. Niter.lt.10) then
            durms = 0.0
            maxchange = 0.

```

```

do NN = 1, Nmax
  do i = 1, 4
    durms = durms + dU(1,NN)**2
    if (abs(dU(1,NN)).gt.abs(maxchange)) then
      maxchange = dU(1,NN)
      maxnode = NN
      maxeqn = 1
    endif
  enddo
enddo

durms = sqrt(durms/(4.*Nmax))

c* print diagnostics to screen
call flowio(0)
open(unit=50, status='unknown', form='unformatted')
write(50) Cmax, (vol(CC),CC=1,Cmax)
close(unit=50)

10      open(unit=35, status='old',access='append',err=10)
      &   write(35,1) Niter, durms, maxchange, x(maxnode),
      &       y(maxnode), maxeqn
      close(unit=35)
      &   write(6,1) Niter, durms, maxchange, x(maxnode),
      &       y(maxnode), maxeqn
      endif
    enddo
1      format('Niter=',i4,' rms=',f9.7,' max=',f9.7,' x=',
      &         f6.3,' y=',f6.3,' eqn=',i1)

c* write out data to file
call flowio(0)

stop
end

```

```

C*****
C*
C* calculate areas of cells and distribute to nodal area *
C* which is made up of the sum of the cell areas *
C* surrounding the node *
C*
C*****

```

```

subroutine calcaarea
implicit none

include 'TRI.INC'
integer CC
integer N1, N2, N3
real dx12, dx31, dy12, dy31
!pointer
!nodes at corner of cell
!length of cell edges

c* calculate area of cells
do CC = 1, Cmax
  N1 = cell(CC,4)
  N2 = cell(CC,5)
  N3 = cell(CC,6)
  dx12 = x(N1) - x(N2)
  dx31 = x(N3) - x(N1)
  dy12 = y(N1) - y(N2)
  dy31 = y(N3) - y(N1)
  areaC(CC) = abs(dx31*dy12 - dy31*dx12)
enddo

return
end

```

```

c*****
c*
c*      calculate time step for nodes
c*
c*****

      subroutine timestep
      implicit none

      include 'TRI.IWC'
      real onethird
      integer NE, CC, PN, EN
      integer N1, N2, N3
      integer P1, P2
      real delx, dely
      real delside
      real uu, vv, aa, rr, ww2
      integer CC1, CC2, col

      !one divided by three
      !pointer
      !nodes related to face
      !periodic nodes
      !x and y length of side
      !length of side
      !values at node
      !pointers

      onethird = 1./3.

      do NN = 1, Nmax
         deltN(NN) = 0.
      enddo

c** find time step for each cell
c** delt is really delt/area
      CC2 = 0

      do col = 1, Ccolormax
         CC1 = CC2 + 1
         CC2 = CC1 - 1 + NCcolor(col)

CVD$      NODEPCHK
            do CC = CC1, CC2
               N1 = cell(CC,4)
               N2 = cell(CC,5)
               N3 = cell(CC,6)

               rr = Uc(1,CC)
               uu = Uc(2,CC)/rr
               vv = Uc(3,CC)/rr
               ww2 = uu**2 + vv**2
               aa = sqrt(gam*gam*(Uc(4,CC)/rr - 0.5*ww2))

               delx = x(N2) - x(N1)
               dely = y(N2) - y(N1)
               delside = sqrt(delx**2 + dely**2)

               deltC(CC) = abs(uu*dely-vv*delx) + aa*delside

               delx = x(N3) - x(N2)
               dely = y(N3) - y(N2)
               delside = sqrt(delx**2 + dely**2)

               deltC(CC) = deltC(CC) + abs(uu*dely-vv*delx) + aa*delside

               delx = x(N1) - x(N3)
               dely = y(N1) - y(N3)
               delside = sqrt(delx**2 + dely**2)

               deltC(CC) = deltC(CC) + abs(uu*dely-vv*delx) + aa*delside

               deltC(CC) = CFL*2./deltC(CC)

               deltN(N1) = deltN(N1) + onethird/deltC(CC)
               deltN(N2) = deltN(N2) + onethird/deltC(CC)
            enddo
      enddo

```

```

        deltN(N3) = deltN(N3) + onethird/deltC(CC)
    enddo
enddo

CVD$  NODEPCHZ
do PH = 1, Pmax
    P1 = pnode(PH,1)
    P2 = pnode(PH,2)
    deltN(P1) = deltN(P1) + deltN(P2)
    deltN(P2) = deltN(P1)
enddo

do NN = 1, Nmax
    deltN(NN) = 1./deltN(NN)
enddo

return
end

c*****
c*
c*   calculate flux vector values at nodes
c*
c*****

subroutine nodeflux
implicit none

include 'TRI.INC'
integer NN                !pointers
real WW                   !kinetic energy

do NN = 1, Nmax
c* calculate f and g at nodes
    WW = 0.5*(U(2,NN)**2 + U(3,NN)**2)/U(1,NN)
    F(1,NN) = U(2,NN)
    F(2,NN) = U(2,NN)**2/U(1,NN) + gam1*(U(4,NN) - WW)
    F(3,NN) = U(2,NN)*U(3,NN)/U(1,NN)
    F(4,NN) = (U(2,NN)/U(1,NN))*(gam*U(4,NN) - gam1*WW)

    G(1,NN) = U(3,NN)
    G(2,NN) = U(2,NN)*U(3,NN)/U(1,NN)
    G(3,NN) = U(3,NN)**2/U(1,NN) + gam1*(U(4,NN) - WW)
    G(4,NN) = (U(3,NN)/U(1,NN))*(gam*U(4,NN) - gam1*WW)
enddo

return
end

c*****
c*
c*   this subroutine finds the change in the fluxes
c*   F and G and the change in the state U at each cell
c*
c*****

subroutine delcell
implicit none

include 'TRI.INC'
integer CC                !pointers
integer N1, N2, N3        !nodes at corners of cells

```

```

      real u1, u2, u3, v1, v2, v3      !values at nodes
      real coef
      real dx23, dx31, dx12             !change in x on cell edges
      real dy23, dy31, dy12             !change in y on cell edges

c* find change in state vector for cell
do CC = 1, Cmax
  N1 = cell(CC,4)
  N2 = cell(CC,5)
  N3 = cell(CC,6)

  u1 = U(2,N1)/U(1,N1)
  u2 = U(2,N2)/U(1,N2)
  u3 = U(2,N3)/U(1,N3)
  v1 = U(3,N1)/U(1,N1)
  v2 = U(3,N2)/U(1,N2)
  v3 = U(3,N3)/U(1,N3)

  dy23 = y(N2) - y(N3)
  dy31 = y(N3) - y(N1)
  dy12 = y(N1) - y(N2)
  dx23 = x(N2) - x(N3)
  dx31 = x(N3) - x(N1)
  dx12 = x(N1) - x(N2)

  coef = 0.5*deltC(CC)

  dUc(1,CC) = coef*(-F(1,N1)*dy23 + G(1,N1)*dx23
&               -F(1,N2)*dy31 + G(1,N2)*dx31
&               -F(1,N3)*dy12 + G(1,N3)*dx12)

  dUc(2,CC) = coef*(-F(2,N1)*dy23 + G(2,N1)*dx23
&               -F(2,N2)*dy31 + G(2,N2)*dx31
&               -F(2,N3)*dy12 + G(2,N3)*dx12)

  dUc(3,CC) = coef*(-F(3,N1)*dy23 + G(3,N1)*dx23
&               -F(3,N2)*dy31 + G(3,N2)*dx31
&               -F(3,N3)*dy12 + G(3,N3)*dx12)

  dUc(4,CC) = coef*(-F(4,N1)*dy23 + G(4,N1)*dx23
&               -F(4,N2)*dy31 + G(4,N2)*dx31
&               -F(4,N3)*dy12 + G(4,N3)*dx12)

c** shock smoothing
  vol(CC) = u1*dy23 - v1*dx23 +
&          u2*dy31 - v2*dx31 +
&          u3*dy12 - v3*dx12
  enddo

c* implement wall boundary conditions
  call bwall

  return
end

c*****
c*
c*   this subroutine applies the wall boundary condition
c*
c*****

subroutine bwall
  implicit none

  include 'TRI.INC'
  integer EF, FF, CC      !pointer
  integer N1, N3           !nodes on edge
  real u1, u3, v1, v3      !values at nodes

```

```

      real coef
      real dy13, dx13      !change in x and y
      real pres1, pres3    !pressure
      integer EF1, EF2, col

c*  upper and lower boundary
      EF2 = 0

      do col = 1, Eicolormax
        EF1 = EF2 + 1
        EF2 = EF1 - 1 + NEcolor(col)

CVD$  NODEPCHK
      do EF = EF1, EF2
        FF = eface(EF)
        CC = face(FF,1)
        N1 = face(FF,4)
        N3 = face(FF,3)

        u1 = U(2,N1)/U(1,N1)
        u3 = U(2,N3)/U(1,N3)
        v1 = U(3,N1)/U(1,N1)
        v3 = U(3,N3)/U(1,N3)

        coef = 0.5*deltC(CC)

        dy13 = y(N1) - y(N3)
        dx13 = x(N1) - x(N3)

        pres1 = gam1*(U(4,N1) - 0.5*(U(2,N1)**2 + U(3,N1)**2)/
&              U(1,N1))
        pres3 = gam1*(U(4,N3) - 0.5*(U(2,N3)**2 + U(3,N3)**2)/
&              U(1,N3))

        dUc(1,CC) = dUc(1,CC) + coef*(
&              (F(1,N3) + F(1,N1))*dy13 -
&              (G(1,N3) + G(1,N1))*dx13)

        dUc(2,CC) = dUc(2,CC) + coef*(
&              (F(2,N3) + F(2,N1) - pres1 - pres3)*dy13 -
&              (G(2,N3) + G(2,N1))*dx13)

        dUc(3,CC) = dUc(3,CC) + coef*(
&              (F(3,N3) + F(3,N1))*dy13 -
&              (G(3,N3) + G(3,N1) - pres1 - pres3)*dx13)

        dUc(4,CC) = dUc(4,CC) + coef*(
&              (F(4,N3) + F(4,N1))*dy13 -
&              (G(4,N3) + G(4,N1))*dx13)

c** shock smoothing
        vol(CC) = vol(CC) - (u1 + u3)*dy13 + (v1 + v3)*dx13

      enddo
    enddo

    return
  end

c*****
c*
c*      this subroutine calculates the change at each node
c*
c*****

      subroutine delstate
      implicit none

```

```

include 'TRI.INC'
integer NH, CC, FF, EF           !pointers
integer N1, N2, N3               !nodes at corners of cell
real rr, uu, vv, ww2            !values at cell
real rdu, rdv                   !part of dFc and dGc
real HH, dp
real dFc(4), dGc(4)
real dy21, dy32, dy13           !change in y on cell edges
real dx21, dx32, dx13           !change in x on cell edges
real CON                         !coefficient
integer CC1, CC2, col           !pointers
integer EF1, EF2
real coef                       !coefficient
real onethird                   !one divided by three

do NH = 1, Nmax
  dU(1,NH) = 0.
  dU(2,NH) = 0.
  dU(3,NH) = 0.
  dU(4,NH) = 0.
enddo

onethird = 1./3.

CC2 = 0

do col = 1, Ccolormax
  CC1 = CC2 + 1
  CC2 = CC1 - 1 + NCcolor(col)
CVD$ NODEPCHK
  do CC = CC1, CC2
    N1 = cell(CC,4)
    N2 = cell(CC,5)
    N3 = cell(CC,6)

    rr = Uc(1,CC)
    uu = Uc(2,CC)/rr
    vv = Uc(3,CC)/rr
    ww2 = 0.5*(uu*uu + vv*vv)

    dy21 = y(N2) - y(N1)
    dy32 = y(N3) - y(N2)
    dy13 = y(N1) - y(N3)
    dx21 = x(N2) - x(N1)
    dx32 = x(N3) - x(N2)
    dx13 = x(N1) - x(N3)

c* find second order change in flux vector at cells
    HH = gam*Uc(4,CC)/Uc(1,CC) - gam1*ww2
    rdu = dUc(2,CC) - uu*dUc(1,CC)
    rdv = dUc(3,CC) - vv*dUc(1,CC)
    dp = gam1*(dUc(4,CC) - uu*dUc(2,CC) - vv*dUc(3,CC)
    & + ww2*dUc(1,CC))

    dFc(1) = dUc(2,CC)
    dFc(2) = uu*(dUc(2,CC) + rdu) + dp
    dFc(3) = vv*dUc(2,CC) + uu*rdv
    dFc(4) = uu*(dUc(4,CC) + dp) + HH*rdu

    dGc(1) = dUc(3,CC)
    dGc(2) = uu*dUc(3,CC) + vv*rdu
    dGc(3) = vv*(dUc(3,CC) + rdv) + dp
    dGc(4) = vv*(dUc(4,CC) + dp) + HH*rdv

c** shock smoothing
c  CON = 0.25*min(0.,vol(CC))*vol(CC)*ww2/areaC(CC)
    vol(CC) = vol(CC)/sqrt(2.*areaC(CC))
    CON = epsicoef*0.5*max(-0.1,min(0.,vol(CC)))*vol(CC)*ww2
c  vol(CC) = -vol(CC)/sqrt(2.*areaC(CC))
c  CON = epsicoef*0.5*max(-0.1,min(0.,vol(CC)))*vol(CC)*ww2
    dFc(2) = dFc(2) + CON

```



```

dGc(3) = dGc(3) + CON

coef = onethird/deltC(CC)

dU(1,N1) = dU(1,N1) + coef*dUc(1,CC)
*      - 0.25*(dFc(1)*dy32 - dGc(1)*dx32)
dU(1,N2) = dU(1,N2) + coef*dUc(1,CC)
*      - 0.25*(dFc(1)*dy13 - dGc(1)*dx13)
dU(1,N3) = dU(1,N3) + coef*dUc(1,CC)
*      - 0.25*(dFc(1)*dy21 - dGc(1)*dx21)

dU(2,N1) = dU(2,N1) + coef*dUc(2,CC)
*      - 0.25*(dFc(2)*dy32 - dGc(2)*dx32)
dU(2,N2) = dU(2,N2) + coef*dUc(2,CC)
*      - 0.25*(dFc(2)*dy13 - dGc(2)*dx13)
dU(2,N3) = dU(2,N3) + coef*dUc(2,CC)
*      - 0.25*(dFc(2)*dy21 - dGc(2)*dx21)

dU(3,N1) = dU(3,N1) + coef*dUc(3,CC)
*      - 0.25*(dFc(3)*dy32 - dGc(3)*dx32)
dU(3,N2) = dU(3,N2) + coef*dUc(3,CC)
*      - 0.25*(dFc(3)*dy13 - dGc(3)*dx13)
dU(3,N3) = dU(3,N3) + coef*dUc(3,CC)
*      - 0.25*(dFc(3)*dy21 - dGc(3)*dx21)

dU(4,N1) = dU(4,N1) + coef*dUc(4,CC)
*      - 0.25*(dFc(4)*dy32 - dGc(4)*dx32)
dU(4,N2) = dU(4,N2) + coef*dUc(4,CC)
*      - 0.25*(dFc(4)*dy13 - dGc(4)*dx13)
dU(4,N3) = dU(4,N3) + coef*dUc(4,CC)
*      - 0.25*(dFc(4)*dy21 - dGc(4)*dx21)

enddo
enddo

EF2 = 0
do col = 1, E2colormax
  EF1 = EF2 + 1
  EF2 = EF1 - 1 + Ncolor(col)
  do EF = EF1, EF2
    FF = eface(EF)
    CC = face(FF,1)
    N1 = face(FF,3)
    N2 = face(FF,4)

    dy21 = y(N2) - y(N1)
    dx21 = x(N2) - x(N1)

    uu = Uc(2,CC)/Uc(1,CC)
    vv = Uc(3,CC)/Uc(1,CC)
    ww2 = 0.5*(uu*uu + vv*vv)

    dp = gam1*(dUc(4,CC) - uu*dUc(2,CC) - vv*dUc(3,CC)
*      + ww2*dUc(1,CC))

    dFc(2) = dp
    dGc(3) = dp

    dU(2,N1) = dU(2,N1) - 0.25*dFc(2)*dy21
    dU(2,N2) = dU(2,N2) - 0.25*dFc(2)*dy21

    dU(3,N1) = dU(3,N1) + 0.25*dGc(3)*dx21
    dU(3,N2) = dU(3,N2) + 0.25*dGc(3)*dx21

  enddo
enddo

do NN = 1, Nmax
  dU(1,NN) = dU(1,NN)*deltN(NN)
  dU(2,NN) = dU(2,NN)*deltN(NN)
  dU(3,NN) = dU(3,NN)*deltN(NN)
  dU(4,NN) = dU(4,NN)*deltN(NN)

```

```

enddo

return
end

```

```

C*****
C*
C*      subroutine to calculate forth difference smoothing
C*
C*****

      subroutine smooth
      implicit none

      include 'TRI.INC'
      integer N1, N2, N3          !nodes at corners of cell
      integer NN, CC1, CC2, col, CC !pointers
      integer EF, EF1, EF2
      integer FF, FF1, FF2, PN
      integer P1, P2              !periodic nodes
      real dx31, dx12, dx23       !change in x
      real dy31, dy12, dy23       !change in y
      real dxC1, dxC2, dxC3, dxC4 !change in x in cell
      real dyC1, dyC2, dyC3, dyC4 !change in y in cell
      real delN(4,Maxnodes)       !gradient at node
      real coef, coef1, coef2      !coefficient

      do NN = 1, Nmax
         delN(1,NN) = 0.
         delN(2,NN) = 0.
         delN(3,NN) = 0.
         delN(4,NN) = 0.
      enddo

      CC2 = 0

      do col = 1, Ccolormax
         CC1 = CC2 + 1
         CC2 = CC1 - 1 + NCcolor(col)

CVD$      NODEPCHK
         do CC = CC1, CC2
            N1 = cell(CC,4)
            N2 = cell(CC,5)
            N3 = cell(CC,6)

            dx31 = x(N3) - x(N1)
            dx12 = x(N1) - x(N2)
            dx23 = x(N2) - x(N3)
            dy31 = y(N3) - y(N1)
            dy12 = y(N1) - y(N2)
            dy23 = y(N2) - y(N3)

            coef = ((max(0.,min(1.,(1.+10.*vol(CC)))-1.)*eps1coef +1.)
&               *(0.5/areaC(CC))

            dxC1 = (U(1,N1)*dy23 + U(1,N2)*dy31 + U(1,N3)*dy12)
&               *coef
            dxC2 = (U(2,N1)*dy23 + U(2,N2)*dy31 + U(2,N3)*dy12)
&               *coef
            dxC3 = (U(3,N1)*dy23 + U(3,N2)*dy31 + U(3,N3)*dy12)
&               *coef
            dxC4 = (U(4,N1)*dy23 + U(4,N2)*dy31 + U(4,N3)*dy12)
&               *coef

            dyC1 = (U(1,N1)*dx23 + U(1,N2)*dx31 + U(1,N3)*dx12)
&               *coef
            dyC2 = (U(2,N1)*dx23 + U(2,N2)*dx31 + U(2,N3)*dx12)

```

```

*      *coef
dyC3 = (U(3,N1)*dx23 + U(3,N2)*dx31 + U(3,N3)*dx12)
*      *coef
*      dyC4 = (U(4,N1)*dx23 + U(4,N2)*dx31 + U(4,N3)*dx12)
*      *coef

delN(1,N1) = delN(1,N1) + (dxC1*dy23 + dyC1*dx23)
delN(1,N2) = delN(1,N2) + (dxC1*dy31 + dyC1*dx31)
delN(1,N3) = delN(1,N3) + (dxC1*dy12 + dyC1*dx12)

delN(2,N1) = delN(2,N1) + (dxC2*dy23 + dyC2*dx23)
delN(2,N2) = delN(2,N2) + (dxC2*dy31 + dyC2*dx31)
delN(2,N3) = delN(2,N3) + (dxC2*dy12 + dyC2*dx12)

delN(3,N1) = delN(3,N1) + (dxC3*dy23 + dyC3*dx23)
delN(3,N2) = delN(3,N2) + (dxC3*dy31 + dyC3*dx31)
delN(3,N3) = delN(3,N3) + (dxC3*dy12 + dyC3*dx12)

delN(4,N1) = delN(4,N1) + (dxC4*dy23 + dyC4*dx23)
delN(4,N2) = delN(4,N2) + (dxC4*dy31 + dyC4*dx31)
delN(4,N3) = delN(4,N3) + (dxC4*dy12 + dyC4*dx12)
enddo
enddo

c* upper and lower boundary
EF2 = 0

do col = 1, Eicolormax
  EF1 = EF2 + 1
  EF2 = EF1 - 1 + NEcolor(col)
enddo

CVD$ NODEPCHK
do EF = EF1, EF2
  FF = eface(EF)
  CC = face(FF,1)
  N1 = face(FF,3)
  N2 = face(FF,4)
  N3 = face(FF,6)

  dx31 = x(N3) - x(N1)
  dx12 = x(N1) - x(N2)
  dx23 = x(N2) - x(N3)
  dy31 = y(N3) - y(N1)
  dy12 = y(N1) - y(N2)
  dy23 = y(N2) - y(N3)

  coef = ((max(0.,min(1.,(1.+10.*vol(CC))))-1.)*eps1coef +1.)
*      *(0.5/areaC(CC))

  dxC1 = (U(1,N1)*dy23 + U(1,N2)*dy31 + U(1,N3)*dy12)
*      *coef
  dxC2 = (U(2,N1)*dy23 + U(2,N2)*dy31 + U(2,N3)*dy12)
*      *coef
  dxC3 = (U(3,N1)*dy23 + U(3,N2)*dy31 + U(3,N3)*dy12)
*      *coef
  dxC4 = (U(4,N1)*dy23 + U(4,N2)*dy31 + U(4,N3)*dy12)
*      *coef

  dyC1 = (U(1,N1)*dx23 + U(1,N2)*dx31 + U(1,N3)*dx12)
*      *coef
  dyC2 = (U(2,N1)*dx23 + U(2,N2)*dx31 + U(2,N3)*dx12)
*      *coef
  dyC3 = (U(3,N1)*dx23 + U(3,N2)*dx31 + U(3,N3)*dx12)
*      *coef
  dyC4 = (U(4,N1)*dx23 + U(4,N2)*dx31 + U(4,N3)*dx12)
*      *coef

  delN(1,N1) = delN(1,N1) + (dxC1*dy12 + dyC1*dx12)
  delN(1,N2) = delN(1,N2) + (dxC1*dy12 + dyC1*dx12)

  delN(2,N1) = delN(2,N1) + (dxC2*dy12 + dyC2*dx12)
  delN(2,N2) = delN(2,N2) + (dxC2*dy12 + dyC2*dx12)

```

```

        delN(3,N1) = delN(3,N1) + (dxC3*dy12 + dyC3*dx12)
        delN(3,N2) = delN(3,N2) + (dxC3*dy12 + dyC3*dx12)

        delN(4,N1) = delN(4,N1) + (dxC4*dy12 + dyC4*dx12)
        delN(4,N2) = delN(4,N2) + (dxC4*dy12 + dyC4*dx12)

    enddo
enddo

CVD$ NODEPCHK
do PN = 1, Pmax
    P1 = pnode(PN,1)
    P2 = pnode(PN,2)
    delN(1,P1) = delN(1,P1) + delN(1,P2)
    delN(2,P1) = delN(2,P1) + delN(2,P2)
    delN(3,P1) = delN(3,P1) + delN(3,P2)
    delN(4,P1) = delN(4,P1) + delN(4,P2)
    delN(1,P2) = delN(1,P1)
    delN(2,P2) = delN(2,P1)
    delN(3,P2) = delN(3,P1)
    delN(4,P2) = delN(4,P1)
enddo

FF2 = 0

do col = 1, Fcolormax
    FF1 = FF2 + 1
    FF2 = FF1 - 1 + NFcolor(col)
enddo

CVD$ NODEPCHK
do FF = FF1, FF2
    N1 = face(FF,3)
    N2 = face(FF,4)

    coef1 = 0.5*eps2*(1./deltN(N1) + 1./deltN(N2))*deltN(N1)
    coef2 = 0.5*eps2*(1./deltN(N1) + 1./deltN(N2))*deltN(N2)

    dU(1,N1) = dU(1,N1) + coef1*(delN(1,N2) - delN(1,N1))
    dU(1,N2) = dU(1,N2) + coef2*(delN(1,N1) - delN(1,N2))

    dU(2,N1) = dU(2,N1) + coef1*(delN(2,N2) - delN(2,N1))
    dU(2,N2) = dU(2,N2) + coef2*(delN(2,N1) - delN(2,N2))

    dU(3,N1) = dU(3,N1) + coef1*(delN(3,N2) - delN(3,N1))
    dU(3,N2) = dU(3,N2) + coef2*(delN(3,N1) - delN(3,N2))

    dU(4,N1) = dU(4,N1) + coef1*(delN(4,N2) - delN(4,N1))
    dU(4,N2) = dU(4,N2) + coef2*(delN(4,N1) - delN(4,N2))
enddo
enddo

return
end

c*****
c*
c*      this subroutine accounts for periodic nodes
c*
c*****

subroutine bperiodic
implicit none

include 'TRI.INC'
integer PN          !pointer
integer P1, P2      !periodic nodes

```

```

CVD$  NODEPCHK
      do PN = 1, Pmax
        P1 = pnode(PN,1)
        P2 = pnode(PN,2)
        dU(1,P1) = dU(1,P1) + dU(1,P2)
        dU(2,P1) = dU(2,P1) + dU(2,P2)
        dU(3,P1) = dU(3,P1) + dU(3,P2)
        dU(4,P1) = dU(4,P1) + dU(4,P2)
        dU(1,P2) = dU(1,P1)
        dU(2,P2) = dU(2,P1)
        dU(3,P2) = dU(3,P1)
        dU(4,P2) = dU(4,P1)
      enddo

      return
      end

c*****
c*
c*      adjust inlet state vector for boundary condition
c*
c*****

      subroutine binlet
      implicit none

      include 'TRI.INC'
      real rrinl, uuinl, vvinnl, ppinnl !average values at inlet
      real vw2innl, aainl
      real rOinf, aOinf                !stag. density and speed of sound
      real HOpres                       !stagnation enthalpy and pressure
      real spres                        !entropy
      real coef
      real aaa1, aaa2, aaa3, aaa4      !coef. of inverse of matrix
      real bbb1, bbb2, bbb3, bbb4
      real ccc1, ccc2, ccc3, ccc4
      real ddd1, ddd2, ddd3, ddd4
      real vw2, uu, vv, pp, rr, aa    !values at next interior node
      real dHO, ds, dtan, dw4
      real drr, duu, dvv, dpp
      integer IN, NN                  !boundary face and its nodes

c** average values at inlet
      rrinl = U(1,innode(Imax/2))
      uuinl = U(2,innode(Imax/2))/rrinl
      vvinnl = U(3,innode(Imax/2))/rrinl
      vw2innl = uuinl**2 + vvinnl**2
      ppinnl = gam1*(U(4,innode(Imax/2))
      &      - 0.5*rrinl*vw2innl)
      aainl = sqrt(gam*ppinnl/rrinl)

c* subsonic inlet nodes
      if ((sqrt(vw2innl)/aainl).lt.1. .or. pitch.ne.0.) then
c** prescribed values
          HOpres = 1./gam1
          spres = 0.

c** coefficient of matrix
          coef = 1./(uuinnl*(aainl+uuinnl) + vvinnl**2)
          aaa1 = (rrinl+uuinnl/aainl)*coef
          aaa2 = -(ppinnl/aainl)*(uuinnl*gam/gam1 + vw2innl/aainl)*coef
          aaa3 = -(rrinl*vvinnl/aainl)*coef
          aaa4 = (vw2innl/aainl**2)*coef
          bbb1 = uuinl*coef
          bbb2 = -(uuinnl*ppinnl/(gam1*rrinl))*coef
          bbb3 = -vvinnl*coef
          bbb4 = -(uuinnl/rrinl)*coef
          ccc1 = vvinnl*coef

```

```

ccc2 = -(vv1nl*pp1nl/(gam1*rr1nl))*coef
ccc3 = (aa1nl + uu1nl)*coef
ccc4 = -(vv1nl/rr1nl)*coef
ddd1 = rr1nl*aa1nl*uu1nl*coef
ddd2 = -(pp1nl*aa1nl*uu1nl/gam1)*coef
ddd3 = -rr1nl*aa1nl*vv1nl*coef
ddd4 = vv21nl*coef

CVD$  NODEPCHK
      do IN = 1, Imax
        NN = innode(IN)
        rr = U(1,NN)
        uu = U(2,NN)/rr
        vv = U(3,NN)/rr
        ww2 = uu**2 + vv**2
        pp = gam1*(U(4,NN) - 0.5*rr*ww2)
        aa = sqrt(gam*pp/rr)
        dHO = HOpres - ((gam/gam1)*pp/rr + 0.5*ww2)
        ds = spres - (log(gam*pp) - gam*log(rr))
        dtan = (S1nl - vv/uu)*uu1nl**2
        dw4 = dU(1,NN)*(aa*uu + gam1*ww2)
        &      - dU(2,NN)*(aa + gam1*uu)
        &      - dU(3,NN)*gam1*vv
        &      + dU(4,NN)*gam1
        drr = aaa1*dHO + aaa2*ds + aaa3*dtan + aaa4*dw4
        duu = bbb1*dHO + bbb2*ds + bbb3*dtan + bbb4*dw4
        dvv = ccc1*dHO + ccc2*ds + ccc3*dtan + ccc4*dw4
        dpp = ddd1*dHO + ddd2*ds + ddd3*dtan + ddd4*dw4

        dU(1,NN) = drr
        dU(2,NN) = rr*duu + uu*drr
        dU(3,NN) = rr*dvv + vv*drr
        dU(4,NN) = dpp/gam1 + 0.5*ww2*drr + rr*(uu*duu + vv*dvv)
      enddo

c*  supersonic inlet nodes
      else if ((sqrt(ww21nl)/aa1nl).ge.1.) then
c*  state vector components far from body
        r0inf = 1.
        a0inf = 1.

CVD$  NODEPCHK
      do IN = 1, Imax
        NN = innode(IN)
        rr = r0inf*(1.0+0.5*gam1*Minl**2)**(-1./gam1)
        aa = a0inf*(1.0+0.5*gam1*Minl**2)**(-0.5)
        uu = Minl*aa
        pp = rr*aa**2/gam1

        dU(1,NN) = rr - U(1,NN)
        dU(2,NN) = rr*uu - U(2,NN)
        dU(3,NN) = - U(3,NN)
        dU(4,NN) = pp/gam1 + .5*(uu**2)*rr - U(4,NN)
      enddo
    endif

    return
  end

C*****
c*
c*      adjust outlet state vector for boundary condition
c*
C*****

      subroutine boutlet
      implicit none

      include 'TRI.INC'

```

```

real rrout, uuout, vvout, ppout !average values at outlet
real aaout, ww2out
real aaa1, aaa4           !coef. of inverse of matrix
real bbb3, bbb4
real ccc2
real ddd4
real ww2, uu, vv, pp, rr, aa !values at next interior node
real dw1, dw2, dw3, dp
real drr, duu, dvv, dpp
integer ON, NN           !boundary face and its nodes

rrout = U(1,outnode(Onmax/2))
uuout = U(2,outnode(Onmax/2))/rrout
vvout = U(3,outnode(Onmax/2))/rrout
ww2out = uuout**2 + vvout**2
ppout = gam1*(U(4,outnode(Onmax/2))
      - 0.5*rrout*ww2out)
*
aaout = sqrt(gam*ppout/rrout)

aaa1 = -1./aaout**2
aaa4 = 1./aaout**2
bbb3 = .1/(rrout*aaout)
bbb4 = -1./(rrout*aaout)
ccc2 = .1/(rrout*aaout)
ddd4 = 1.

c* set boundary values of state vector for nodes
CVD$ NODEPCHK
do ON = 1, Onmax
  NN = outnode(ON)
  rr = U(1,NN)
  uu = U(2,NN)/rr
  vv = U(3,NN)/rr
  ww2 = uu**2 + vv**2
  pp = gam1*(U(4,NN) - 0.5*rr*ww2)
  aa = sqrt(gam*pp/rr)
  if (sqrt(ww2) .lt. aa) then
    dw1 = dU(1,NN)*(0.5*ww2*gam1 - aa**2)
    *
    *
    *
    dw2 = - dU(1,NN)*aa*vv
    *
    *
    dw3 = dU(1,NN)*(0.5*ww2*gam1 - aa*uu)
    *
    *
    *
    dp = pout - pp
    drr = aaa1*dw1 + aaa4*dp
    duu = bbb3*dw3 + bbb4*dp
    dvv = ccc2*dw2
    dpp = ddd4*dp
    dU(1,NN) = drr
    dU(2,NN) = rr*duu + uu*drr
    dU(3,NN) = rr*dvv + vv*drr
    dU(4,NN) = dpp/gam1 + 0.5*ww2*drr + rr*(uu*duu + vv*dvv)

  endif
enddo

return
end

c*****
c*
c*   this subroutine changes the momentum change to make
c*   flow tangent to wall
c*
c*

```

```

c*****
      subroutine tangent
      implicit none

      include 'TRI.INC'
      integer EN                      !pointer
      integer NN                      !node on wall
      real drwn                       !change in momentum normal to wall

CVD$  NODEPCHK
      do EN = 1, bnode
         NN = enode(EN)
         drwn = -(U(2,NN) + dU(2,NN))*senode(EN) +
&             (U(3,NN) + dU(3,NN))*cenode(EN)
         dU(2,NN) = dU(2,NN) + drwn*senode(EN)
         dU(3,NN) = dU(3,NN) - drwn*cenode(EN)
      enddo

      return
      end

```

A.1.4 Jameson Scheme

```

c*****
c*
c*      main program for triangular Jameson scheme
c*
c*****

      program triangle
      implicit none

      include 'TRI.INC'
      integer Niter                   !number of iterations
      real maxchange                  !max change in state vector
      integer CC                      !pointer
      integer maxnode, maxeqn         !where max change occurs

c*  read in data from file
      call gridio(1)
      call flowio(1)
      call input

      Niter = 0
      durms = 999.

c*  start L-story file from the top
      open(unit=35,status='unknown',form='formatted')
      write(35,2) Minl
      close(unit=35)
2    format(' inlet Mach number = ',f5.3)

c*  loop until converged
      do while ((Niter.lt.Maxiter) .and. (durms.gt.2.e-7))
         Niter = Niter + 1

         call update(maxchange, maxnode, maxeqn)

         if (mod(Niter,10).eq.0 .or. Niter.lt.10) then
            call flowio(0)
            open(unit=50, status='unknown', form='unformatted')
            write(50) Cmax, (vol(CC),CC=1,Cmax)
            close(unit=50)

10        open(unit=35, status='old',access='append',err=10)
            write(35,1) Niter, durms, maxchange, x(maxnode),

```



```

&          y(maxnode), maxeqn
      close(unit=35)
      write(6,1) Niter, durms, maxchange, x(maxnode),
&          y(maxnode), maxeqn
      endif
1      format('Niter=',i4,' rms=',f9.7,' max=',f9.7,' x=',f6.3,
&          ' y=',f6.3,' eqn=',i1)

      enddo

c* write out data to file
      call flowio(0)

      stop
      end

C*****
c*
c*      update state vectors at next time step
c*
c*
C*****

      subroutine update(maxchange, maxnode, maxeqn)

      include 'TRI.INC'
      integer NN, i
      real UO(4,Maxnodes)
      real alpha1, alpha2, alpha3
      real alpha4
      real maxchange
      integer maxnode, maxeqn

      !pointer
      !starting values of state vector
      !coefficients
      !max change in state vector
      !where max change occurs

      alpha1 = 0.25
      alpha2 = 1./3.
      alpha3 = 0.5
      alpha4 = 1.

c* find timestep at each node
      call timestep

c* advance to next time step in four steps

c* step one
      call calcflux
      call dissipation

      do NN = 1, Nmax
          UO(1,NN) = U(1,NN)
          UO(2,NN) = U(2,NN)
          UO(3,NN) = U(3,NN)
          UO(4,NN) = U(4,NN)
          dU(1,NN) = U(1,NN)
          dU(2,NN) = U(2,NN)
          dU(3,NN) = U(3,NN)
          dU(4,NN) = U(4,NN)
          U(1,NN) = UO(1,NN) - (alpha1*deltN(NN))
&          *(flux(1,NN) - dis(1,NN))
          U(2,NN) = UO(2,NN) - (alpha1*deltN(NN))
&          *(flux(2,NN) - dis(2,NN))
          U(3,NN) = UO(3,NN) - (alpha1*deltN(NN))
&          *(flux(3,NN) - dis(3,NN))
          U(4,NN) = UO(4,NN) - (alpha1*deltN(NN))
&          *(flux(4,NN) - dis(4,NN))
          dU(1,NN) = U(1,NN) - dU(1,NN)
          dU(2,NN) = U(2,NN) - dU(2,NN)

```

```

        dU(3,NN) = U(3,NN) - dU(3,NN)
        dU(4,NN) = U(4,NN) - dU(4,NN)
    enddo

    call binlet
    call boutlet
    call tangent

c* step two
    call calcflux
    call dissipation

    do NN = 1, Nmax
        dU(1,NN) = U(1,NN)
        dU(2,NN) = U(2,NN)
        dU(3,NN) = U(3,NN)
        dU(4,NN) = U(4,NN)
        U(1,NN) = UO(1,NN) - (alpha2*deltN(NN))
        &      *(flux(1,NN) - dis(1,NN))
        &      U(2,NN) = UO(2,NN) - (alpha2*deltN(NN))
        &      *(flux(2,NN) - dis(2,NN))
        &      U(3,NN) = UO(3,NN) - (alpha2*deltN(NN))
        &      *(flux(3,NN) - dis(3,NN))
        &      U(4,NN) = UO(4,NN) - (alpha2*deltN(NN))
        &      *(flux(4,NN) - dis(4,NN))
        dU(1,NN) = U(1,NN) - dU(1,NN)
        dU(2,NN) = U(2,NN) - dU(2,NN)
        dU(3,NN) = U(3,NN) - dU(3,NN)
        dU(4,NN) = U(4,NN) - dU(4,NN)
    enddo

    call binlet
    call boutlet
    call tangent

c* step three
    call calcflux

    do NN = 1, Nmax
        dU(1,NN) = U(1,NN)
        dU(2,NN) = U(2,NN)
        dU(3,NN) = U(3,NN)
        dU(4,NN) = U(4,NN)
        U(1,NN) = UO(1,NN) - (alpha3*deltN(NN))
        &      *(flux(1,NN) - dis(1,NN))
        &      U(2,NN) = UO(2,NN) - (alpha3*deltN(NN))
        &      *(flux(2,NN) - dis(2,NN))
        &      U(3,NN) = UO(3,NN) - (alpha3*deltN(NN))
        &      *(flux(3,NN) - dis(3,NN))
        &      U(4,NN) = UO(4,NN) - (alpha3*deltN(NN))
        &      *(flux(4,NN) - dis(4,NN))
        dU(1,NN) = U(1,NN) - dU(1,NN)
        dU(2,NN) = U(2,NN) - dU(2,NN)
        dU(3,NN) = U(3,NN) - dU(3,NN)
        dU(4,NN) = U(4,NN) - dU(4,NN)
    enddo

    call binlet
    call boutlet
    call tangent

c* step four
    call calcflux

    do NN = 1, Nmax
        dU(1,NN) = U(1,NN)
        dU(2,NN) = U(2,NN)
        dU(3,NN) = U(3,NN)
        dU(4,NN) = U(4,NN)
        U(1,NN) = UO(1,NN) - (alpha4*deltN(NN))
        &      *(flux(1,NN) - dis(1,NN))
        &      U(2,NN) = UO(2,NN) - (alpha4*deltN(NN))

```

```

      &      U(3,NN) = UO(3,NN) - (flux(2,NN) - dis(2,NN))
      &      U(4,NN) = UO(4,NN) - (flux(3,NN) - dis(3,NN))
      &      dU(1,NN) = U(1,NN) - dU(1,NN)
      &      dU(2,NN) = U(2,NN) - dU(2,NN)
      &      dU(3,NN) = U(3,NN) - dU(3,NN)
      &      dU(4,NN) = U(4,NN) - dU(4,NN)
    enddo

    call binlet
    call boutlet
    call tangent

c* find root mean square difference in state vector
    durms = 0.0
    maxchange = 0.

    do i = 1, 4
      do NN = 1, Nmax
        durms = durms + (U(i,NN) - UO(i,NN))**2
        if (abs(U(i,NN)-UO(i,NN)).gt.abs(maxchange)) then
          maxchange = U(i,NN) - UO(i,NN)
          maxnode = NN
          maxeqn = i
        endif
      enddo
    enddo

    durms = sqrt(durms/(4.*Nmax))

    return
    end

c*****
c*
c* calculate time step for nodes
c*
c*****

      subroutine timestep
      implicit none

      include 'TRI.INC'
      integer EF, FF, NN          !pointer
      integer CC, CC1, CC2
      integer N1, N2, N3, N4      !nodes related to face
      integer PN, P1, P2          !periodic nodes
      real delx, dely             !x and y length of side
      real delside                !length of side
      real uu, vv, aa, rr, ww2    !values at node
      integer FF1, FF2, col        !color pointers
      integer EF1, EF2
      real onethird

c* set values for false node 0
      U(1,0) = 1.
      U(2,0) = 0.
      U(3,0) = 0.
      U(4,0) = 0.

c* zero out delt
      do NN = 1, Nmax
        deltN(NN) = 0.
      enddo

c* find time step for each node

```

```

FF2 = 0

do col = 1, Fcolormax
  FF1 = FF2 + 1
  FF2 = FF1 - 1 + NFcolor(col)
  do FF = FF1, FF2
    N1 = face(FF,3)
    N2 = face(FF,4)
    N3 = face(FF,5)
    N4 = face(FF,6)

    delx = x(N2) - x(N1)
    dely = y(N2) - y(N1)
    delside = sqrt(delx**2 + dely**2)

    rr = U(1,N3)
    uu = U(2,N3)/rr
    vv = U(3,N3)/rr
    ww2 = uu**2 + vv**2
    aa = sqrt(gam*gam1*(U(4,N3)/rr - 0.5*ww2))

    deltn(N3) = deltn(N3) + abs(uu*dely-vv*delx) + aa*delside

    rr = U(1,N4)
    uu = U(2,N4)/rr
    vv = U(3,N4)/rr
    ww2 = uu**2 + vv**2
    aa = sqrt(gam*gam1*(U(4,N4)/rr - 0.5*ww2))

    deltn(N4) = deltn(N4) + abs(uu*dely-vv*delx) + aa*delside

  enddo
enddo

c* add to timestep for boundary nodes
EF2 = 0

do col = 1, E2colormax
  EF1 = EF2 + 1
  EF2 = EF1 - 1 + NEcolor(col)

  do EF = EF1, EF2
    FF = eface(EF)
    N1 = face(FF,3)
    N2 = face(FF,4)

    delx = x(N2) - x(N1)
    dely = y(N2) - y(N1)
    delside = sqrt(delx**2 + dely**2)

    rr = U(1,N2)
    uu = U(2,N2)/rr
    vv = U(3,N2)/rr
    ww2 = uu**2 + vv**2
    aa = sqrt(gam*gam1*(U(4,N2)/rr - 0.5*ww2))

    deltn(N2) = deltn(N2) + abs(uu*dely-vv*delx) + aa*delside

    rr = U(1,N1)
    uu = U(2,N1)/rr
    vv = U(3,N1)/rr
    ww2 = uu**2 + vv**2
    aa = sqrt(gam*gam1*(U(4,N1)/rr - 0.5*ww2))

    deltn(N1) = deltn(N1) + abs(uu*dely-vv*delx) + aa*delside

  enddo
enddo

c** find time step at periodic nodes
do PH = 1, Pmax
  P1 = pnode(PH,1)

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```

        P2 = pnode(PH,2)
        deltn(P1) = deltn(P1) + deltn(P2)
        deltn(P2) = deltn(P1)
    enddo

c** deltn is actually deltn/areaH
    do NN = 1, Nmax
        deltn(NN) = CFL*2./deltn(NN)
    enddo

    return
end

c*****
c*
c*      calculate flux vector values at nodes
c*
c*****

      subroutine calcflux
      implicit none

      include 'TRI.INC'
      integer CC, CC1, CC2
      real dx23, dx31, dx12           !change in x on cell edges
      real dy23, dy31, dy12           !change in y on cell edges
      integer EF, NN, FF, i           !pointers
      real delx, dely                 !change in x and y on face
      real WW                         !kinetic energy
      integer N1, N2, N3, N4           !nodes around edge
      real dflux                      !fluxes through face
      integer FF1, FF2, col           !color pointers

      do NN = 1, Nmax
c* set flux to zero
        flux(1,NN) = 0.
        flux(2,NN) = 0.
        flux(3,NN) = 0.
        flux(4,NN) = 0.

c* calculate f and g at nodes
        WW = 0.5*(U(2,NN)**2 + U(3,NN)**2)/U(1,NN)
        F(1,NN) = U(2,NN)
        F(2,NN) = U(2,NN)**2/U(1,NN) + gam1*(U(4,NN) - WW)
        F(3,NN) = U(2,NN)*U(3,NN)/U(1,NN)
        F(4,NN) = (U(2,NN)/U(1,NN))*(gam*U(4,NN) - gam1*WW)

        G(1,NN) = U(3,NN)
        G(2,NN) = U(2,NN)*U(3,NN)/U(1,NN)
        G(3,NN) = U(3,NN)**2/U(1,NN) + gam1*(U(4,NN) - WW)
        G(4,NN) = (U(3,NN)/U(1,NN))*(gam*U(4,NN) - gam1*WW)
      enddo

      do CC = 1, Cmax
        N1 = cell(CC,4)
        N2 = cell(CC,5)
        N3 = cell(CC,6)

        dy23 = y(N2) - y(N3)
        dy31 = y(N3) - y(N1)
        dy12 = y(N1) - y(N2)
        dx23 = x(N2) - x(N3)
        dx31 = x(N3) - x(N1)
        dx12 = x(N1) - x(N2)

        dUc(1,CC) = 0.5*(-F(1,N1)*dy23 + G(1,N1)*dx23
&                -F(1,N2)*dy31 + G(1,N2)*dx31

```

```

&          -F(1,N3)*dy12 + G(1,N3)*dx12)
      dUc(2,CC) = 0.5*(-F(2,N1)*dy23 + G(2,N1)*dx23
&          -F(2,N2)*dy31 + G(2,N2)*dx31
&          -F(2,N3)*dy12 + G(2,N3)*dx12)
      dUc(3,CC) = 0.5*(-F(3,N1)*dy23 + G(3,N1)*dx23
&          -F(3,N2)*dy31 + G(3,N2)*dx31
&          -F(3,N3)*dy12 + G(3,N3)*dx12)
      dUc(4,CC) = 0.5*(-F(4,N1)*dy23 + G(4,N1)*dx23
&          -F(4,N2)*dy31 + G(4,N2)*dx31
&          -F(4,N3)*dy12 + G(4,N3)*dx12)
    enddo

c* implement wall boundary conditions
    call bwall

    CC2 = 0

    do col = 1, Ccolormax
      CC1 = CC2 + 1
      CC2 = CC1 - 1 + NCcolor(col)

CVD$ NODEPCHK
      do CC = CC1, CC2
        N1 = cell(CC,4)
        N2 = cell(CC,5)
        N3 = cell(CC,6)

        flux(1,N1) = flux(1,N1) - dUc(1,CC)
        flux(1,N2) = flux(1,N2) - dUc(1,CC)
        flux(1,N3) = flux(1,N3) - dUc(1,CC)

        flux(2,N1) = flux(2,N1) - dUc(2,CC)
        flux(2,N2) = flux(2,N2) - dUc(2,CC)
        flux(2,N3) = flux(2,N3) - dUc(2,CC)

        flux(3,N1) = flux(3,N1) - dUc(3,CC)
        flux(3,N2) = flux(3,N2) - dUc(3,CC)
        flux(3,N3) = flux(3,N3) - dUc(3,CC)

        flux(4,N1) = flux(4,N1) - dUc(4,CC)
        flux(4,N2) = flux(4,N2) - dUc(4,CC)
        flux(4,N3) = flux(4,N3) - dUc(4,CC)

      enddo
    enddo

    return
end

c*****
c*
c*      this subroutine applies the wall boundary condition
c*
c*****

      subroutine bwall
      implicit none

      include 'TRI.INC'
      integer EF, FF, CC          !pointer
      integer N1, N3              !nodes on edge
      real coef
      real dy13, dx13             !change in x and y
      real p1, p3                 !pressure
      integer EF1, EF2, col

```

```

c* upper and lower boundary
  EF2 = 0

  do col = 1, Elcolormax
    EF1 = EF2 + 1
    EF2 = EF1 - 1 + NEcolor(col)

CVD$ NODEPCHK
  do EF = EF1, EF2
    FF = eface(EF)
    CC = face(FF,1)
    N1 = face(FF,4)
    N3 = face(FF,3)

    dy13 = y(N1) - y(N3)
    dx13 = x(N1) - x(N3)

    p1 = gam1*(U(4,N1) - 0.5*(U(2,N1)**2 + U(3,N1)**2)/
      & U(1,N1))
    & p3 = gam1*(U(4,N3) - 0.5*(U(2,N3)**2 + U(3,N3)**2)/
      & U(1,N3))

    dUc(1,CC) = dUc(1,CC) + 0.5*(
      & (F(1,N3) + F(1,N1))*dy13 -
      & (G(1,N3) + G(1,N1))*dx13)

    dUc(2,CC) = dUc(2,CC) + 0.5*(
      & (F(2,N3) + F(2,N1) - p1 - p3)*dy13 -
      & (G(2,N3) + G(2,N1))*dx13)

    dUc(3,CC) = dUc(3,CC) + 0.5*(
      & (F(3,N3) + F(3,N1))*dy13 -
      & (G(3,N3) + G(3,N1) - p1 - p3)*dx13)

    dUc(4,CC) = dUc(4,CC) + 0.5*(
      & (F(4,N3) + F(4,N1))*dy13 -
      & (G(4,N3) + G(4,N1))*dx13)

    enddo
  enddo

  return
end

C*****
C*
C* calculate the dissipation at each of the nodes for
C* the current values of the state vector
C*
C*****

subroutine dissipation
implicit none

include 'TRI.INC'
integer N1, N2, N3
integer PN, P1, P2
real change
integer NN, FF, 1
real dx31, dx12, dx23
real dy31, dy12, dy23
real dxC1, dxC2, dxC3, dxC4
real dyC1, dyC2, dyC3, dyC4
real del2(4,Maxnodes)
real pres
real delp
real eps1(Maxnodes)

!nodes at end of face
!periodic nodes
!change in dissipation
!pointers
!change in x
!change in y
!change in x in cell
!change in y in cell
!second order changes
!pressure at nodes
!change in pressure
!dissipation coefficients

```

```

real coef, coef2           !combination of eps1
integer FF1, FF2, col      !color pointers
integer CC1, CC2, CC       !pointers
integer EF, EF1, EF2

do NN = 1, Nmax
  dis(1,NN) = 0.
  dis(2,NN) = 0.
  dis(3,NN) = 0.
  dis(4,NN) = 0.

  del2(1,NN) = 0.
  del2(2,NN) = 0.
  del2(3,NN) = 0.
  del2(4,NN) = 0.

  eps1(NN) = 0.
enddo

c* undivided Laplacian of pressure for eps1 coefficient
FF2 = 0

do col = 1, Fcolormax
  FF1 = FF2 + 1
  FF2 = FF1 - 1 + NFcolor(col)

CVD$ NODEPCHK
  do FF = FF1, FF2
    N1 = face(FF,3)
    N2 = face(FF,4)

    delp = gam1*(U(4,N1) - U(4,N2)
      & + .5*(U(2,N2)**2 + U(3,N2)**2)/U(1,N2)
      & - .5*(U(2,N1)**2 + U(3,N1)**2)/U(1,N1))
    eps1(N1) = eps1(N1) - epsicoef*delp
    eps1(N2) = eps1(N2) + epsicoef*delp

  enddo
enddo

c** find eps1 and del2 at periodic nodes
do PN = 1, Pmax
  P1 = pnode(PN,1)
  P2 = pnode(PN,2)
  eps1(P1) = eps1(P1) + eps1(P2)
  eps1(P2) = eps1(P1)
enddo

do NN = 1, Nmax
  pres = gam1*(U(4,NN) - 0.5*(U(2,NN)**2 +
    & U(3,NN)**2)/U(1,NN))
  eps1(NN) = abs(eps1(NN)/pres)
enddo

c** low-accuracy smoothing
if (sigE .eq. 0.) then

  FF2 = 0

  do col = 1, Fcolormax
    FF1 = FF2 + 1
    FF2 = FF1 - 1 + NFcolor(col)

CVD$ NODEPCHK
  do FF = FF1, FF2
    N1 = face(FF,3)
    N2 = face(FF,4)

    del2(1,N1) = del2(1,N1) - U(1,N2) + U(1,N1)
    del2(1,N2) = del2(1,N2) - U(1,N1) + U(1,N2)

    del2(2,N1) = del2(2,N1) - U(2,N2) + U(2,N1)

```



```

        del2(2,N2) = del2(2,N2) - U(2,N1) + U(2,N2)

        del2(3,N1) = del2(3,N1) - U(3,N2) + U(3,N1)
        del2(3,N2) = del2(3,N2) - U(3,N1) + U(3,N2)

        del2(4,N1) = del2(4,N1) - U(4,N2) + U(4,N1)
        del2(4,N2) = del2(4,N2) - U(4,N1) + U(4,N2)
    enddo
enddo

c** high-accuracy smoothing
    else if (sigE .eq. 1.) then

        CC2 = 0

        do col = 1, Ccolormax
            CC1 = CC2 + 1
            CC2 = CC1 - 1 + NCcolor(col)
        enddo

CVD$ NODEPCHK
    do CC = CC1, CC2
        N1 = cell(CC,4)
        N2 = cell(CC,5)
        N3 = cell(CC,6)

        dx31 = x(N3) - x(N1)
        dx12 = x(N1) - x(N2)
        dx23 = x(N2) - x(N3)
        dy31 = y(N3) - y(N1)
        dy12 = y(N1) - y(N2)
        dy23 = y(N2) - y(N3)

        coef = 0.5/abs(-dx12*dy31 + dy12*dx31)

        & dxC1 = (U(1,N1)*dy23 + U(1,N2)*dy31 + U(1,N3)*dy12)
        & *coef
        & dxC2 = (U(2,N1)*dy23 + U(2,N2)*dy31 + U(2,N3)*dy12)
        & *coef
        & dxC3 = (U(3,N1)*dy23 + U(3,N2)*dy31 + U(3,N3)*dy12)
        & *coef
        & dxC4 = (U(4,N1)*dy23 + U(4,N2)*dy31 + U(4,N3)*dy12)
        & *coef

        & dyC1 = (U(1,N1)*dx23 + U(1,N2)*dx31 + U(1,N3)*dx12)
        & *coef
        & dyC2 = (U(2,N1)*dx23 + U(2,N2)*dx31 + U(2,N3)*dx12)
        & *coef
        & dyC3 = (U(3,N1)*dx23 + U(3,N2)*dx31 + U(3,N3)*dx12)
        & *coef
        & dyC4 = (U(4,N1)*dx23 + U(4,N2)*dx31 + U(4,N3)*dx12)
        & *coef

        del2(1,N1) = del2(1,N1) + (dxC1*dy23 + dyC1*dx23)
        del2(1,N2) = del2(1,N2) + (dxC1*dy31 + dyC1*dx31)
        del2(1,N3) = del2(1,N3) + (dxC1*dy12 + dyC1*dx12)

        del2(2,N1) = del2(2,N1) + (dxC2*dy23 + dyC2*dx23)
        del2(2,N2) = del2(2,N2) + (dxC2*dy31 + dyC2*dx31)
        del2(2,N3) = del2(2,N3) + (dxC2*dy12 + dyC2*dx12)

        del2(3,N1) = del2(3,N1) + (dxC3*dy23 + dyC3*dx23)
        del2(3,N2) = del2(3,N2) + (dxC3*dy31 + dyC3*dx31)
        del2(3,N3) = del2(3,N3) + (dxC3*dy12 + dyC3*dx12)

        del2(4,N1) = del2(4,N1) + (dxC4*dy23 + dyC4*dx23)
        del2(4,N2) = del2(4,N2) + (dxC4*dy31 + dyC4*dx31)
        del2(4,N3) = del2(4,N3) + (dxC4*dy12 + dyC4*dx12)
    enddo
enddo

c* upper and lower boundary
    EF2 = 0

```

```

do col = 1, Eicolormax
  EF1 = EF2 + 1
  EF2 = EF1 - 1 + NEcolor(col)
CVD$ NODEPCHK
  do EF = EF1, EF2
    FF = eface(EF)
    CC = face(FF,1)
    N1 = face(FF,3)
    N2 = face(FF,4)
    N3 = face(FF,5)

    dx31 = x(N3) - x(N1)
    dx12 = x(N1) - x(N2)
    dx23 = x(N2) - x(N3)
    dy31 = y(N3) - y(N1)
    dy12 = y(N1) - y(N2)
    dy23 = y(N2) - y(N3)

    coef = 0.5/abs(-dx12*dy31 + dy12*dx31)

    dxC1 = (U(1,N1)*dy23 + U(1,N2)*dy31 + U(1,N3)*dy12)
    *coef
    dxC2 = (U(2,N1)*dy23 + U(2,N2)*dy31 + U(2,N3)*dy12)
    *coef
    dxC3 = (U(3,N1)*dy23 + U(3,N2)*dy31 + U(3,N3)*dy12)
    *coef
    dxC4 = (U(4,N1)*dy23 + U(4,N2)*dy31 + U(4,N3)*dy12)
    *coef

    dyC1 = (U(1,N1)*dx23 + U(1,N2)*dx31 + U(1,N3)*dx12)
    *coef
    dyC2 = (U(2,N1)*dx23 + U(2,N2)*dx31 + U(2,N3)*dx12)
    *coef
    dyC3 = (U(3,N1)*dx23 + U(3,N2)*dx31 + U(3,N3)*dx12)
    *coef
    dyC4 = (U(4,N1)*dx23 + U(4,N2)*dx31 + U(4,N3)*dx12)
    *coef

    del2(1,N1) = del2(1,N1) + (dxC1*dy12 + dyC1*dx12)
    del2(1,N2) = del2(1,N2) + (dxC1*dy12 + dyC1*dx12)

    del2(2,N1) = del2(2,N1) + (dxC2*dy12 + dyC2*dx12)
    del2(2,N2) = del2(2,N2) + (dxC2*dy12 + dyC2*dx12)

    del2(3,N1) = del2(3,N1) + (dxC3*dy12 + dyC3*dx12)
    del2(3,N2) = del2(3,N2) + (dxC3*dy12 + dyC3*dx12)

    del2(4,N1) = del2(4,N1) + (dxC4*dy12 + dyC4*dx12)
    del2(4,N2) = del2(4,N2) + (dxC4*dy12 + dyC4*dx12)

  enddo
enddo

endif
CVD$ NODEPCHK
do PH = 1, Pmax
  P1 = pnode(PH,1)
  P2 = pnode(PH,2)
  del2(1,P1) = del2(1,P1) + del2(1,P2)
  del2(2,P1) = del2(2,P1) + del2(2,P2)
  del2(3,P1) = del2(3,P1) + del2(3,P2)
  del2(4,P1) = del2(4,P1) + del2(4,P2)
  del2(1,P2) = del2(1,P1)
  del2(2,P2) = del2(2,P1)
  del2(3,P2) = del2(3,P1)
  del2(4,P2) = del2(4,P1)
enddo

FF2 = 0

```

```

do col = 1, Fcolormax
  FF1 = FF2 + 1
  FF2 = FF1 - 1 + NFcolor(col)
CVD$ NCDEPCHK
do FF = FF1, FF2
  N1 = face(FF,3)
  N2 = face(FF,4)

  coef = 0.5*(eps1(N1) + eps1(N2))
  coef2 = 0.5*max(0.,eps2-coef)*(1./deltN(N1) + 1./deltN(N2))

  change = coef2*(del2(1,N2) - del2(1,N1)) +
  & coef*(U(1,N2) - U(1,N1))
  dis(1,N1) = dis(1,N1) + change
  dis(1,N2) = dis(1,N2) - change

  change = coef2*(del2(2,N2) - del2(2,N1)) +
  & coef*(U(2,N2) - U(2,N1))
  dis(2,N1) = dis(2,N1) + change
  dis(2,N2) = dis(2,N2) - change

  change = coef2*(del2(3,N2) - del2(3,N1)) +
  & coef*(U(3,N2) - U(3,N1))
  dis(3,N1) = dis(3,N1) + change
  dis(3,N2) = dis(3,N2) - change

  change = coef2*(del2(4,N2) - del2(4,N1)) +
  & coef*(U(4,N2) - U(4,N1))
  dis(4,N1) = dis(4,N1) + change
  dis(4,N2) = dis(4,N2) - change
enddo
enddo

c** find eps1 and del2 at periodic nodes
do PN = 1, Pmax
  P1 = pnode(PN,1)
  P2 = pnode(PN,2)
  dis(1,P1) = dis(1,P1) + dis(1,P2)
  dis(2,P1) = dis(2,P1) + dis(2,P2)
  dis(3,P1) = dis(3,P1) + dis(3,P2)
  dis(4,P1) = dis(4,P1) + dis(4,P2)
  dis(1,P2) = dis(1,P1)
  dis(2,P2) = dis(2,P1)
  dis(3,P2) = dis(3,P1)
  dis(4,P2) = dis(4,P1)
enddo

return
end

c*****
c*
c* adjust inlet state vector for boundary condition
c*
c*****

subroutine binlet
implicit none

include 'TRI.INC'
real rrinl, uuinl, vvlinl, ppinl !average values at inlet
real ww2inl, aainl
real r0inf, a0inf !stag. density and speed of sound
real h0pres !stagnation enthalpy and pressure
real spres !entropy
real coef

```

```

real aaa1, aaa2, aaa3, aaa4      !coef. of inverse of matrix
real bbb1, bbb2, bbb3, bbb4
real ccc1, ccc2, ccc3, ccc4
real ddd1, ddd2, ddd3, ddd4
real ww2, uu, vv, pp, rr, aa      !values at next interior node
real dHO, ds, dtan, dw4
real drr, duu, dvv, dpp
integer IN, NN                    !boundary face and its nodes

c** average values at inlet
rrinl = U(1,innode(Imax/2)) - dU(1,innode(Imax/2))
uvinl = (U(2,innode(Imax/2)) - dU(2,innode(Imax/2)))/rrinl
vvinl = (U(3,innode(Imax/2)) - dU(3,innode(Imax/2)))/rrinl
ww2inl = uvinl**2 + vvinl**2
ppinl = gam1*((U(4,innode(Imax/2)) - dU(4,innode(Imax/2)))
*      - 0.5*rrinl*ww2inl)
aaainl = sqrt(gam*ppinl/rrinl)

c* subsonic inlet nodes
if ((sqrt(ww2inl)/aaainl).lt.1. .or. pitch.ne.0.) then
c** prescribed values
HOpres = 1./gam1
spres = 0.

c** coefficient of matrix
coef = 1./(uvinl*(aaainl+uvinl) + vvinl**2)
aaa1 = (rrinl*uvinl/aaainl)*coef
aaa2 = -(ppinl/aaainl)*(uvinl*gam/gam1 + ww2inl/aaainl)*coef
aaa3 = -(rrinl*vvinl/aaainl)*coef
aaa4 = (ww2inl/aaainl**2)*coef
bbb1 = uvinl*coef
bbb2 = -(uvinl*ppinl/(gam1*rrinl))*coef
bbb3 = -vvinl*coef
bbb4 = -(uvinl/rrinl)*coef
ccc1 = vvinl*coef
ccc2 = -(vvinl*ppinl/(gam1*rrinl))*coef
ccc3 = (aaainl + uvinl)*coef
ccc4 = -(vvinl/rrinl)*coef
ddd1 = rrinl*aaainl*uvinl*coef
ddd2 = -(ppinl*aaainl*uvinl/gam1)*coef
ddd3 = -rrinl*aaainl*vvinl*coef
ddd4 = ww2inl*coef

CVD$ NODEPCHK
do IN = 1, Imax
NN = innode(IN)
rr = U(1,NN) - dU(1,NN)
uu = (U(2,NN) - dU(2,NN))/rr
vv = (U(3,NN) - dU(3,NN))/rr
ww2 = uu**2 + vv**2
pp = gam1*((U(4,NN) - dU(4,NN)) - 0.5*rr*ww2)
aa = sqrt(gam*pp/rr)
dHO = HOpres - ((gam/gam1)*pp/rr + 0.5*ww2)
ds = spres - (log(gam*pp) - gam*log(rr))
dtan = (S1nl - vv/uu)*uvinl**2
dw4 = dU(1,NN)*(aa*uu + gam1*ww2)
*      - dU(2,NN)*(aa + gam1*uu)
*      - dU(3,NN)*gam1*vv
*      + dU(4,NN)*gam1
drr = aaa1*dHO + aaa2*ds + aaa3*dtan + aaa4*dw4
duu = bbb1*dHO + bbb2*ds + bbb3*dtan + bbb4*dw4
dvv = ccc1*dHO + ccc2*ds + ccc3*dtan + ccc4*dw4
dpp = ddd1*dHO + ddd2*ds + ddd3*dtan + ddd4*dw4

U(1,NN) = U(1,NN) - dU(1,NN) + drr
U(2,NN) = U(2,NN) - dU(2,NN) + rr*duu + uu*drr
U(3,NN) = U(3,NN) - dU(3,NN) + rr*dvv + vv*drr
U(4,NN) = U(4,NN) - dU(4,NN) + dpp/gam1 + 0.5*ww2*drr
*      + rr*(uu*duu + vv*dvv)
enddo

c* supersonic inlet nodes

```

```

      else if ((sqrt(vw2inl)/aainl).ge.1.) then
c* state vector components far from body
      r0inf = 1.
      a0inf = 1.

CVD$ NODEPCHK
      do IN = 1, Imax
        NN = innode(IN)
        rr = r0inf*(1.0+0.5*gam1*Minl**2)**(-1./gam1)
        aa = a0inf*(1.0+0.5*gam1*Minl**2)**(-0.5)
        uu = Minl*aa
        pp = rr*aa**2/gam

        U(1,NN) = rr
        U(2,NN) = rr*uu
        U(3,NN) = 0.
        U(4,NN) = pp/gam1 + .5*(uu**2)*rr
      enddo
    endif

    return
  end

c*****
c*
c*      adjust outlet state vector for boundary condition
c*
c*****

      subroutine boutlet
      implicit none

      include 'TRI.INC'
      real rrout, uuout, vvout, ppout !average values at outlet
      real aaout, vw2out
      real aaa1, aaa4 !coef. of inverse of matrix
      real bbb3, bbb4
      real ccc2
      real ddd4
      real vw2, uu, vv, pp, rr, aa !values at next interior node
      real dw1, dw2, dw3, dp
      real drr, duu, dvv, dpp
      integer ON, NN !boundary face and its nodes

      rrout = U(1,outnode(Onmax/2)) - dU(1,outnode(Onmax/2))
      uuout = (U(2,outnode(Onmax/2)) - dU(2,outnode(Onmax/2)))/rrout
      vvout = (U(3,outnode(Onmax/2)) - dU(3,outnode(Onmax/2)))/rrout
      vw2out = uuout**2 + vvout**2
      ppout = gam1*((U(4,outnode(Onmax/2)) - dU(4,outnode(Onmax/2)))
      * - 0.5*rrout*vw2out)
      aaout = sqrt(gam*ppout/rrout)

      aaa1 = -1./aaout**2
      aaa4 = 1./aaout**2
      bbb3 = .1/(rrout*aaout)
      bbb4 = -1./(rrout*aaout)
      ccc2 = .1/(rrout*aaout)
      ddd4 = 1.

c* set boundary values of state vector for nodes
CVD$ NODEPCHK
      do ON = 1, Onmax
        NN = outnode(ON)
        rr = U(1,NN) - dU(1,NN)
        uu = (U(2,NN) - dU(2,NN))/rr
        vv = (U(3,NN) - dU(3,NN))/rr
        vw2 = uu**2 + vv**2
        pp = gam1*((U(4,NN) - dU(4,NN)) - 0.5*rr*vw2)
        aa = sqrt(gam*pp/rr)

```

```

      if (sqrt(ww2) .lt. aa) then
        dw1 = dU(1,NN)*(0.5*ww2*gami - aa**2)
&         - dU(2,NN)*gami*uu
&         - dU(3,NN)*gami*vv
&         + dU(4,NN)*gami
        dw2 = - dU(1,NN)*aa*vv
&         + dU(3,NN)*aa
        dw3 = dU(1,NN)*(0.5*ww2*gami - aa*uu)
&         - dU(2,NN)*(gami*uu - aa)
&         - dU(3,NN)*gami*vv
&         + dU(4,NN)*gami
        dp = pout - pp
        drr = aaa1*dw1 + aaa4*dp
        duu = bbb3*dw3 + bbb4*dp
        dvv = ccc2*dw2
        dpp = ddd4*dp
        U(1,NN) = U(1,NN) - dU(1,NN) + drr
        U(2,NN) = U(2,NN) - dU(2,NN) + rr*duu + uu*drr
        U(3,NN) = U(3,NN) - dU(3,NN) + rr*dvv + vv*drr
        U(4,NN) = U(4,NN) - dU(4,NN) + dpp/gami + 0.5*ww2*drr
&         + rr*(uu*duu + vv*dvv)
      endif
    enddo

  return
end

```

```

C*****
C*
C*   this subroutine changes the momentum change to make
C*   flow tangent to wall
C*
C*****

```

```

      subroutine tangent
      implicit none

      include 'TRI.INC'
      integer EN                      !pointer
      integer NN                      !node on wall
      real drwn                       !change in momentum normal to wall

      do EN = 1, bnode
        NN = enode(EN)
        drwn = -U(2,NN)*senode(EN) + U(3,NN)*cerode(EN)
        U(2,NN) = U(2,NN) + drwn*senode(EN)
        U(3,NN) = U(3,NN) - drwn*cerode(EN)
      enddo

      return
end

```

A.1.5 Plotting Package

```

      program plotgen
      implicit none

      include 'TRI.INC'

      integer NN, EN, CC              !pointers
      integer N1, N2, N3

```

```

integer ptype          !type of plot
integer ctype          !type of contour
integer stype          !type of surface distributions
character*80 TITLE, ITITLE !title for plots
integer NTITL          !number of letters in title
external g2pltg, g2pltc !plotting subroutines
integer indgr
integer a4, a5, a6, a7, a8, a9, a10 !dummy variables
real uu, vv, pp, rr, mm, mm2 !from state vectors
real pt, ptinf         !total pressure
real zz, z(Maxnodes)   !contour values
real zmax, zmin        !max and min of z array
integer NCONT          !contour level info
real CBASE, CSTEP
real Cinc              !contour increment
character*6 NUM        !same as CSTEP but character
integer NLINE, IOPT(2) !indicators for line plots
integer points, npts(2) !number of points on line
real xline(Maxedges)   !points on line to plot
real yline(Maxedges)

c* read data from file
call gridio(1)
call flowio(1)

c* initialize GRAFIC
write(ITITLE,1) Min1
1   format('INLET MACH NUMBER = ',F5.3)
call grinit(5,6,ITITLE)

do while (1)

c* proxpt user for type of plot
type*, 'Type of plot '
type*, ' 0) STOP '
type*, ' 1) grid '
type*, ' 2) contour'
type*, ' 3) surface distribution'
type*, ' 4) data'
type i1
11  format(1,' selection = ')
accept i11, ptype
111 format(I)

if (ptype.eq.0) then
stop
else if (ptype.eq.1) then

TITLE = ' X Y COMPUTATIONAL GRID'
indgr = 23
call gr_control(g2pltg, indgr, TITLE, x, y, Nmax, a4,
& a5, a6, a7, a8, a9, a10)

else if (ptype.eq.2) then
c* choose type of contour plot
do while (1)

Cinc = 0.

2   type*, 'Type of contour'
type*, ' 0) TOP LEVEL'
type*, ' 1) density'
type*, ' 2) Mach number'
type*, ' 3) normal velocity'
type*, ' 4) pressure'
type*, ' 5) total pressure loss'
type*, ' 6) speed of sound'
type*, ' 7) entropy'
type*, ' 8) stagnation enthalpy'
type*, ' 9) vol'
type*, ' 10) CONTOUR INCREMENT'
type 22

```

```

22      format(9,' selection = ')
      accept 222, ctype
222     format(I)

c*  exit from contour loop
      if (ctype.eq.0) goto 999

      if (ctype.eq.10) then
        type 21
21      format(9,' CONTOUR INCREMENT = ')
        accept 221, Cinc
221     format(F)
        type*, ' '
        goto 2
      endif

c*  set up nodal contour values
      if (ctype.eq.5) then
        ptinf = 1./gam
      endif

      zmin = 1.e20
      zmax = -1.e20

      do NN = 1, Nmax
        rr = U(1,NN)
        uu = U(2,NN)/rr
        vv = U(3,NN)/rr

        if (ctype.eq.1) then
          zz = rr
        else if (ctype.eq.2) then
          pp = gam1*(U(4,NN) - 0.5*rr*(uu**2 + vv**2))
          mm = sqrt(rr*(uu**2 + vv**2)/(gam*pp))
          zz = mm
        else if (ctype.eq.3) then
          zz = sqrt(uu**2 + vv**2)
        else if (ctype.eq.4) then
          pp = gam1*(U(4,NN) - 0.5*rr*(uu**2 + vv**2))
          zz = pp
        else if (ctype.eq.5) then
          pp = gam1*(U(4,NN) - 0.5*rr*(uu**2 + vv**2))
          mm2 = rr*(uu**2 + vv**2)/(gam*pp)
          pt = pp*(1. + 5*gam1*mm2)**(gam/gam1)
          zz = (1. - pt/ptinf)*100.
        else if (ctype.eq.6) then
          pp = gam1*(U(4,NN) - 0.5*rr*(uu**2 + vv**2))
          zz = sqrt(gam*pp/rr)
        else if (ctype.eq.7) then
          pp = gam1*(U(4,NN) - 0.5*rr*(uu**2 + vv**2))
          zz = (log(gam*pp) - gam*log(rr))*100.
        else if (ctype.eq.8) then
          pp = gam1*(U(4,NN) - 0.5*rr*(uu**2 + vv**2))
          zz = ((gam/gam1)*(pp/rr) + 0.5*(uu**2 + vv**2))*100.
        endif

        zmax = max(zmax,zz)
        zmin = min(zmin,zz)
        z(NN) = zz
      enddo

      if (ctype.eq.9) then
        open(unit=50, status='unknown', form='unformatted')
        read(50) Cmax, (vol(CC),CC=1,Cmax)
        close(unit=50)
        zmin = 1.e20
        zmax = -1.e20
        do NN = 1, Nmax
          z(NN) = 0.
        enddo
        do CC = 1, Cmax
          N1 = cell(CC,4)

```



```

        N3 = cell(CC,5)
        N3 = cell(CC,6)
        zz = vol(CC)
        zmax = max(zmax,zz)
        zmin = min(zmin,zz)
        z(N1) = z(N1) + zz/3.
        z(N2) = z(N2) + zz/3.
        z(N3) = z(N3) + zz/3.
    enddo
endif

c* set title and extra variables needed for GRAFIC
if (ctype.eq.1) then
    TITLE = '      X      Y      DENSITY '
    NTITL = 24
    NCONT = 20
else if (ctype.eq.2) then
    TITLE = '      X      Y      MACH NUMBER '
    NTITL = 28
    NCONT = 20
else if (ctype.eq.3) then
    TITLE = '      X      Y      NORMAL VELOCITY '
    NTITL = 32
    NCONT = 20
else if (ctype.eq.4) then
    TITLE = '      X      Y      PRESSURE '
    NTITL = 25
    NCONT = 20
else if (ctype.eq.5) then
    TITLE = '      X      Y      % TOTAL PRESSURE LOSS '
    NTITL = 38
    NCONT = 20
else if (ctype.eq.6) then
    TITLE = '      X      Y      SPEED OF SOUND '
    NTITL = 31
    NCONT = 20
else if (ctype.eq.7) then
    TITLE = '      X      Y      % ENTROPY '
    NTITL = 24
    NCONT = 20
else if (ctype.eq.8) then
    TITLE = '      X      Y      % STAGNATION ENTHALPY '
    NTITL = 38
    NCONT = 20
else if (ctype.eq.9) then
    TITLE = '      X      Y      VOL '
    NTITL = 20
    NCONT = 20
endif

c* find contour levels
if (Cinc.ne.0.) then
    NCONT = int((zmax-zmin)/Cinc + 2.)
    CSTEP = Cinc
    CBASE = (real(int((zmin/Cinc)-1.)))*Cinc
else
    call GR_SCALE(zmin, zmax, NCONT-1, CBASE, CSTEP)
endif

z(Nmax+1) = NCONT
z(Nmax+2) = CBASE + 0.01*CSTEP
z(Nmax+3) = CSTEP

c* finish up title
TITLE(NTITL+1:NTITL+24) = 'CONTOURS WITH INCREMENT '
write(NUM,10) CSTEP
format(F6.4)
TITLE(NTITL+25:80) = NUM

c* plot the contour lines
indgr = 23

```

```

      call gr_control(g2pltc, indgr, TITLE, z, Nmax, x, y,
*          a5, a6, a7, a8, a9, a10)

      enddo

c* plot surface distribution
      else if (ptype.eq.3) then
        do while (1)

          type*, 'Type of surface distribution'
          type*, ' 0) TOP LEVEL'
          type*, ' 1) density'
          type*, ' 2) Mach number'
          type*, ' 3) normal velocity'
          type*, ' 4) surface flow angle'
          type*, ' 5) analytical surface flow angle'
          type*, ' 6) pressure'
          type*, ' 7) entropy'
          type*, ' 8) stagnation enthalpy'
          type*, ' 9) total pressure loss'
          type 33
          33      format(9,' selection = ')
          accept 333, stype
          333      format(I)

c* exit from contour loop
          if (stype.eq.0) goto 999

c* set up array with points on line for upper surface
          if (stype.eq.9) then
            ptinf = 1./gam
            endif

          points = 0
          do EN = 1, tnode
            points = points + 1
            NN = enode(EN)
            rr = U(1,NN)
            uu = U(2,NN)/rr
            vv = U(3,NN)/rr

            if (stype.eq.1) then
              zz = rr
            else if (stype.eq.2) then
              pp = gam1*(U(4,NN) - 0.5*rr*(uu**2 + vv**2))
              mm = sqrt(rr*(uu**2 + vv**2)/(gam*pp))
              zz = mm
            else if (stype.eq.3) then
              zz = sqrt(uu**2 + vv**2)
            else if (stype.eq.4) then
              zz = atan(vv/uu)
            else if (stype.eq.5) then
              zz = atan(senode(EN)/cenode(EN))
            else if (stype.eq.6) then
              pp = gam1*(U(4,NN) - 0.5*rr*(uu**2 + vv**2))
              zz = pp
            else if (stype.eq.7) then
              pp = gam1*(U(4,NN) - 0.5*rr*(uu**2 + vv**2))
              zz = (log(gam*pp) - gam*log(rr))*100.
            else if (stype.eq.8) then
              pp = gam1*(U(4,NN) - 0.5*rr*(uu**2 + vv**2))
              zz = ((gam/gam1)*(pp/rr) + 0.5*(uu**2 + vv**2))*100.
            else if (stype.eq.9) then
              pp = gam1*(U(4,NN) - 0.5*rr*(uu**2 + vv**2))
              mm2 = rr*(uu**2 + vv**2)/(gam*pp)
              pt = pp*(1.+5*gam1*mm2)**(gam/gam1)
              zz = (1. - pt/ptinf)*100.
            endif

            xline(points) = x(NN)
            yline(points) = zz
          enddo
        enddo
      enddo

```

```

npts(1) = points

c* set up array with points on line for lower surface
points = 0
do EN = tnode+1, bnode
  points = points + 1
  NN = enode(EN)
  rr = U(1,NN)
  uu = U(2,NN)/rr
  vv = U(3,NN)/rr

  if (stype.eq.1) then
    zz = rr
  else if (stype.eq.2) then
    pp = gam1*(U(4,NN) - 0.5*rr*(uu**2 + vv**2))
    mm = sqrt(rr*(uu**2 + vv**2)/(gam*pp))
    zz = mm
  else if (stype.eq.3) then
    zz = sqrt(uu**2 + vv**2)
  else if (stype.eq.4) then
    zz = atan(vv/uu)
  else if (stype.eq.5) then
    zz = atan(senode(EN)/cenode(EN))
  else if (stype.eq.6) then
    pp = gam1*(U(4,NN) - 0.5*rr*(uu**2 + vv**2))
    zz = pp
  else if (stype.eq.7) then
    pp = gam1*(U(4,NN) - 0.5*rr*(uu**2 + vv**2))
    zz = (log(gam*pp) - gam*log(rr))*100.
  else if (stype.eq.8) then
    pp = gam1*(U(4,NN) - 0.5*rr*(uu**2 + vv**2))
    zz = ((gam/gam1)*(pp/rr) + 0.5*(uu**2 + vv**2))*100.
  else if (stype.eq.9) then
    pp = gam1*(U(4,NN) - 0.5*rr*(uu**2 + vv**2))
    mm2 = rr*(uu**2 + vv**2)/(gam*pp)
    pt = pp*(1.+5*gam1*mm2)**(gam/gam1)
    zz = (1. - pt/ptinf)*100.
  endif

  xline(npts(1)+points) = x(NN)
  yline(npts(1)+points) = zz
enddo

npts(2) = points

c** set plot indicators
NLINE = 2
IOPT(1) = 14
IOPT(2) = 14

c** set title for plot
if (stype.eq.1) then
  TITLE = ' X DENSITY DENSITY'
else if (stype.eq.2) then
  TITLE = ' X MACH NO.MACH NUMBER'
else if (stype.eq.3) then
  TITLE = ' X VELOCITYNORMAL VELOCITY'
else if (stype.eq.4) then
  TITLE = ' X ANGLE SURFACE FLOW ANGLE'
else if (stype.eq.5) then
  TITLE = ' X ANGLE ANALYTICAL SURFACE FLOW ANGLE'
else if (stype.eq.6) then
  TITLE = ' X PRESSUREPRESSURE'
else if (stype.eq.7) then
  TITLE = ' X ENTROPY~% ENTROPY'
else if (stype.eq.8) then
  TITLE =
& ' X ~STAGNATION ENTHALPY~% STAGNATION ENTHALPY'
else if (stype.eq.9) then
  TITLE =
& ' X ~TOTAL PRESSURE LOSS~% TOTAL PRESSURE LOSS'

```

```

endif

c* plot the line
indgr = 21
call gr_line(IOPT,NLINE,TITLE,INDGR,xline,yline,npts)

enddo

else if (ptype.eq.4) then
c* calculate mean of total pressure loss
ptinf = 1./gam
zz = 0.
zmax = 0.
do NN = 1, Nmax
rr = U(1,NN)
uu = U(2,NN)/rr
vv = U(3,NN)/rr
pp = gam1*(U(4,NN) - 0.5*rr*(uu**2 + vv**2))
mm2 = rr*(uu**2 + vv**2)/(gam*pp)
pt = pp*(1.+5*gam1*mm2)**(gam/gam1)
zz = zz + (1. - pt/ptinf)**2
zmax = max(zmax,abs(1.-pt/ptinf))
enddo
zz = sqrt(zz/real(Nmax))

type*, ' '
type*, 'inlet Mach number      = ', Minl
type*, 'rms total pres. loss   = ', zz, ' (',log10(zz),')'
type*, 'max total pres. loss   = ', zmax, ' (',log10(zmax),')'
type*, 'cells                  = ', Cmax
type*, 'faces                  = ', Fmax
type*, 'nodes                  = ', Nmax
type*, 'inlet nodes            = ', Imax
type*, 'outlet nodes           = ', Omax
type*, 'periodic nodes         = ', Pmax
type*, 'edge nodes             = ', Emax
type*, 'face colors            = ', Fcolormax
type*, 'cell colors            = ', Ccolormax
type*, 'edge colors            = ', E2colormax
type*, ' '
999 endif
enddo
end

subroutine g2pltg(ifun, indgr, TITLE, alimits, info_string,
& a1, a2, a3, a4, a5, a6, a7, a8, a9, a10)
implicit none

include 'TRI.INC'

integer FF, EF, NN
integer indgr
character*80 TITLE
integer ifun
real alimits(4)
integer a1, a2, a3, a4, a5, a6, a7, a8, a9, a10
character*80 info_string
real x1, x2, y1, y2

if (ifun.eq.0) then
return
elseif (ifun.eq.1) then
call gr_get_limits(x, y, Nmax, alimits)
if (pitch.ne.0.) then
alimits(4) = alimits(4) + pitch
endif
elseif (ifun.eq.2) then
info_string = ' '
elseif (ifun.eq.3) then

```

```

do FF = 1, Fmax
  NN = face(FF,3)
  x1 = x(NN)
  y1 = y(NN)
  NN = face(FF,4)
  x2 = x(NN)
  y2 = y(NN)
  call gr_move(x1, y1, 0)
  call gr_draw(x2, y2, 0)
  if (pitch.ne.0.) then
    y1 = y1 + pitch
    y2 = y2 + pitch
    call gr_move(x1, y1, 0)
    call gr_draw(x2, y2, 0)
  endif
enddo
endif

return
end

```

```

subroutine g2pltc(ifun, indgr, TITLE, alimits, info_string,
&               z, a2, a3, a4, a5, a6, a7, a8, a9, a10)
implicit none

include 'TRI.INC'

integer FF, EF, NN, CC, NC      !pointers
integer node                    !pointer
integer indgr
character*80 TITLE
integer ifun
real alimits(4)
real z(Maxnodes)
integer a2, a3, a4, a5, a6, a7, a8, a9, a10
character*80 info_string
real y1, y2, y3                !location of cell nodes
integer ncont                   !determine contour levels
real cbase, cstep
real zn12, zn32, zn31           !contour crossing on sides
integer N1, N2, N3              !nodes at end of edge face
real xn(3), yn(3), zn(3)        !x, y, and contour value at nodes
real xpoint, ypoint            !x, y of pointer
real atriangle                 !area of triangle
real value                     !value of contour at pointer
real cn(2,3)                   !x, y at cell nodes
integer IIN                     !is pointer in cell?
real aa1, aa2, aa3, aa         !areas
real zcont, cont(80)           !contour levels
real zmax, zmin                !max & min contour values in cell

if (ifun.eq.0) then
  return
elseif (ifun.eq.1) then
  call gr_get_limits(x, y, Nmax, alimits)
  if (pitch.ne.0.) then
    alimits(4) = alimits(4) + pitch
  endif
elseif (ifun.eq.2) then
c* find contour value at point
  xpoint = alimits(1)
  ypoint = alimits(2)
c* look for cell (x,y) is in

```

```

do CC = 1, Cmax
  N1 = cell(CC,4)
  N2 = cell(CC,5)
  N3 = cell(CC,6)
  cn(1,1) = x(N1)
  cn(1,2) = x(N2)
  cn(1,3) = x(N3)
  cn(2,1) = y(N1)
  cn(2,2) = y(N2)
  cn(2,3) = y(N3)
  call gr_inside(IIN,cn,3,xpoint,ypoint)
  if (IIN.eq.1) then

c* linear interpolation for value
    aa2 = atriangle(x(N1),y(N1),x(N3),y(N3),xpoint,ypoint)
    aa3 = atriangle(x(N2),y(N2),x(N1),y(N1),xpoint,ypoint)
    aa1 = atriangle(x(N3),y(N3),x(N2),y(N2),xpoint,ypoint)
    aa = atriangle(x(N3),y(N3),x(N2),y(N2),x(N1),y(N1))
    if (aa.eq.0) then
      aa1 = 1.
      aa2 = 1.
      aa3 = 1.
      aa = 3.
    endif
    value = (aa1*z(N1) + aa2*z(N2) + aa3*z(N3))/aa
    goto 30
  endif
  cn(2,1) = cn(2,1) + pitch
  cn(2,2) = cn(2,2) + pitch
  cn(2,3) = cn(2,3) + pitch
  call gr_inside(IIN,cn,3,xpoint,ypoint)
  if (IIN.eq.1) then

c* linear interpolation for value
    y1 = y(N1) + pitch
    y2 = y(N2) + pitch
    y3 = y(N3) + pitch
    aa2 = atriangle(x(N1),y1,x(N3),y3,xpoint,ypoint)
    aa3 = atriangle(x(N2),y2,x(N1),y1,xpoint,ypoint)
    aa1 = atriangle(x(N3),y3,x(N2),y2,xpoint,ypoint)
    aa = atriangle(x(N3),y3,x(N2),y2,x(N1),y1)
    if (aa.eq.0) then
      aa1 = 1.
      aa2 = 1.
      aa3 = 1.
      aa = 3.
    endif
    value = (aa1*z(N1) + aa2*z(N2) + aa3*z(N3))/aa
    goto 30
  endif
onddo
30  write (INFO_STRING,31) value
31  format(' function value = ',f15.6)

elseif (ifun.eq.3) then

c* plot boundary
  do EF = 1, Emax
    FF = eface(EF)
    N1 = face(FF,3)
    N2 = face(FF,4)
    call gr_move(x(N1), y(N1), 0)
    call gr_draw(x(N2), y(N2), 0)
  enddo

  if (pitch.ne.0.) then
    do Ef = 1, Emax
      FF = eface(EF)
      N1 = face(FF,3)
      N2 = face(FF,4)
      y1 = y(N1) + pitch
      y2 = y(N2) + pitch

```

```

        call gr_move(x(N1), y1, 0)
        call gr_draw(x(N2), y2, 0)
    enddo
endif

c* set up contour levels
ncont = z(Nmax+1)
cbase = z(Nmax+2)
cstep = z(Nmax+3)
do NC = 1, ncont
    cont(NC) = cbase + (NC-1)*cstep
enddo

c* loop through cells
do CC = 1, Cmax

    do NN = 1, 3
        node = cell(CC, NN+3)
        xn(NN) = x(node)
        yn(NN) = y(node)
        zn(NN) = z(node)
    enddo
    zmax = max(zn(1), zn(2), zn(3))
    zmin = min(zn(1), zn(2), zn(3))

c* find contour crossing on triangle for each contour level
    do NC = 1, ncont
        zcont = cont(NC)
        if (zcont.lt.zmin .or. zcont.gt.zmax) then
c* no need to compute
        else
c* find contour crossing in triangle
            call gr_cross(zn(1), zn(2), zcont, zn12)
            call gr_cross(zn(3), zn(2), zcont, zn32)
            call gr_cross(zn(3), zn(1), zcont, zn31)

c* plot contour levels in triangle
            call gr_ctriangle(xn(1), yn(1),
&                             xn(2), yn(2),
&                             xn(3), yn(3),
&                             zn31, zn32, zn12)
            if (pitch.ne.0.) then
                y1 = yn(1) + pitch
                y2 = yn(2) + pitch
                y3 = yn(3) + pitch
                call gr_ctriangle(xn(1), y1,
&                             xn(2), y2,
&                             xn(3), y3,
&                             zn31, zn32, zn12)
            endif
        endif
    enddo
enddo

endif

return
end

c*****
c*
c* function to find the area of a triangle
c*
c*****

function atriangle(x1,y1,x2,y2,x3,y3)

real x1,y1,x2,y2,x3,y3
real*8 distance,a,b,c,s

```

```

distance(xa,ya,xb,yb) = dsqrt(dble(xa-xb)**2 + dble(ya-yb)**2)

a = distance(x1,y1,x2,y2)
b = distance(x3,y3,x2,y2)
c = distance(x1,y1,x3,y3)

s = .5*(a + b + c)
a = dsqrt(s*(s-a)*(s-b)*(s-c))
atriangle = a
return
end

```

A.2 Quadrilateral Schemes

A.2.1 Common Files

This is file QUAD.INC which includes many declarations and common block statements and is included in all subroutines for the quadrilateral schemes.

```

parameter Maxnodes = 5000
parameter Maxfaces = 10000
parameter Maxcells = 5000
parameter Maxedges = 400

integer Nmax           !number of nodes
integer Fmax           !number of faces (including edges)
integer Cmax           !number of cells (including edge cells)
integer Imax, Omax     !number of inlet and outlet nodes
integer Wmax           !number of wall faces
integer Emax           !number of edges
integer tedge, bedge, iedge !where different types of edges are
integer Pmax           !number of periodic nodes
integer ENmax          !number of edge nodes
integer tnode, bnode, inode !location of diff. types of edge nodes

real pitch             !blade pitch
real pout              !outlet pressure for blades
real Sinl              !tan of inlet flow angle
real x(Maxnodes)       !x values of nodes
real y(Maxnodes)       !y values of nodes
integer eface(Maxedges) !array of edge faces
integer enode(Maxedges) !array of edge nodes
real senode(Maxedges)  !sin and cos at edge nodes
real cenode(Maxedges)

integer face(Maxfaces,8) !array of faces
integer cell(Maxcells,8) !array of cells
integer innode(Maxedges) !inlet nodes
integer outnode(Maxedges) !outlet nodes
integer pnode(Maxedges,2) !periodic nodes
integer NCcolor(50)      !number of cells colored each color
integer Ccolormax        !max number of colors used
integer NFcolor(50)      !number of faces colored each color
integer Fcolormax        !max number of colors used
integer NEcolor(10)      !number of edges colored each color
integer E1colormax, E2colormax !max number of colors used

common /quad/ Imax, Omax, Wmax, Pmax
common /quad/ Nmax, Fmax, Cmax, Emax, tedge, bedge, iedge
common /quad/ ENmax, tnode, bnode, inode

```



```

common /quad/ pitch, pout, Sinl
common /quad/ x, y, eface, enode, senode, cenode, face, cell
common /quad/ innode, outnode, pnode
common /color/ NCcolor, Ccolormax, NFcolor, Fcolormax
common /color/ NEcolor, Ecolormax, E2colormax

real Minl                      !inlet Mach number
integer Maxiter                 !max number of iterations
real CFL                       !CFL number
real epsicoef, eps2            !smoothing coef
real sigE, sigV                !smoothing coef
real vol(Maxcells)             !for vorticity smoothing
real durms                     !rms difference in state vector
real U(4,0:Maxnodes)          !state vector
real dU(4,Maxnodes)           !change in state vector at nodes
real Uc(4,Maxcells)            !state vector at cells
real dUc(4,0:Maxcells)         !change in state vector at cells
real F(4,Maxnodes)            !flux vectors at nodes
real G(4,Maxnodes)
real areaN(0:Maxnodes)        !control area around each node
real areaC(0:Maxcells)        !area of each cell
real deltC(0:Maxcells)        !time step at each cell
real deltN(0:Maxnodes)        !time step at each node
real dis(4,Maxnodes)          !dissipation at each node
real flux(4,0:Maxnodes)       !flux at each node

common /flo/ Minl, durms, Maxiter, CFL
common /flo/ epsicoef, eps2, sigE, sigV, vol
common /flo/ U, dU, Uc, dUc
common /flo/ F, G
common /flo/ deltC, deltN, areaC, areaN, dis, flux

parameter gam = 1.4
parameter gam1 = 0.4

```

This file contains the subroutines for input and output and is linked with all the quadrilateral schemes.

```

c*****
c*
c*   write or read grid data from file
c*
c*****

subroutine gridio(process)
implicit none

include 'QUAD.INC'
integer process                !read or write?
integer i, NN, FF, CC, EN, EF, col !pointers
integer IN, ON, WN, PN

if (process.eq.1) then

c** read in data from file
open(unit=30, status='unknown', form='unformatted')

read(30) Nmax, (x(NN),NN=1,Nmax), (y(NN),NN=1,Nmax)
read(30) Fmax, ((face(FF,i),i=1,8),FF=1,Fmax)
read(30) tedge, bedge, iedge
read(30) Wmax, Emax, (eface(EF),EF=1,Emax)
read(30) tnode, bnode, ilode, ENmax, (enode(EN),EN=1,ENmax)
read(30) (senode(EN),EN=1,ENmax), (cenode(EN),EN=1,ENmax)
read(30) Imax, (innode(IN),IN=1,Imax)

```

```

        read(30) Omax, (outnode(ON),ON=1,Omax)
        read(30) Pmax, ((pnode(PN,1),i=1,2),PN=1,Pmax)
        read(30) Cmax, ((cell(CC,1),i=1,8),CC=1,Cmax)
        read(30) Ccolormax, (NCcolor(col),col=1,Ccolormax)
        read(30) Fcolormax, (NFcolor(col),col=1,Fcolormax)
        read(30) Ecolormax, E2colormax
        read(30) (NEcolor(col),col=1,E2colormax)
        read(30) pitch

        close(unit=30)

    else if (process.eq.0) then

c*  write data to file
        open(unit=30, status='unknown', form='unformatted')

        write(30) Nmax, (x(NN),NN=1,Nmax), (y(NN),NN=1,Nmax)
        write(30) Fmax, ((face(FF,1),i=1,8),FF=1,Fmax)
        write(30) tedge, bedge, iedge
        write(30) Wmax, Emax, (eface(EF),EF=1,Emax)
        write(30) tnode, bnode, inode, ENmax, (enode(EN),EN=1,ENmax)
        write(30) (senode(EN),EN=1,ENmax), (cenode(EN),EN=1,ENmax)
        write(30) Imax, (innode(IN),IN=1,Imax)
        write(30) Omax, (outnode(ON),ON=1,Omax)
        write(30) Pmax, ((pnode(PN,1),i=1,2),PN=1,Pmax)
        write(30) Cmax, ((cell(CC,1),i=1,8),CC=1,Cmax)
        write(30) Ccolormax, (NCcolor(col),col=1,Ccolormax)
        write(30) Fcolormax, (NFcolor(col),col=1,Fcolormax)
        write(30) Ecolormax, E2colormax
        write(30) (NEcolor(col),col=1,E2colormax)
        write(30) pitch

        close(unit=30)

    endif

    return
end

c*****
c*
c*      write or read flow data from file
c*
c*****

subroutine flowio(process)
    implicit none

    include 'QUAD.INC'
    integer process          !read or write?
    integer i, NN            !pointers

    if (process.eq.1) then

c** read in data from file
        open(unit=25, status='unknown', form='unformatted')

        read(25) Min1
        read(25) Nmax, ((U(1,NN),i=1,4),NN=1,Nmax)

        close(unit=25)

    else if (process.eq.0) then

c*  write data to file
        open(unit=25, status='unknown', form='unformatted')

        write(25) Min1

```

```

        write(25) Hmax, ((U(1,NN),i=1,4),NN=1,Nmax)

        close(unit=25)

endif

return
end

c*****
c*
c*      read input data from file
c*
c*****

      subroutine input
      implicit none

      include 'QUAD.INC'
      real pi                      !the one and only

      pi = 3.14159

      open(unit=20, status='old')

      read(20,*) Maxiter, CFL
      read(20,*) Minl, Sinl, pout
      read(20,*) sigE, sigV, epsicoef, eps2

      close (unit=20)

      pout = pout/gam
      Sinl = tan(pi*Sinl/180.0)

      return
      end

```

A.2.2 Mesh Generator

This is file GRID.INC which includes many declarations and common block statements for the mesh generator.

```

      parameter Maxdim = 200

      real xx(0:Maxdim,0:Maxdim)    !x coordinate of grid points
      real yy(0:Maxdim,0:Maxdim)    !y coordinate of grid points
      integer Nx                     !number cells on x axis
      integer Ny                     ! " " " y "
      integer ILE, ITE               !leading and trailing edge of blade

      common /grid/ xx, yy, Nx, Ny
      common /grid/ ILE, ITE

      parameter IBX=251

      real GSINL, GSOUT, CHINL, CHOUT
      integer II, JJ
      real XB(IBX), XPB(IBX), YB(IBX), YPB(IBX), SB(IBX)

```

```

integer IIB, IBLE
real SBLE, SBLULD, SS, SP
real XMIN, XMAX, YMIN, XLE, YLE, XPOS(Maxdim), YPOS(Maxdim)
integer NINL, NOUT, NBELD

common /c01/ GSINL, GSOUT, CHINL, CHOUT, II, JJ,
&          XB, XPB, YB, YPB, SB,
&          IIB, IBLE, SBLE, SBLULD, SS, SP,
&          XMIN, XMAX, YMIN, XLE, YLE, XPOS, YPOS,
&          NINL, NOUT, NBELD

c***** FACE ARRAY *****
c
c
c  N6-----N1-----N3      face(F,1) = C1
c  |         |         |      face(F,2) = C2
c  |         |         |      face(F,3) = N1
c  |         |         |      face(F,4) = N2
c  |         |         |      face(F,5) = N3
c  |         |         |      face(F,6) = N4
c  |         |         |      face(F,7) = N5
c  |         |         |      face(F,8) = N6
c  N5-----N2-----N4
c
c
c***** CELL ARRAY *****
c
c
c  N3-----F3-----N2      cell(C,1) = F1
c  |         |         |      cell(C,2) = F2
c  |         |         |      cell(C,3) = F3
c  |         |         |      cell(C,4) = F4
c  |         |         |      cell(C,5) = N1
c  |         |         |      cell(C,6) = N2
c  |         |         |      cell(C,7) = N3
c  |         |         |      cell(C,8) = N4
c  N4-----F1-----N1
c
c
c***** EDGE ARRAY *****
c
c
c  N4-----N2
c  |         |
c  |         |      F(edge)
c  |         |
c  N3-----N1
c
c          eface(EF) = F
c
c*****

```

```

program gridgen
implicit none

include 'QUAD.INC'
include 'GRID.INC'
integer gtype !type of grid
integer EN, EF, FF, NN, CC, col !pointers
integer ON, IN, PN
integer i, j !pointers
integer JJJ

c** determine geometry
type*, 'Type of grid '
type*, ' 1) N1 bump '
type*, ' 2) blade'
type 100
100 format(%, ' selection = ')
accept 110, gtype
110 format(I)

write(6,*) ' '

```

```

        write(6,*) 'Generating grid ...'
        write(6,*) ' '

        if (gtype.eq.1) then
c** create rectangular mesh
            call rectangle
            pitch = 0.

            else if (gtype.eq.2) then
c** Read in, normalize and spline blade data
                call readin

c** Initialize grid
                call grinit

c** Fix up grid
                call ellip(Maxdim,Maxdim,II,JJ,JJJ,XX,YY,YPOS,XPOS)
                call improv

c** change pointers
                do i = 1, ii
                    do j = 1, jj
                        xx(i-1,j-1) = xx(i,j)
                        yy(i-1,j-1) = yy(i,j)
                    enddo
                enddo
                Nx = ii-1
                Ny = jj-1
            endif

c** change rectangular mesh to triangular mesh
                call pointers
                call bpointers

c** rearrange node numbers on edges
                call edgenumber

c** color cells and faces
                call cellcolor
                call facecolor
                call edgcolor

c** write out data to file
                call gridio(0)

                stop
                end

subroutine rectangle
implicit none

include 'GRID.INC'

real*4 yymax, xxmax, xxmin      !coordinates of grid boundaries
real*4 delta_x                  !grid spacing in x direction
real*4 delta_y                  !grid spacing in y direction
integer n, i, j, kk, m         !counters
real*4 num
real*4 delta_xbot
real*4 omega                    !relaxation constant for
                                !interior point SLOR
real*4 pi                       !the one and only
real*4 tau                      !height of bump
real*4 rr                      !radius of bump
real*4 yc                      !rr-tau
real*4 ang, delta_ang          !angle of bump,angle between nodes
real*4 angplus
real*4 delx, dely              !help define initial conditions
real*4 alpha(Maxdim), beta(Maxdim) !coef. in modified equation

```

```

real*4 gamma(Maxdim)
real*4 AA(Maxdim), BB(Maxdim) !values for system of equations
real*4 CC(Maxdim), DD(Maxdim)
real*4 EPS !allowable error
integer Niter !number of iterations so far
real*4 error !largest error for an iteration
real*4 x_xi(Maxdim), x_eta(Maxdim) !derivatives on boundary
real*4 x_xi_xi(Maxdim), x_eta_eta(Maxdim)
real*4 x_eta_xi(Maxdim)
real*4 y_xi(Maxdim), y_eta(Maxdim)
real*4 y_xi_xi(Maxdim), y_eta_eta(Maxdim)
real*4 y_eta_xi(Maxdim)
real*4 theta, AR !angle and aspect ratio of edge cells
real*4 R1, R2 !part of source term
real*4 Q1(Maxdim), P1(Maxdim) !part of source term, function of xi
real*4 a1(Maxdim), b1(Maxdim) !alpha, beta, gamma on lower boundary
real*4 c1(Maxdim)
real*4 omega_P, omega_Q !relaxation conts. for source terms
real*4 Jacobi1(Maxdim) !Jacobian on the lower boundary
real*4 Jacobi2(Maxdim) !Jacobian squared in region
real*4 xxi, yxi, xeta, yeta !derivatives in region
real*4 a, b !exponents in source terms

pi = 3.14159

c** number of nodes on x and y axes
xxmax = 2.
xxmin = -1.
yymin = 1.

call Irequest(' Nx',Nx)
call Irequest(' Ny',Ny)
call Rrequest(' tau',tau)
c call Rrequest(' omega',omega)
c call Rrequest(' omega_P',omega_P)
c call Rrequest(' omega_Q',omega_Q)
c call Rrequest(' a',a)
c call Rrequest(' b',b)
c call Rrequest(' AR',AR)
c call Rrequest(' EPS',EPS)

omega = 1.
a = 0.8
b = 0.8
AR = 0.5
EPS = 0.0005

if (tau.eq.0.) then
  omega_P = 0.
  omega_Q = 0.
else if (tau.lt.0.) then
  AR = 1.
  omega_P = 0.02
  omega_Q = 0.02
else
  omega_P = 0.02
  omega_Q = 0.02
endif

theta = .5*pi
delta_x = (xxmax-xxmin)/real(Nx)

c** set initial and boundary conditions for x and y
if (tau.gt.0.) then
  yc = 0.5*(-tau**2 + 0.25)/tau
  rr = sqrt(0.25 + yc**2)
  ang = asin(.5/rr)
  if (mod(Nx,3).ne.0) then
    num = real(Nx - mod(Nx,3))
    delta_xbot = (xxmax-xxmin)/num
    angplus = 2.*ang/(num/3. + real(mod(Nx,3)))
  else

```

```

        angplus = 2.*ang*delta_x
        delta_xbot = delta_x
    endif
    delta_ang = 0.
    do i = 0, Nx
        yy(i,Ny) = yymax
        xx(i,Ny) = xxmin + i*delta_x
        if (xx(i,Ny).le.0.) then
            xx(i,0) = xxmin + i*delta_xbot
            yy(i,0) = 0.
        else if (xx(i,Ny).gt.0. .and. xx(i,Ny).lt.1.) then
            delta_ang = delta_ang + angplus
            xx(i,0) = 0.5 - rr*sin(ang-delta_ang)
            yy(i,0) = rr*cos(ang-delta_ang) - yc
        else if (xx(i,Ny).ge.1.) then
            xx(i,0) = xxmin + (i-mod(Nx,3))*delta_xbot
            yy(i,0) = 0.
        endif
        delx = (xx(i,Ny) - xx(i,0))/real(Ny)
        dely = (yy(i,Ny) - yy(i,0))/real(Ny)
        do j = 1, Ny-1
            xx(i,j) = delx*j + xx(i,0)
            yy(i,j) = dely*j + yy(i,0)
        enddo
    enddo
else if (tau.lt.0.) then
    delta_xbot = delta_x
    do i = 0, Nx
        yy(i,Ny) = yymax
        xx(i,Ny) = xxmin + i*delta_x
        if (xx(i,Ny).le.0.) then
            xx(i,0) = xxmin + i*delta_xbot
            yy(i,0) = 0.
        else if (xx(i,Ny).gt.0. .and. xx(i,Ny).lt.1.) then
            xx(i,0) = xxmin + i*delta_xbot
            yy(i,0) = -tau*(sin(pi*xx(i,0)))*2
        else if (xx(i,Ny).ge.1.) then
            xx(i,0) = xxmin + i*delta_xbot
            yy(i,0) = 0.
        endif
        delx = (xx(i,Ny) - xx(i,0))/real(Ny)
        dely = (yy(i,Ny) - yy(i,0))/real(Ny)
        do j = 1, Ny-1
            xx(i,j) = delx*j + xx(i,0)
            yy(i,j) = dely*j + yy(i,0)
        enddo
    enddo
else if (tau.eq.0) then
    delta_y = yymax/real(Ny)
    do i = 0, Nx
        do j = 0, Ny
            xx(i,j) = i*delta_x + xxmin
            yy(i,j) = j*delta_y
        enddo
    enddo
endif

c* Solve for source terms
do n = 1, Nx-1
    P1(n) = 0.
    Q1(n) = 0.
    x_xi(n) = .5*(xx(n+1,0) - xx(n-1,0))
    y_xi(n) = .5*(yy(n+1,0) - yy(n-1,0))

    x_xi_xi(n) = (xx(n+1,0) - 2.*xx(n,0) + xx(n-1,0))
    y_xi_xi(n) = (yy(n+1,0) - 2.*yy(n,0) + yy(n-1,0))

    if (xx(n,0).eq.0. .or. xx(n,0).eq.1.) then
        x_eta(n) = max(5.*tau,1.)*AR*(-x_xi(n)*cos(theta)
&          - y_xi(n)*sin(theta))
        y_eta(n) = max(5.*tau,1.)*AR*(-y_xi(n)*cos(theta)
&          + x_xi(n)*sin(theta))
    endif
enddo

```

```

else
    x_eta(n) = AR*(-x_xi(n)*cos(theta) - y_xi(n)*sin(theta))
    y_eta(n) = AR*(-y_xi(n)*cos(theta) + x_xi(n)*sin(theta))
endif

x_eta_xi(n) = 0.5*(x_eta(n+1)-x_eta(n-1))
y_eta_xi(n) = 0.5*(y_eta(n+1)-y_eta(n-1))

Jacobi(n) = x_xi(n)*y_eta(n) - x_eta(n)*y_xi(n)

a1(n) = x_eta(n)**2 + y_eta(n)**2
b1(n) = x_xi(n)*x_eta(n) + y_xi(n)*y_eta(n)
c1(n) = x_xi(n)**2 + y_xi(n)**2
enddo

c* SOR by lines
Niter = 0
error = 9999.
do while (error.gt.EPS)
    error = 0.
    Niter = Niter + 1

c** solve for source terms
    if (tau.ne.0.) then
        do n = 1, Nx-1
            x_eta_eta(n) = 0.5*(-7.*xx(n,0) + 8.*xx(n,1) - xx(n,2))
            & - 3.*x_eta(n)
            y_eta_eta(n) = 0.5*(-7.*yy(n,0) + 8.*yy(n,1) - yy(n,2))
            & - 3.*y_eta(n)
            R1 = (-a1(n)*x_xi_xi(n) + 2.*b1(n)*x_eta_xi(n)
            & - c1(n)*x_eta_eta(n))/Jacobi(n)**2
            R2 = (-a1(n)*y_xi_xi(n) + 2.*b1(n)*y_eta_xi(n)
            & - c1(n)*y_eta_eta(n))/Jacobi(n)**2
            P1(n) = P1(n) + omega_P*((y_eta(n)*R1 - x_eta(n)*R2)
            & /Jacobi(n) - P1(n))
            Q1(n) = Q1(n) + omega_Q*((-y_xi(n)*R1 + x_xi(n)*R2)
            & /Jacobi(n) - Q1(n))
        enddo
    endif

c* solve for each line
    do kk = 1, Ny-1

c* evaluate alpha, beta, and gamma
        do i = 1, Nx-1
            alpha(i) = .25*(xx(i,kk+1)-xx(i,kk-1))**2
            & + .25*(yy(i,kk+1)-yy(i,kk-1))**2
            beta(i) = .25*(xx(i+1,kk)-xx(i-1,kk))*(xx(i,kk+1)
            & -xx(i,kk-1)) + .25*(yy(i+1,kk)-yy(i-1,kk))
            & *(yy(i,kk+1)-yy(i,kk-1))
            gamma(i) = .25*(xx(i+1,kk)-xx(i-1,kk))**2
            & + .25*(yy(i+1,kk)-yy(i-1,kk))**2
            Jacobi2(i) = (.25*(xx(i+1,kk)-xx(i-1,kk))*
            & (yy(i,kk+1)-yy(i,kk-1))
            & - .25*(yy(i+1,kk)-yy(i-1,kk))*
            & (xx(i,kk+1)-xx(i,kk-1)))**2
        enddo

c* Solve for x
c* set up matrix for tridiagonal system of equations
        do i = 1, Nx-1
            AA(i) = omega*alpha(i)
            DD(i) = -2.*(gamma(i) + alpha(i))
            BB(i) = omega*alpha(i)
        enddo

c* set up vector of constants for tridiagonal system of equations
        do i = 1, Nx-1
            xxi = xx(i+1,kk) - xx(i-1,kk)
            xeta = xx(i,kk+1) - xx(i,kk-1)

            CC(i) = -omega*gamma(i)*(xx(i,kk+1) + xx(i,kk-1))

```



```

&          + omega*0.5*beta(i)*(xx(i+1,kk+1)
&          - xx(i+1,kk-1) - xx(i-1,kk+1) + xx(i-1,kk-1))
&          + 2.*(omega-1.)*(alpha(i) + gamma(i))*xx(i,kk)
&          - Jacobi2(i)*(P1(i)*xxi*exp(-a*kk) +
&          Q1(i)*xeta*exp(-b*kk))
      enddo
      CC(1) = CC(1) - BB(1)*xx(0,kk)
      CC(Nx-1) = CC(Nx-1) - AA(Nx-1)*xx(Nx,kk)

c* solve tridiagonal system of equations
call tridiag(1, Nx-1, BB, DD, AA, CC)
do m = 1, Nx-1
  if (abs(CC(m)-xx(m,kk)).gt.error) then
    error = abs(CC(m) - xx(m,kk))
  endif
  xx(m,kk) = CC(m)
enddo

c* Solve for y
c* set up matrix for tridiagonal system of equations
do i = 1, Nx-1
  AA(i) = omega*alpha(i)
  DD(i) = -2.*(gamma(i) + alpha(i))
  BB(i) = omega*alpha(i)
enddo

c* set up vector of constants for tridiagonal system of equations
do i = 1, Nx-1
  yxi = yy(i+1,kk) - yy(i-1,kk)
  yeta = yy(i,kk+1) - yy(i,kk-1)

  CC(i) = -omega*gamma(i)*(yy(i,kk+1) + yy(i,kk-1))
&          + 0.5*omega*beta(i)*(yy(i+1,kk+1)
&          -yy(i+1,kk-1) - yy(i-1,kk+1) + yy(i-1,kk-1))
&          + 2.*(omega-1.)*(alpha(i)+gamma(i))*yy(i,kk)
&          - Jacobi2(i)*(P1(i)*yxi*exp(-a*kk) +
&          Q1(i)*yeta*exp(-b*kk))
      enddo
      CC(1) = CC(1) - BB(1)*yy(0,kk)
      CC(Nx-1) = CC(Nx-1) - AA(Nx-1)*yy(Nx,kk)

c* solve tridiagonal system of equations
call tridiag(1, Nx-1, BB, DD, AA, CC)
do m = 1, Nx-1
  if (abs(CC(m)-yy(m,kk)).gt.error) then
    error = abs(CC(m) - yy(m,kk))
  endif
  yy(m,kk) = CC(m)
enddo

enddo

c* set x=xxmax and x=xxmin boundary conditions to next interior point
do kk = 1, Ny-1
  yy(0,kk) = yy(1,kk)
  yy(Nx,kk) = yy(Nx-1,kk)
enddo

type*, 'iteration number = ',Niter,' error = ',error
enddo

c** write out data to data file
C   write(10,*) Nx+1, Ny+1
C   do i = 0, Nx
C     do j = 0, Ny
C       write(10,*) xx(i,j), yy(i,j)
C     enddo
C   enddo
C   close(unit=10)

c** plot the grid
C   call plot

```

```

return
end

```

```

C*****
C*
C*      Subroutine to solve a tridiagonal system of equations.      *
C*      Taken from "Computational Fluid Mechanics and Heat Transfer" *
C*      by Anderson, Tannehill and Pletcher.                       *
C*                                                                    *
C*****

```

```

SUBROUTINE TRIDIAG(IL,IU,BB,DD,AA,RR)
IMPLICIT NONE

```

```

include 'GRID.INC'

```

```

INTEGER IL                      !SUBSCRIPT OF FIRST EQUATION
INTEGER IU                      !SUBSCRIPT OF LAST EQUATION
REAL*4 BB(Maxdim)              !COEFFICIENT BEHIND DIAGONAL
REAL*4 DD(Maxdim)              !COEFFICIENT ON DIAGONAL
REAL*4 AA(Maxdim)              !COEFFICIENT AHEAD OF DIAGONAL
REAL*4 RR(Maxdim)              !ELEMENT OF CONSTANT VECTOR
INTEGER LP
INTEGER I, J                    !POINTERS
REAL*4 R

```

```

C**      ESTABLISH UPPER TRIANGULAR MATRIX

```

```

LP = IL + 1
DO I = LP, IU
    R = BB(I)/DD(I-1)
    DD(I) = DD(I)-R*AA(I-1)
    RR(I) = RR(I)-R*RR(I-1)
ENDDO

```

```

C**      BACK SUBSTITUTION

```

```

RR(IU) = RR(IU)/DD(IU)
DO I = LP, IU
    J = IU - I + 1
    RR(J) = (RR(J)-AA(J)*RR(J+1))/DD(J)
ENDDO

```

```

C**      SOLUTION STORED IN RR

```

```

RETURN
END

```

```

C*****
C*
C*      this subroutine requests the user to input the value of a    *
C*      real variable                                                 *
C*                                                                    *
C*****

```

```

subroutine Rrequest (name,var)
character*10 name
real*4 var

```

```

1  write(6,1) name
   format(9,' ',A,' = ')
   accept 11, var
11 format(F)

```

```

return

```

```

end

C*****
C*
C*   this subroutine requests the user to input the value of a   *
C*   integer variable                                           *
C*                                                                 *
C*****

      subroutine Irequest (name,var)
      character*10 name
      integer var

      write(6,1) name
1      format(9,' ',A,' = ')
      accept 11, var
11     format(I)

      return
      end

SUBROUTINE READIN

C---- Read in, normalize and spline blade data

      INCLUDE 'QUAD.INC'
      INCLUDE 'GRID.INC'

      CHARACTER*32 NAMEXT
      CHARACTER*80 NAME

C---- Read in blade data
      OPEN(UNIT=3,STATUS='OLD')
1000 FORMAT(A32)
      READ(3,1000) NAME
      READ(3,*) G SINL, GSOUT, CHINL, CHOUT, PITCH

      WRITE(6,1001) NAME
1001 FORMAT(/,' Blade name: ',A60)

      READ(3,*) XB(1), YB(1)
      XMIN = XB(1)
      XMAX = XB(1)
      YMIN = YB(1)
      DO 1 IB = 2, 12345
      READ(3,*,END=11) XB(IB),YB(IB)
      XMAX = AMAX1(XMAX,XB(IB))
      IF(XMIN.GT.XB(IB)) THEN
      XMIN = XB(IB)
      YMIN = YB(IB)
      ENDIF
1      CONTINUE
11     IIB = IB - 1
      CLOSE(UNIT=3)

      IF(IIB.GT.IBX) STOP 'Array overflow: IBX too small'

C---- Normalize blade and calculate surface arc length array
      PITCH = PITCH/(XMAX-XMIN)
      DO 2 IB = 1, IIB
      XB(IB) = (XB(IB)-XMIN) / (XMAX-XMIN)
      YB(IB) = (YB(IB)-YMIN) / (XMAX-XMIN)
2      CONTINUE

C---- close t.e. if open
      IF( XB(1).NE.XB(IIB) .OR. YB(1).NE.YB(IIB) ) THEN
      ABOUT = ATAN( (YB(2) -YB(1)) / (XB(2) -XB(1)) )

```

```

APOUT = ATAN( (YB(IIB)-YB(IIB-1)) / (XB(IIB)-XB(IIB-1)) )
DSTE = SQRT( (XB(1)-XB(IIB))**2 + (YB(1)-YB(IIB))**2 )
AOUT = 0.5*(APOUT+ASOUT)
XOUT = 0.5*(XB(1)+XB(IIB)) + 3.0*DSTE*COS(AOUT)
YOUT = 0.5*(YB(1)+YB(IIB)) + 3.0*DSTE*SIN(AOUT)
XS1 = (XB(1)-XOUT)*COS(AOUT) + (YB(1)-YOUT)*SIN(AOUT)
YS1 = -(XB(1)-XOUT)*SIN(AOUT) + (YB(1)-YOUT)*COS(AOUT)
YPS1 = TAN(ASOUT-AOUT) * XS1
XP1 = (XB(IIB)-XOUT)*COS(AOUT) + (YB(IIB)-YOUT)*SIN(AOUT)
YP1 = -(XB(IIB)-XOUT)*SIN(AOUT) + (YB(IIB)-YOUT)*COS(AOUT)
YPP1 = TAN(APOUT-AOUT) * XP1

WRITE(6,1002)
1002 FORMAT(/,' Input flap deflection angle (degrees): ',%)
READ(6,*) AFLAP
YPFLAP = TAN(3.14159*AFLAP/180.0)

WRITE(NAMEXT,1003) AFLAP
1003 FORMAT(' (flap deflection angle = ',F4.1,')')
LENSTART = INDEX(NAME,')
NAME(LENSTART:LENSTART+31) = NAMEXT

IIB = IIB+20
IF(IIB.GT.IBX) STOP 'Array overflow: IBX too small'

DO 3 IB = IIB-20, 1, -1
  XB(IB+10) = XB(IB)
  YB(IB+10) = YB(IB)
3 CONTINUE

DO 4 IB = 1, 10
  ETA = 0.1*(IB-1)
  XXS = XS1*ETA
  YYS = YS1*ETA*ETA*(3.0-2.0*ETA) + YPS1*ETA*ETA*(ETA-1.0)
  * - YPFLAP*XS1*0.5*(ETA-1.0)**2
  XB(IB) = XOUT + XXS*COS(AOUT) - YYS*SIN(AOUT)
  YB(IB) = YOUT + XXS*SIN(AOUT) + YYS*COS(AOUT)
4 CONTINUE

DO 5 IB = IIB-9, IIB
  ETA = 0.1*(IIB-IB)
  XXP = XP1*ETA
  YYP = YP1*ETA*ETA*(3.0-2.0*ETA) + YPP1*ETA*ETA*(ETA-1.0)
  * - YPFLAP*XP1*0.5*(ETA-1.0)**2
  XB(IB) = XOUT + XXP*COS(AOUT) - YYP*SIN(AOUT)
  YB(IB) = YOUT + XXP*SIN(AOUT) + YYP*COS(AOUT)
5 CONTINUE

ENDIF

C---- Spline blade surface(s) and find leading edge position
SB(1) = 0.
DO 6 IB = 2, IIB
  ALF = FLOAT( MIN(IB-1,IIB-IB) ) / FLOAT(IIB/2)
  SB(IB) = SB(IB-1) +
  * SQRT( (XB(IB)-XB(IB-1))**2 + (ALF*(YB(IB)-YB(IB-1)))**2 )
6 CONTINUE

CALL SPLINE(XB,XPB,SB,IIB)
CALL SPLINE(YB,YPB,SB,IIB)

DO 7 IB=2, IIB
  DP1 = XPB(IB-1) + GSINL*YPB(IB-1)
  DP2 = XPB(IB) + GSINL*YPB(IB)
  IF(DP1.LT.0.0 .AND. DP2.GE.0.0) GO TO 71
7 CONTINUE

STOP 'Leading edge not found'
71 DSB = SB(IB) - SB(IB-1)
SBLE = SB(IB-1) + DSB*DP1/(DP1-DP2)
XLE = SEVAL(SBLE,XB,XPB,SB,IIB)
YLE = SEVAL(SBLE,YB,YPB,SB,IIB)

```

```

RETURN
END

```

SUBROUTINE GRINIT

C---- Fix grid points on boundary of domain, and initialize interior

```

INCLUDE 'QUAD.INC'
INCLUDE 'GRID.INC'

```

C---- Input and check grid size

```

WRITE(6,1000)
1000 FORMAT(/,' Input II, JJ: ',%)
READ(6,*) II,JJ
IF(II.GT.Maxdim) STOP 'Array overflow: Maxdim too small'
IF(JJ.GT.Maxdim) STOP 'Array overflow: Maxdim too small'

```

C---- Set various parameters

```

SLEN = CHINL + 0.5*SB(IIB) + CHOUT
NINL = INT( FLOAT(II)*CHINL/SLEN )
NOUT = INT( FLOAT(II)*CHOUT/SLEN )
NBLD = II - NOUT - NINL + 2
ILE = NINL
ITE = II - NOUT + 1

```

C---- Set inlet stagnation streamline

```

DO 1 K=1, NINL
  XX(K,1) = XLE + CHINL * FLOAT(K-NINL) / FLOAT(NINL-1)
  YY(K,1) = YLE + (XX(K,1)-XLE) * GSINL
  XX(K,JJ) = XX(K,1)
  YY(K,JJ) = YY(K,1) + PITCH
1 CONTINUE

```

C---- Set outlet stagnation streamline

```

XTE = XB(1)
YTE = YB(1)

DO 2 K= 1, NOUT
  I = II-NOUT+K
  XX(I,1) = XTE + CHOUT * FLOAT(K-1) / FLOAT(NOUT-1)
  YY(I,1) = YTE + (XX(I,1)-XTE) * GSOUT
  XX(I,JJ) = XX(I,1)
  YY(I,JJ) = YY(I,1) + PITCH
2 CONTINUE

```

C---- Set points on blade suction surface

```

DO 3 K=1, NBLD
  I = NINL + K - 1
  S = SBLE - SBLE*FLOAT(K-1)/FLOAT(NBLD-1)
  XX(I,1) = SEVAL(S,XB,XPB,SB,IIB)
  YY(I,1) = SEVAL(S,YB,YPB,SB,IIB)
3 CONTINUE

```

C---- Set points on blade pressure surface

```

DO 4 K=1, NBLD
  I = NINL + K - 1
  S = SBLE + (SB(IIB)-SBLE)*FLOAT(K-1)/FLOAT(NBLD-1)
  XX(I,JJ) = SEVAL(S,XB,XPB,SB,IIB)
  YY(I,JJ) = SEVAL(S,YB,YPB,SB,IIB) + PITCH
4 CONTINUE

```

C---- set up metrics

```

DO 5 I=1, II
  XPOS(I) = FLOAT(I-1)/FLOAT(II-1)
5 CONTINUE

DO 6 J = 1, JJ

```

```

      RJ = FLOAT(J-1)/FLOAT(JJ-1)
      YPOS(J) = RJ - 1.8 * ( (RJ-0.5) * ((RJ-0.5)**2-0.25) )
6      CONTINUE

C---- Initialize interior grid
DO 7 I = 1, II
  DO 71 J = 2, JJ-1
    XX(I,J) = XX(I,1) + YPOS(J)*(XX(I,JJ)-XX(I,1))
    YY(I,J) = YY(I,1) + YPOS(J)*(YY(I,JJ)-YY(I,1))
71  CONTINUE
7  CONTINUE

RETURN
END

```

SUBROUTINE IMPROV

C---- Improves grid after elliptic grid generation

```

INCLUDE 'QUAD.INC'
INCLUDE 'GRID.INC'

DIMENSION SUM(Maxdim), XT1(Maxdim), YT1(Maxdim)
DIMENSION XT2(Maxdim), YT2(Maxdim)

DO 1 I = 1, II
  IM = I-1
  IP = I+1
  IF(I.EQ.1) IM=1
  IF(I.EQ.II) IP=II

  SUM(1) = 0.
  DO 11 J = 1, JJ-1
    JP = J+1
    XS = XX(IP,J)+XX(IP,JP) - XX(IM,J)-XX(IM,JP)
    YS = YY(IP,J)+YY(IP,JP) - YY(IM,J)-YY(IM,JP)
    SS = SQRT(XS*XS + YS*YS)
    XS = XS/SS
    YS = YS/SS
    SUM(JP) = SUM(J) + ABS( (XX(I,J)-XX(I,JP))*YS
      * - (YY(I,J)-YY(I,JP))*XS )
11  CONTINUE

  J = 1
  DO 12 JO = 2, JJ-1
    SUMJ = FLOAT(JO-1)/FLOAT(JJ-1) * SUM(JJ)
121  IF(SUMJ.GT.SUM(J+1)) THEN
    J = J+1
    GOTO 121
  ENDIF
  ALPHA = (SUMJ-SUM(J)) / (SUM(J+1)-SUM(J))
  XT2(JO) = XX(I,J) + ALPHA*(XX(I,J+1)-XX(I,J))
  YT2(JO) = YY(I,J) + ALPHA*(YY(I,J+1)-YY(I,J))
12  CONTINUE

  DO 13 J = 2, JJ-1
    IF(I.NE.1) THEN
      XX(IM,J) = XT1(J)
      YY(IM,J) = YT1(J)
    ENDIF
    XT1(J) = XT2(J)
    YT1(J) = YT2(J)
    IF(I.EQ.II) THEN
      XX(I,J) = XT1(J)
      YY(I,J) = YT1(J)
    ENDIF
  ENDIF

```

13 CONTINUE

1 CONTINUE
 RETURN
 END

```

C*****C
C                                     C
C   ISES - an Integrated Steamtube Euler Solver   C
C                                     C
C   Written by M. Giles and M.Drela               C
C                                     C
C   Copyright M.I.T. (1985)                       C
C*****C
C
C   SUBROUTINE ELLIP(IMAX,JMAX,II,JJ,JJJ,X,Y,YPOS,XPOS)
C   DIMENSION X(0:IMAX,0:JMAX), Y(0:IMAX,0:JMAX)
C   DIMENSION YPOS(JMAX), XPOS(IMAX)
C   CHARACTER*1 ANS
C
C   DIMENSION C(400),D(2,400)
C   IF(II.GT.400) STOP 'ELLIP dimensions must be increased'
C
C   ITMAX = 50
C
C   DSET1 = 1.0E-1
C   DSET2 = 5.0E-3
C   DSET3 = 2.0E-4
C
C   RLX1 = 1.30      !           DMAX > DSET1
C   RLX2 = 1.50      !   DSET1 > DMAX > DSET2
C   RLX3 = 1.60      !   DSET2 > DMAX > DSET3
CCC  STOP           !   DSET3 > DMAX
C
C   RLX = RLX1
C
C   DO 1 ITER = 1, ITMAX
C
C       DMAX = 0.
C       DO 5 JO=2, JJ-1
C           JM = JO-1
C           JP = JO+1
C
C           IF(JO.EQ.JJJ) THEN
C               DO 2 IO=2, II-1
C                   X(IO,JO) = X(IO,JM)
C               CONTINUE
C               GO TO 6
C           ELSE IF(JO.EQ.JJJ+1) THEN
C               DO 3 IO=2, II-1
C                   X(IO,JO) = X(IO,JP)
C               CONTINUE
C               GO TO 6
C           ENDIF
C
C       DO 6 IO=2, II-1
C           IM = IO-1
C           IP = IO+1
C
C           XMM = X(IM,JM)
C           XOM = X(IO,JM)
C           XPM = X(IP,JM)
C           XMO = X(IM,JO)
C           XOO = X(IO,JO)
C           XPO = X(IP,JO)
C           XMP = X(IM,JP)
C           XOP = X(IO,JP)

```

```

XPP = X(IP,JP)
YMM = Y(IM,JM)
YQM = Y(IO,JM)
YPM = Y(IP,JM)
YMO = Y(IM,JO)
YOO = Y(IO,JO)
YPO = Y(IP,JO)
YMP = Y(IM,JP)
YOP = Y(IO,JP)
YPP = Y(IP,JP)

C
DXIM = XPOS(IO)-XPOS(IM)
DXIP = XPOS(IP)-XPOS(IO)
DXIAV = 0.5*(DXIM+DXIP)

C
DETM = YPOS(JO)-YPOS(JM)
DETP = YPOS(JP)-YPOS(JO)
DETAV = 0.5*(DETM+DETP)

C
DXDET = ( XOP - XOM ) / DETAV
DYDET = ( YOP - YOM ) / DETAV
DXDXI = ( XPO - XMO ) / DXIAV
DYDXI = ( YPO - YMO ) / DXIAV

C
ALF = DXDET**2 + DYDET**2
BET = DXDET*DXDXI + DYDET*DYDXI
GAM = DXDXI**2 + DYDXI**2

C
CXIM = 1.0 / (DXIM*DXIAV)
CXIP = 1.0 / (DXIP*DXIAV)
CETM = 1.0 / (DETM*DETAV)
CETP = 1.0 / (DETP*DETAV)

C
B = -ALF*CXIM
A = ALF*(CXIM+CXIP) + GAM*(CETM+CETP)
C(IO) = -ALF*CXIP
IF(IO.EQ.2) B = 0

C
* D(1,IO) = ALF*((XMO-XOO)*CXIM + (XPO-XOO)*CXIP)
* - 2.0*BET*(XPP-XMP-XPM+XMM) / (4.0*DXIAV*DETAV)
* + GAM*((XQM-XOO)*CETM + (XOP-XOO)*CETP)

C
* D(2,IO) = ALF*((YMO-YOO)*CXIM + (YPO-YOO)*CXIP)
* - 2.0*BET*(YPP-YMP-YPM+YMM) / (4.0*DXIAV*DETAV)
* + GAM*((YQM-YOO)*CETM + (YOP-YOO)*CETP)

C
AINV = 1.0/(A - B*C(IM))
C(IO) = C(IO) * AINV
D(1,IO) = ( D(1,IO) - B*D(1,IM) ) * AINV
D(2,IO) = ( D(2,IO) - B*D(2,IM) ) * AINV

C
6 CONTINUE

C
D(1,II) = 0.
D(2,II) = 0.

C
IFIN = II-1
DO 8 IBACK=2, IFIN
  IO = II-IBACK+1
  IP = IO+1
  D( 1,IO) = D( 1,IO) - C(IO)*D(1,IP)
  D( 2,IO) = D( 2,IO) - C(IO)*D(2,IP)
  X(IO,JO) = X(IO,JO) + RLX*D(1,IO)
  Y(IO,JO) = Y(IO,JO) + RLX*D(2,IO)
  AD1 = ABS(D(1,IO))
  AD2 = ABS(D(2,IO))
  DMAX = AMAX1(DMAX,AD1,AD2)
8 CONTINUE

C
5 CONTINUE

C
WRITE(8,*) 'Dmax = ', DMAX, RLX

```



```

C      RLX = RLX1
      IF(DMAX.LT.DSET1) RLX = RLX2
      IF(DMAX.LT.DSET2) RLX = RLX3
      IF(DMAX.LT.DSET3) RETURN
C
C 1 CONTINUE
C
      RETURN
      END ! ELLIP

      SUBROUTINE SPLINE(X,XP,S,II)
C
      DIMENSION X(1),XP(1),S(1)
      DIMENSION A(480),B(480),C(480)
C
      IF(II.GT.480) STOP 'SPLINE: Array overflow'
      DO 1 I = 1, II
C
C----- Beginning points
      IF(I.EQ.1 .OR. S(I).EQ.S(I-1)) THEN
          DSMI = 0.
          DXM = 0.
          DSPI = 1.0 / (S(I+1)-S(I))
          DXP = X(I+1) - X(I)
C
C----- End points
      ELSE IF(I.EQ.II .OR. S(I).EQ.S(I+1)) THEN
          DSMI = 1.0 / (S(I) - S(I-1))
          DXM = X(I) - X(I-1)
          DSPI = 0.
          DXP = 0.
C
C----- Interior points
      ELSE
          DSMI = 1.0 / (S(I) - S(I-1))
          DXM = X(I) - X(I-1)
          DSPI = 1.0 / (S(I+1) - S(I))
          DXP = X(I+1) - X(I)
C
      ENDIF
C
      B(I) = DSMI
      A(I) = 2.0 * (DSMI + DSPI)
      C(I) = DSPI
      XP(I) = 3.0 * (DXP*DSPI**2 + DXM*DSMI**2)
C
C 1 CONTINUE
C
      CALL TRISOL(A,B,C,XP,II)
C
      RETURN
      END ! SPLINE

      SUBROUTINE TRISOL(A,B,C,D,KK)
C
      DIMENSION A(1),B(1),C(1),D(1)
C
      DO 1 K = 2, KK
          KM = K - 1
          C(KM) = C(KM) / A(KM)
          D(KM) = D(KM) / A(KM)
          A(K) = A(K) - B(K) * C(KM)

```

```

      D(K) = D(K) - E(K) * D(KM)
1  CONTINUE
C
      D(KK) = D(KK) / A(KK)
C
      DO 2 K = KK-1, 1, -1
        D(K) = D(K) - C(K) * D(K+1)
2  CONTINUE
C
      RETURN
END ! TRISOL

```

```

      FUNCTION SEVAL(SS,X,XP,S,N)
      REAL X(1), XP(1), S(1)
C
      ILOW = 1
      I = N
C
10 IF(I-ILOW .LE. 1) GO TO 11
C
      IMID = (I+ILOW)/2
      IF(SS .LT. S(IMID)) THEN
        I = IMID
      ELSE
        ILOW = IMID
      ENDIF
      GO TO 10
C
11 DS = S(I) - S(I-1)
      T = (SS-S(I-1)) / DS
      CX1 = DS*XP(I-1) - X(I) + X(I-1)
      CX2 = DS*XP(I) - X(I) + X(I-1)
      SEVAL = T*X(I) + (1.0-T)*X(I-1) + (T-T*T)*((1.0-T)*CX1 - T*CX2)
      RETURN
END ! SEVAL

```

```

      FUNCTION DEVAL(SS,X,XP,S,N)
      REAL X(1), XP(1), S(1)
C
      ILOW = 1
      I = N
C
10 IF(I-ILOW .LE. 1) GO TO 11
C
      IMID = (I+ILOW)/2
      IF(SS .LT. S(IMID)) THEN
        I = IMID
      ELSE
        ILOW = IMID
      ENDIF
      GO TO 10
C
11 DS = S(I) - S(I-1)
      T = (SS-S(I-1)) / DS
      CX1 = DS*XP(I-1) - X(I) + X(I-1)
      CX2 = DS*XP(I) - X(I) + X(I-1)
      DEVAL = (X(I)-X(I-1) + (1.-4.*T+3.*T*T)*CX1 + T*(3.*T-2.)*CX2)/DS
      RETURN
END ! DEVAL

```

```

subroutine pointers
implicit none

include 'GRID.INC'
include 'QUAD.INC'
integer i, j                !pointers
integer NN, FF, CC          !pointers

NN = 0
CC = 0
FF = 0

c** assign values to node arrays
do j = 0, Ny
  do i = 0, Nx
    NN = NN + 1
    x(NN) = xx(i,j)
    y(NN) = yy(i,j)
  enddo
enddo
Nmax = NN
if (Nmax .gt. Maxnodes) then
  write(6,*) 'NN = ', NN
  write(6,*) 'more than Maxnodes required'
endif

c** assign horizontal and vertical faces and cells
do j = 0, Ny-1
  do i = 1, Nx
    FF = FF + 1
    face(FF,1) = Nx*j + 1
    face(FF,2) = Nx*(j-1) + 1
    face(FF,3) = (Nx+1)*j + 1
    face(FF,4) = (Nx+1)*j + 1 + 1
    face(FF,5) = (Nx+1)*(j+1) + 1
    face(FF,6) = (Nx+1)*(j+1) + 1 + 1
    face(FF,7) = (Nx+1)*(j-1) + 1 + 1
    face(FF,8) = (Nx+1)*(j-1) + 1
    CC = CC + 1
    cell(CC,1) = FF
    cell(CC,2) = FF + Nx + 1
    cell(CC,3) = FF + 2*Nx + 1
    cell(CC,4) = FF + Nx
    cell(CC,5) = (Nx+1)*j + 1 + 1
    cell(CC,6) = (Nx+1)*(j+1) + 1 + 1
    cell(CC,7) = (Nx+1)*(j+1) + 1
    cell(CC,8) = (Nx+1)*j + 1
  enddo
  do i = 1, Nx
    FF = FF + 1
    face(FF,1) = Nx*j + 1
    face(FF,2) = Nx*j - 1 + 1
    face(FF,3) = (Nx+1)*(j+1) + 1
    face(FF,4) = (Nx+1)*j + 1
    face(FF,5) = (Nx+1)*(j+1) + 1 + 1
    face(FF,6) = (Nx+1)*j + 1 + 1
    face(FF,7) = (Nx+1)*j - 1 + 1
    face(FF,8) = (Nx+1)*(j+1) - 1 + 1
  enddo
  FF = FF + 1
  face(FF,1) = 0
  face(FF,2) = Nx*(j+1)
  face(FF,3) = (Nx+1)*(j+2)
  face(FF,4) = (Nx+1)*(j+1)
  face(FF,5) = 0
  face(FF,6) = 0
  face(FF,7) = (Nx+1)*j + Nx
  face(FF,8) = (Nx+1)*(j+1) + Nx
enddo

```

```

      Cmax = CC
      if (Cmax .gt. Maxcells) then
        write(6,*) 'CC = ', CC
        write(6,*) 'more than Maxcells required'
      endif

c** assign top boundary faces
      do i = 1, Nx
        FF = FF + 1
        face(FF,1) = 0
        face(FF,2) = Nx*(Ny-1) + 1
        face(FF,3) = (Nx+1)*Ny + 1
        face(FF,4) = (Nx+1)*Ny + 1 + 1
        face(FF,5) = 0
        face(FF,6) = 0
        face(FF,7) = (Nx+1)*(Ny-1) + 1 + 1
        face(FF,8) = (Nx+1)*(Ny-1) + 1
      enddo

      Fmax = FF
      if (Fmax .gt. Maxfaces) then
        write(6,*) 'FF = ', FF
        write(6,*) 'more than Maxfaces required'
      endif

c** set bottom edge face array
      do i = 1, Nx
        face(i,2) = 0
        face(i,7) = 0
        face(i,8) = 0
      enddo

c** set inlet edge face array
      do j = 1, Ny
        face((2*Nx+1)*j-Nx,2) = 0
        face((2*Nx+1)*j-Nx,7) = 0
        face((2*Nx+1)*j-Nx,8) = 0
      enddo

c** set outlet edge face array
      do j = 1, Ny
        face((2*Nx+1)*j,1) = 0
        face((2*Nx+1)*j,5) = 0
        face((2*Nx+1)*j,6) = 0
      enddo

      return
      end

subroutine bpointers
implicit none

include 'GRID.INC'
include 'QUAD.INC'
integer i, j
integer n
integer EN, EF, IN, ON, WN, PN
integer CC, FF
integer F1, F2
real dxM, dyM, dxP, dyP
real dsM, dsP, dx, dy
integer node

!pointers
!pointer
!pointers
!pointers
!periodic faces
!change in x & y near node
!current node processed

EN = 0
EF = 0
IN = 0
ON = 0
WN = 0

```

```

PN = 0

c** wall boundaries
if (pitch.eq.0.) then
  do i = 1, Nx
    EF = EF + 1
    eface(EF) = Ny*(2*Nx+1) + 1
    EN = EN + 1
    node = (Nx+1)*Ny + 1
    enode(EN) = node
    dxP = x(node+1) - x(node)
    dyP = y(node+1) - y(node)
    if (i .eq. 1) then
      dxM = -dxP
      dyM = -dyP
    else
      dxM = x(node-1) - x(node)
      dyM = y(node-1) - y(node)
    endif
    if ((dxM*dxP + dyM*dyP) .gt. 0.) then
      dsM = sqrt(dxM**2 + dyM**2)
      dsP = sqrt(dxP**2 + dyP**2)
      dx = dxM/dsM + dxP/dsP
      dy = dyM/dsM + dyP/dsP
    else
      dx = dxP - dxM
      dy = dyP - dyM
    endif
    senode(EN) = dy/sqrt(dx**2 + dy**2)
    cenode(EN) = dx/sqrt(dx**2 + dy**2)
  enddo

  EN = EN + 1
  node = (Nx+1)*(Ny+1)
  enode(EN) = node
  dxM = x(node-1) - x(node)
  dyM = y(node-1) - y(node)
  dxP = -dxM
  dyP = -dyM
  dx = dxP - dxM
  dy = dyP - dyM
  senode(EN) = dy/sqrt(dx**2 + dy**2)
  cenode(EN) = dx/sqrt(dx**2 + dy**2)

  tnode = EN

  do i = 1, Nx
    EF = EF + 1
    eface(EF) = 1
    EN = EN + 1
    node = 1
    enode(EN) = node
    dxP = x(node+1) - x(node)
    dyP = y(node+1) - y(node)
    if (i .eq. 1) then
      dxM = -dxP
      dyM = -dyP
    else
      dxM = x(node-1) - x(node)
      dyM = y(node-1) - y(node)
    endif
    if ((dxM*dxP + dyM*dyP) .gt. 0.) then
      dsM = sqrt(dxM**2 + dyM**2)
      dsP = sqrt(dxP**2 + dyP**2)
      dx = dxM/dsM + dxP/dsP
      dy = dyM/dsM + dyP/dsP
    else
      dx = dxP - dxM
      dy = dyP - dyM
    endif
    senode(EN) = dy/sqrt(dx**2 + dy**2)
    cenode(EN) = dx/sqrt(dx**2 + dy**2)
  enddo

```

```

        enddo

        EN = EN + 1
        node = Nx + 1
        enode(EN) = node
        dxM = x(node-1) - x(node)
        dyM = y(node-1) - y(node)
        dxP = -dxM
        dyP = -dyM
        dx = dxP - dxM
        dy = dyP - dyM
        senode(EN) = dy/sqrt(dx**2 + dy**2)
        cenode(EN) = dx/sqrt(dx**2 + dy**2)

        bnode = EN
        Wmax = 2*Nx
    else
c** periodic nodes
        do i = 1, ILE
            PN = PN + 1
            pnode(PN,1) = 1
            pnode(PN,2) = (Nx+1)*Ny + 1
        enddo

        do i = ITE, Nx+1
            PN = PN + 1
            pnode(PN,1) = 1
            pnode(PN,2) = (Nx+1)*Ny + 1
        enddo
        Pmax = ILE + Nx + 2 - ITE

        do i = 1, ILE-1
            F1 = 1
            F2 = Ny*(2*Nx+1) + 1
            face(F1,2) = face(F2,2)
            face(F1,7) = face(F2,7)
            face(F1,8) = face(F2,8)
            face(F2,1) = face(F1,1)
            face(F2,5) = face(F1,5)
            face(F2,6) = face(F1,6)
        enddo

        do i = ITE, Nx
            F1 = 1
            F2 = Ny*(2*Nx+1) + 1
            face(F1,2) = face(F2,2)
            face(F1,7) = face(F2,7)
            face(F1,8) = face(F2,8)
            face(F2,1) = face(F1,1)
            face(F2,5) = face(F1,5)
            face(F2,6) = face(F1,6)
        enddo

c** the rest of the edge nodes and faces
        do i = ILE, ITE-1
            EN = EN + 1
            node = (Nx+1)*Ny + 1
            enode(EN) = node
            dxP = x(node+1) - x(node)
            dyP = y(node+1) - y(node)
            if (i .eq. ILE) then
                dxM = x(ILE+1) - x(node)
                dyM = y(ILE+1) + pitch - y(node)
            else
                dxM = x(node-1) - x(node)
                dyM = y(node-1) - y(node)
            endif
            if ((dxM*dxP + dyM*dyP) .gt. 0.) then
                dsM = sqrt(dxM**2 + dyM**2)
                dsP = sqrt(dxP**2 + dyP**2)
                dx = dxM/dsM + dxP/dsP
                dy = dyM/dsM + dyP/dsP
            else
                dx = dxM
                dy = dyM
            endif
        enddo
    enddo

```

```

else
  dx = dxP - dxM
  dy = dyP - dyM
endif
senode(EN) = dy/sqrt(dx**2 + dy**2)
cenode(EN) = dx/sqrt(dx**2 + dy**2)
FF = Ny*(2*Nx+1) + 1
do n = 1, 8
  face(FF-ILE+1,n) = face(FF,n)
enddo
EF = EF + 1
eface(EF) = FF - ILE + 1
nnddo

do FF = Ny*(2*Nx+1)+Nx+1, Fmax
  CC = face(FF,1)
  cell(CC,1) = FF-Nx+ITE-ILE
  do n = 1, 8
    face(FF-Nx+ITE-ILE,n) = face(FF,n)
  enddo
enddo

Fmax = Fmax - Nx + ITE - ILE

EN = EN + 1
node = (Nx+1)*Ny + ITE
enode(EN) = node
dxM = x(node-1) - x(node)
dyM = y(node-1) - y(node)
dxP = x(ITE-1) - x(node)
dyP = y(ITE-1) + pitch - y(node)
if ((dxM*dxP + dyM*dyP) .gt. 0.) then
  dsM = sqrt(dxM**2 + dyM**2)
  dsP = sqrt(dxP**2 + dyP**2)
  dx = dxM/dsM + dxP/dsP
  dy = dyM/dsM + dyP/dsP
else
  dx = dxP - dxM
  dy = dyP - dyM
endif
senode(EN) = dy/sqrt(dx**2 + dy**2)
cenode(EN) = dx/sqrt(dx**2 + dy**2)

tnode = EN

c** upper edge of airfoil
do i = ILE, ITE-1
  EF = EF + 1
  eface(EF) = 1
  EN = EN + 1
  node = i
  enode(EN) = node
  dxP = x(node+1) - x(node)
  dyP = y(node+1) - y(node)
  if (i .eq. ILE) then
    dxM = x((Nx+1)*Ny+ILE+1) - x(node)
    dyM = y((Nx+1)*Ny+ILE+1) - pitch - y(node)
  else
    dxM = x(node-1) - x(node)
    dyM = y(node-1) - y(node)
  endif
  if ((dxM*dxP + dyM*dyP) .gt. 0.) then
    dsM = sqrt(dxM**2 + dyM**2)
    dsP = sqrt(dxP**2 + dyP**2)
    dx = dxM/dsM + dxP/dsP
    dy = dyM/dsM + dyP/dsP
  else
    dx = dxP - dxM
    dy = dyP - dyM
  endif
  senode(EN) = dy/sqrt(dx**2 + dy**2)
  cenode(EN) = dx/sqrt(dx**2 + dy**2)

```

```

        enddo

        EN = EN + 1
        node = ITE
        enode(EN) = node
        dxM = x(node-1) - x(node)
        dyM = y(node-1) - y(node)
        dxP = x((Nx+1)*Ny+ITE-1) - x(node)
        dyP = y((Nx+1)*Ny+ITE-1) - pitch - y(node)
        if ((dxM*dxP + dyM*dyP) .gt. 0.) then
            dsM = sqrt(dxM**2 + dyM**2)
            dsP = sqrt(dxP**2 + dyP**2)
            dx = dxM/dsM + dxP/dsP
            dy = dyM/dsM + dyP/dsP
        else
            dx = dxP - dxM
            dy = dyP - dyM
        endif
        senode(EN) = dy/sqrt(dx**2 + dy**2)
        cenode(EN) = dx/sqrt(dx**2 + dy**2)

        bnode = EN
        Wmax = 2*(ITE-ILE)

    endif

c** inlet and outlet boundaries
    do j = 1, Ny
        EN = EN + 1
        enode(EN) = (j-1)*(Nx+1) + 1
        IN = IN + 1
        innode(IN) = (j-1)*(Nx+1) + 1
        EF = EF + 1
        eface(EF) = (2*Nx+1)*j - Nx
    enddo

    EN = EN + 1
    enode(EN) = Ny*(Nx+1) + 1
    IN = IN + 1
    innode(IN) = Ny*(Nx+1) + 1

    inode = bnode + Ny + 1
    imax = IN

    do j = 1, Ny
        EN = EN + 1
        enode(EN) = (Nx+1)*j
        ON = ON + 1
        outnode(ON) = (Nx+1)*j
        EF = EF + 1
        eface(EF) = (2*Nx+1)*j
    enddo

    EN = EN + 1
    enode(EN) = (Nx+1)*(Ny+1)
    ON = ON + 1
    outnode(ON) = (Nx+1)*(Ny+1)

    ENmax = EN
    Emex = EF
    Omax = ON
    if ((Emex .gt. Maxedges) .or.
        & (ENmax .gt. Maxedges) .or.
        & (Pmax .gt. Maxedges)) then
        write(6,*) 'EF = ', EF
        write(6,*) 'EN = ', EN
        write(6,*) 'PN = ', PN
        write(6,*) 'more than Maxedges required'
    endif

    return
end

```



```

c*****
c*
c*   renumber edges so nodes are consecutive along edges
c*   and N3 is the interior node
c*
c*****

      subroutine edgenumber
      implicit none

      include 'QUAD.INC'
      integer EF, FF
      integer save
      !pointers
      !save value in face

      do EF = 1, Emax
         FF = eFace(EF)
         if (face(FF,1).eq.0) then
            save = face(FF,1)
            face(FF,1) = face(FF,2)
            face(FF,2) = save

            save = face(FF,3)
            face(FF,3) = face(FF,4)
            face(FF,4) = save

            save = face(FF,5)
            face(FF,5) = face(FF,7)
            face(FF,7) = save

            save = face(FF,6)
            face(FF,6) = face(FF,8)
            face(FF,8) = save
         endif
      enddo

      return
      end

      subroutine cellcolor
      implicit none

      include 'QUAD.INC'
      integer CC, NN, FF
      integer Ncells
      integer CC2(0:Maxcells)
      integer Ncolor
      integer cell2(Maxcells,8)
      logical hascolor(Maxcells)
      !pointers
      !number of cells colored
      !new number of cell
      !current color
      !new cell number
      !has node been colored current color?

c** Initialize everything
      Ncells = 0
      do CC = 1, Cmax
         CC2(CC) = 0
      enddo

c** Loop over colors
      do Ncolor = 1, 50

c** Initialize node color array
         NCcolor(Ncolor) = 0
         do NN = 1, Nmax
            hascolor(NN) = .FALSE.
         enddo
      enddo

```

```

c** Loop over cells
    do CC = 1, Cmax

c** Check if cell is already colored with old color
    if(CC2(CC) .NE. 0) GOTO 22
c** Check if cell nodes are already colored with new color
    if( hascolor(cell(CC,5)) .OR.
       & hascolor(cell(CC,6)) .OR.
       & hascolor(cell(CC,7)) .OR.
       & hascolor(cell(CC,8)) ) GOTO 22

c** Set color markers
        Ncells = Ncells + 1
        CC2(CC) = Ncells
        NCcolor(Ncolor) = NCcolor(Ncolor) + 1
        hascolor(cell(CC,5)) = .TRUE.
        hascolor(cell(CC,6)) = .TRUE.
        hascolor(cell(CC,7)) = .TRUE.
        hascolor(cell(CC,8)) = .TRUE.

22      enddo

        if(Ncells.EQ.Cmax) GOTO 23

        enddo

        STOP 'COLOR; more than 50 colors required for cells'

23      Ccclormax= Ncolor

c** Redo pointers
    do CC = 1, Cmax
        cell12(CC2(CC),1) = cell(CC,1)
        cell12(CC2(CC),2) = cell(CC,2)
        cell12(CC2(CC),3) = cell(CC,3)
        cell12(CC2(CC),4) = cell(CC,4)
        cell12(CC2(CC),5) = cell(CC,5)
        cell12(CC2(CC),6) = cell(CC,6)
        cell12(CC2(CC),7) = cell(CC,7)
        cell12(CC2(CC),8) = cell(CC,8)
    enddo

    do CC = 1, Cmax
        cell(CC,1) = cell12(CC,1)
        cell(CC,2) = cell12(CC,2)
        cell(CC,3) = cell12(CC,3)
        cell(CC,4) = cell12(CC,4)
        cell(CC,5) = cell12(CC,5)
        cell(CC,6) = cell12(CC,6)
        cell(CC,7) = cell12(CC,7)
        cell(CC,8) = cell12(CC,8)
    enddo

    CC2(0) = 0

    do FF = 1, Fmax
        face(FF,1) = CC2(face(FF,1))
        face(FF,2) = CC2(face(FF,2))
    enddo

    return
end

subroutine facecolor
implicit none

```

```

include 'QUAD.INC'
integer FF, NN, CC, EF           !pointers
integer Nfaces                   !number of faces colored
integer FF2(Maxfaces)           !new number of face
integer Ncolor                   !current color
integer face2(Maxfaces,8)       !new face number
logical hascolor(0:Maxfaces)    !has node been colored current color?

c** Initialize everything
Nfaces = 0
do FF = 1, Fmax
  FF2(FF) = 0
enddo

c** Loop over colors
do Ncolor = 1, 50

c** Initialize node color array
  NFcolor(Ncolor) = 0
  do NN = 1, Nmax
    hascolor(NN) = .FALSE.
  enddo

c** Loop over faces
  do FF = 1, Fmax

    hascolor(0) = .FALSE.

c** Check if face is already colored with old color
    if(FF2(FF).NE.0) GOTO 22

c** Check if face nodes are already colored with new color
    if( hascolor(face(FF,5)) .OR.
      &   hascolor(face(FF,6)) .OR.
      &   hascolor(face(FF,7)) .OR.
      &   hascolor(face(FF,8)) ) GOTO 22

c** Set color markers
      Nfaces = Nfaces + 1
      FF2(FF) = Nfaces
      NFcolor(Ncolor) = NFcolor(Ncolor) + 1
      hascolor(face(FF,5)) = .TRUE.
      hascolor(face(FF,6)) = .TRUE.
      hascolor(face(FF,7)) = .TRUE.
      hascolor(face(FF,8)) = .TRUE.

22      enddo

      if(Nfaces.EQ.Fmax) GOTO 23

      enddo

      STOP 'COLOR; more than 50 colors required for faces'

23      Fcolormax= Ncolor

c** Redo pointers
      do FF = 1, Fmax
        face2(FF2(FF),1) = face(FF,1)
        face2(FF2(FF),2) = face(FF,2)
        face2(FF2(FF),3) = face(FF,3)
        face2(FF2(FF),4) = face(FF,4)
        face2(FF2(FF),5) = face(FF,5)
        face2(FF2(FF),6) = face(FF,6)
        face2(FF2(FF),7) = face(FF,7)
        face2(FF2(FF),8) = face(FF,8)
      enddo

      do FF = 1, Fmax
        face(FF,1) = face2(FF,1)
        face(FF,2) = face2(FF,2)
        face(FF,3) = face2(FF,3)

```

```

        face(FF,4) = face2(FF,4)
        face(FF,5) = face2(FF,5)
        face(FF,6) = face2(FF,6)
        face(FF,7) = face2(FF,7)
        face(FF,8) = face2(FF,8)
    enddo

    do CC = 1, Cmax
        cell(CC,1) = FF2(cell(CC,1))
        cell(CC,2) = FF2(cell(CC,2))
        cell(CC,3) = FF2(cell(CC,3))
        cell(CC,4) = FF2(cell(CC,4))
    enddo

    do EF = 1, Emax
        eface(EF) = FF2(eface(EF))
    enddo

    return
end

subroutine edgecolor
implicit none

include 'QUAD.INC'
integer FF, NN, EF           !pointers
integer Nedges               !number of faces colored
integer EF2(Maxedges)        !new number of face
integer Ncolor               !current color
integer eface2(Maxedges)     !new face number
logical hascolor(Maxnodes)   !has node been colored current color?

c** Initialize everything
Nedges = 0
do EF = 1, Emax
    EF2(EF) = 0
enddo

c** Loop over colors
do Ncolor = 1, 5

c** Initialize node color array
NEcolor(Ncolor) = 0
do NN = 1, Nmax
    hascolor(NN) = .FALSE.
enddo

c** Loop over faces
do EF = 1, Wmax
    FF = eface(EF)

c** Check if face is already colored with old color
    if(EF2(EF).NE.0) GOTO 22

c** Check if face nodes are already colored with new color
    if( hascolor(face(FF,3)) .OR.
    &    hascolor(face(FF,4)) .OR.
    &    hascolor(face(FF,5)) .OR.
    &    hascolor(face(FF,6)) ) GOTO 22

c** Set color markers
        Nedges = Nedges + 1
        EF2(EF) = Nedges
        NEcolor(Ncolor) = NEcolor(Ncolor) + 1
        hascolor(face(FF,3)) = .TRUE.
        hascolor(face(FF,4)) = .TRUE.
        hascolor(face(FF,5)) = .TRUE.
        hascolor(face(FF,6)) = .TRUE.
    enddo
enddo

```

```

22      enddo

      if(Nedges.EQ.Wmax) GOTO 23

      enddo

      STOP 'COLOR; more than 5 colors required for wall edges'

23      Eicolormax= Ncolor

c** Loop over colors
      do Ncolor = Eicolormax+1, 10

c** Initialize node color array
      NEcolor(Ncolor) = 0
      do NN = 1, Nmax
         hascolor(NN) = .FALSE.
      enddo

c** Loop over faces
      do EF = Wmax+1, Emax
         FF = eface(EF)

c** Check if face is already colored with old color
         if(EF2(EF).NE.0) GOTO 32

c** Check if face nodes are already colored with new color
         if( hascolor(face(FF,3)) .OR.
            &   hascolor(face(FF,4)) .OR.
            &   hascolor(face(FF,5)) .OR.
            &   hascolor(face(FF,6)) ) GOTO 32

c** Set color markers
            Nedges = Nedges + 1
            EF2(EF) = Nedges
            NEcolor(Ncolor) = NEcolor(Ncolor) + 1
            hascolor(face(FF,3)) = .TRUE.
            hascolor(face(FF,4)) = .TRUE.
            hascolor(face(FF,5)) = .TRUE.
            hascolor(face(FF,6)) = .TRUE.

32      enddo

      if(Nedges.EQ.Emax) GOTO 33

      enddo

      STOP 'COLOR; more than 10 colors required for all edges'

33      E2colormax= Ncolor

c** Redo pointers
      do EF = 1, Emax
         eface2(EF2(EF)) = eface(EF)
      enddo

      do EF = 1, Emax
         eface(EF) = eface2(EF)
      enddo

      return
      end

```

C*****

```

c*                                     *
c*   program to create an irregular quadrilateral mesh                       *
c*   from a regular quadrilateral mesh                                       *
c*                                     *
c*****
      program griddis
      implicit none

      include 'QUAD.INC'
      integer NN, EN                                !pointers
      real CC, AA
      integer N1, N2
      real change
      real yc, RR2
      real tau
      real pi

      pi = 3.14159

      write(6,1)
1      format(1,' tau = ')
      accept 11, tau
11     format(F)

c      tau = 0.1

      call gridio(1)

      N1 = face(1,3)
      N2 = face(1,4)
      CC = 0.3*(x(N1) - x(N2))
      AA = (0.3*30.)/CC
      do NN = 1, Nmax
        if ((x(NN).ne.-1.) .and. (x(NN).ne.2.)) then
          change = CC*cos(AA*y(NN)+12.)*cos(AA*x(NN)+43.)
          x(NN) = x(NN) + change
        endif
        if ((y(NN).ne.0.) .and. (y(NN).ne.1.)) then
          change = CC*cos(AA*y(NN)+57.)*cos(AA*x(NN)+33.)
          y(NN) = y(NN) + change
        endif
      enddo

      do EN = tnode, bnode
        NN = enode(EN)
        if (x(NN).gt.0. .and. x(NN).lt.1.) then
          if (tau.lt.0) then
            y(NN) = tau*(sin(pi*x(NN)))*2
          else if (tau.gt.0) then
            yc = (tau**2 - 0.25)/(2.*tau)
            RR2 = 0.25 + yc**2
            y(NN) = yc + sqrt(RR2 - (x(NN)-0.5)**2)
          endif
        endif
      enddo

      call gridio(0)

      stop
      end

```

A.2.3 Ni Scheme

```

c*****o*****c*****

```

```

c*
c*      main program for quadrilateral N1 scheme
c*
c*****

      program quadrilateral
      implicit none

      include 'QUAD.INC'
      integer Niter           !number of iterations
      integer NN, CC, 1       !pointer
      integer N1, N2, N3, N4  !nodes at corners of cell
      real maxchange          !max change in state vector
      integer maxnode, maxeqn !where max change occurs

c*  read in data from file
      call gridio(1)
      call flowio(1)
      call input

c*  calculate control area around each cell
      call calcarea

      Niter = 0
      durms = 999.

c*  start history file from the top
      open(unit=35,status='unknown',form='formatted')
      write(35,2) Min1
      close(unit=35)
      2    format(' inlet Mach number = ',f5.3)

c*  loop until converged
      do while ((Niter.lt.Maxiter) .and. (durms.gt.2.e-7))
        Niter = Niter + 1

c*  set cell values of state vector
        do CC = 1, Cmax
          N1 = cell(CC,5)
          N2 = cell(CC,6)
          N3 = cell(CC,7)
          N4 = cell(CC,8)
          Uc(1,CC) = (U(1,N1) + U(1,N2) + U(1,N3) + U(1,N4))*0.25
          Uc(2,CC) = (U(2,N1) + U(2,N2) + U(2,N3) + U(2,N4))*0.25
          Uc(3,CC) = (U(3,N1) + U(3,N2) + U(3,N3) + U(3,N4))*0.25
          Uc(4,CC) = (U(4,N1) + U(4,N2) + U(4,N3) + U(4,N4))*0.25
        enddo

c*  calculate time step for each cell
        call timestep

c*  calculate flux at each node
        call nodeflux

c*  calculate change in state vector and fluxes at each cell
        call delcell

c*  calculate change in state vector at each node
        call delstate

c*  add smoothing term
        if (eps2.ne. 0.) call smooth

c** account for periodic nodes
        call bperiodic

c*  set inlet and outlet boundary conditions
        call binlet
        call boutlet

c*  change momentum change to make flow tangent at walls
        call tangent

```

```

c* update state vector
do NN = 1, Nmax
  U(1,NN) = U(1,NN) + dU(1,NN)
  U(2,NN) = U(2,NN) + dU(2,NN)
  U(3,NN) = U(3,NN) + dU(3,NN)
  U(4,NN) = U(4,NN) + dU(4,NN)
enddo

c* find root mean square difference in state vector
if (mod(Niter,10).eq.0 .or. Niter.lt.10) then
  durms = 0.0
  maxchange = 0.

  do i = 1, 4
    do NN = 1, Nmax
      durms = durms + dU(i,NN)**2
      if (abs(dU(i,NN)).gt.abs(maxchange)) then
        maxchange = dU(i,NN)
        maxnode = NN
        maxeqn = i
      endif
    enddo
  enddo

  durms = sqrt(durms/(4.*Nmax))

c* print diagnostics to screen and file
call flowio(0)
open(unit=50, status='unknown', form='unformatted')
write(50) Cmax, (vol(CC),CC=1,Cmax)
close(unit=50)

10      open(unit=35, status='old',access='append',err=10)
        write(35,1) Niter, durms, maxchange, x(maxnode),
&          y(maxnode), maxeqn
        close(unit=35)
        write(6,1) Niter, durms, maxchange, x(maxnode),
&          y(maxnode), maxeqn

        endif

      enddo

1      format('Niter=',i4,' rms=',f9.7,' max=',f9.7,' x=',f6.3,
&          ' y=',f6.3,' eqn=',i1)

c* write out data to file
call flowio(0)

stop
end

```

```

c*****
c*
c*      calculate areas of cells
c*
c*****

```

```

subroutine calcarea
implicit none

```

```

include 'QUAD.INC'
integer EF, FF, NN, CC, PN      !pointer
integer N1, N2, N3, N4          !nodes at corner of cell
integer C1                      !cell on boundary
integer P1, P2                  !periodic nodes
real AA                         !area of cell

```



```

integer CC1, CC2, col           !coloring pointers
integer EF1, EF2

c* calculate area of cells and distribute to nodes
do CC = 1, Cmax
  N1 = cell(CC,5)
  N2 = cell(CC,6)
  N3 = cell(CC,7)
  N4 = cell(CC,8)
  AA = -0.5*((x(N2) - x(N4))*(y(N1) - y(N3))
    &      - (x(N1) - x(N3))*(y(N2) - y(N4)))
  areaC(CC) = AA
enddo

return
end

c*****
c*
c*      calculate time step for nodes and cells
c*
c*****

subroutine timestep
implicit none

include 'QUAD.INC'
integer CC, NN, PN              !pointers
integer P1, P2                  !periodic nodes
integer N1, N2, N3, N4          !nodes around cell
real dx1, dy1, d1
real dxm, dym, dm
real rr, uu, vv, pp, aa         !values for cell
real time1, time2

do NN = 1, Nmax
  deltn(NN) = 0.
enddo

c* find time step for each cell
do CC = 1, Cmax
  N1 = cell(CC,5)
  N2 = cell(CC,6)
  N3 = cell(CC,7)
  N4 = cell(CC,8)
  dx1 = .5*(x(N3) + x(N2) - x(N4) - x(N1))
  dy1 = .5*(y(N3) + y(N2) - y(N4) - y(N1))
  d1 = sqrt(dx1**2 + dy1**2)
  dxm = .5*(x(N4) + x(N3) - x(N1) - x(N2))
  dym = .5*(y(N4) + y(N3) - y(N1) - y(N2))
  dm = sqrt(dxm**2 + dym**2)
  rr = Uc(1,CC)
  uu = Uc(2,CC)/rr
  vv = Uc(3,CC)/rr
  pp = gam1*(Uc(4,CC) - 0.5*rr*(uu**2 + vv**2))
  aa = sqrt(gam*pp/rr)

  time1 = 1./(abs(uu*dy1-vv*dx1) + aa*d1)
  time2 = 1./(abs(uu*dym-vv*dxm) + aa*dm)

  deltc(CC) = CFL*min(time1,time2)

  deltn(N1) = deltn(N1) + 0.25/deltC(CC)
  deltn(N2) = deltn(N2) + 0.25/deltC(CC)
  deltn(N3) = deltn(N3) + 0.25/deltC(CC)
  deltn(N4) = deltn(N4) + 0.25/deltC(CC)
enddo

```

```

c** periodic nodes
do PN = 1, Pmax
  P1 = pnode(PN,1)
  P2 = pnode(PN,2)
  deltn(P1) = deltn(P1) + deltn(P2)
  deltn(P2) = deltn(P1)
enddo

do NN = 1, Nmax
  deltn(NN) = 1./deltn(NN)
enddo

return
end

c*****
c*
c*   calculate flux vector values at nodes
c*
c*****

subroutine nodeflux
implicit none

include 'QUAD.INC'
integer NN                !pointers
real WW                   !kinetic energy

do NN = 1, Nmax
c* calculate f and g at nodes
  WW = 0.5*(U(2,NN)**2 + U(3,NN)**2)/U(1,NN)
  F(1,NN) = U(2,NN)
  F(2,NN) = U(2,NN)**2/U(1,NN) + gam1*(U(4,NN) - WW)
  F(3,NN) = U(2,NN)*U(3,NN)/U(1,NN)
  F(4,NN) = (U(2,NN)/U(1,NN))*(gam*U(4,NN) - gam1*WW)

  G(1,NN) = U(3,NN)
  G(2,NN) = U(2,NN)*U(3,NN)/U(1,NN)
  G(3,NN) = U(3,NN)**2/U(1,NN) + gam1*(U(4,NN) - WW)
  G(4,NN) = (U(3,NN)/U(1,NN))*(gam*U(4,NN) - gam1*WW)
enddo

return
end

c*****
c*
c*   this subroutine finds the change in the fluxes
c*   F and G and the change in the state U at each cell
c*
c*****

subroutine delcell
implicit none

include 'QUAD.INC'
integer CC
integer M1, N2, N3, N4    !pointers
                           !nodes at corners of cells
real coef
real dy31, dy24, dx31, dx24 !change in x and y
real dF24, dF13, dG24, dG13 !change in F and G
real u1, u2, u3, u4        !velocity at nodes
real v1, v2, v3, v4

```

```

c* find change in state vector for cell
  do CC = 1, Cmax
    N1 = cell(CC,5)
    N2 = cell(CC,6)
    N3 = cell(CC,7)
    N4 = cell(CC,8)

    u1 = U(2,N1)/U(1,N1)
    u2 = U(2,N2)/U(1,N2)
    u3 = U(2,N3)/U(1,N3)
    u4 = U(2,N4)/U(1,N4)
    v1 = U(3,N1)/U(1,N1)
    v2 = U(3,N2)/U(1,N2)
    v3 = U(3,N3)/U(1,N3)
    v4 = U(3,N4)/U(1,N4)

    dy31 = y(N3) - y(N1)
    dy24 = y(N2) - y(N4)
    dx31 = x(N3) - x(N1)
    dx24 = x(N2) - x(N4)

    coef = 0.5*delC(CC)

    dF24 = F(1,N2) - F(1,N4)
    dF13 = F(1,N1) - F(1,N3)
    dG24 = G(1,N2) - G(1,N4)
    dG13 = G(1,N1) - G(1,N3)
    dUc(1,CC) = coef*(-dF24*dy31 + dG24*dx31 -
*                      dF13*dy24 + dG13*dx24)

    dF24 = F(2,N2) - F(2,N4)
    dF13 = F(2,N1) - F(2,N3)
    dG24 = G(2,N2) - G(2,N4)
    dG13 = G(2,N1) - G(2,N3)
    dUc(2,CC) = coef*(-dF24*dy31 + dG24*dx31 -
*                      dF13*dy24 + dG13*dx24)

    dF24 = F(3,N2) - F(3,N4)
    dF13 = F(3,N1) - F(3,N3)
    dG24 = G(3,N2) - G(3,N4)
    dG13 = G(3,N1) - G(3,N3)
    dUc(3,CC) = coef*(-dF24*dy31 + dG24*dx31 -
*                      dF13*dy24 + dG13*dx24)

    dF24 = F(4,N2) - F(4,N4)
    dF13 = F(4,N1) - F(4,N3)
    dG24 = G(4,N2) - G(4,N4)
    dG13 = G(4,N1) - G(4,N3)
    dUc(4,CC) = coef*(-dF24*dy31 + dG24*dx31 -
*                      dF13*dy24 + dG13*dx24)

c** shock smoothing
    vol(CC) = (u1-u3)*dy24 - (v1-v3)*dx24 +
*             (u2-u4)*dy31 - (v2-v4)*dx31
    enddo

c* implement wall boundary conditions
    call bwall

    return
  end

c*****
c*
c*   this subroutine applies the wall boundary condition
c*
c*****

```

```

subroutine bwall
implicit none

include 'QUAD.INC'
integer EF, FF, CC      !pointer
integer N1, N2          !nodes on edge
real coef
real dy12, dx12        !change in x and y
real presi, pres2      !pressure
real u1, u2, v1, v2    !velocity
integer EF1, EF2, col   !coloring pointers

c* upper and lower boundary
EF2 = 0

do col = 1, Eicolormax
EF1 = EF2 + 1
EF2 = EF1 - 1 + NEcolor(col)
do EF = EF1, EF2

    FF = eface(EF)
    CC = face(FF,1)
    N1 = face(FF,3)
    N2 = face(FF,4)

    coef = 0.5*deltC(CC)

    dy12 = y(N1) - y(N2)
    dx12 = x(N1) - x(N2)

    presi = gam1*(U(4,N1) - 0.5*(U(2,N1)**2 + U(3,N1)**2)/
    &      U(1,N1))
    pres2 = gam1*(U(4,N2) - 0.5*(U(2,N2)**2 + U(3,N2)**2)/
    &      U(1,N2))
    u1 = U(2,N1)/U(1,N1)
    u2 = U(2,N2)/U(1,N2)
    v1 = U(3,N1)/U(1,N1)
    v2 = U(3,N2)/U(1,N2)

    dUc(1,CC) = dUc(1,CC) + coef*(
    &      -(F(1,N2) + F(1,N1))*dy12 +
    &      (G(1,N2) + G(1,N1))*dx12)

    dUc(2,CC) = dUc(2,CC) + coef*(
    &      -(F(2,N2) + F(2,N1) - presi - pres2)*dy12 +
    &      (G(2,N2) + G(2,N1))*dx12)

    dUc(3,CC) = dUc(3,CC) + coef*(
    &      -(F(3,N2) + F(3,N1))*dy12 +
    &      (G(3,N2) + G(3,N1) - presi - pres2)*dx12)

    dUc(4,CC) = dUc(4,CC) + coef*(
    &      -(F(4,N2) + F(4,N1))*dy12 +
    &      (G(4,N2) + G(4,N1))*dx12)

c** shock smoothing
vol(CC) = vol(CC) + (u1+u2)*dy12 - (v1+v2)*dx12

    enddo
enddo

return
end

c*****
c*
c*      this subroutine calculates the change at each node
c*

```

C*****

```

subroutine delstate
implicit none

include 'QUAD.INC'
integer NN, CC, EF, FF           !pointers
integer N1, N2, N3, N4           !nodes at corners of cell
real rr, uu, vv, pp, aa, vv2     !values at cell
real rdu, rdu, dp, HH           !changes
real CON                         !change in y on cell edges
real dy42, dy13                 !change in x on cell edges
real dx42, dx13                 ! " " "
real dy21, dx21                 ! " " "
real coef                       !coefficient
integer CC1, CC2, col           !color pointers
integer EF1, EF2               !color pointers
real dFc(4), dGc(4)

do NN = 1, Nmax
  dU(1,NN) = 0.
  dU(2,NN) = 0.
  dU(3,NN) = 0.
  dU(4,NN) = 0.
enddo

CC2 = 0

do col = 1, Ccolormax
  CC1 = CC2 + 1
  CC2 = CC1 - 1 + NCcolor(col)

CVD$ NODEPCHK
  do CC = CC1, CC2
    N1 = cell(CC,5)
    N2 = cell(CC,6)
    N3 = cell(CC,7)
    N4 = cell(CC,8)

    rr = Uc(1,CC)
    uu = Uc(2,CC)/rr
    vv = Uc(3,CC)/rr
    vv2 = 0.5*(uu*uu + vv*vv)

    dy42 = y(N4) - y(N2)
    dy13 = y(N1) - y(N3)
    dx42 = x(N4) - x(N2)
    dx13 = x(N1) - x(N3)

c* find second order change in flux vector at cells
    HH = gam*Uc(4,CC)/Uc(1,CC) - gam1*vv2
    rdu = dUc(2,CC) - uu*dUc(1,CC)
    rdv = dUc(3,CC) - vv*dUc(1,CC)
    dp = gam1*(dUc(4,CC) - uu*dUc(2,CC) - vv*dUc(3,CC)
    & + vv2*dUc(1,CC))

    dFc(1) = dUc(2,CC)
    dFc(2) = uu*(dUc(2,CC) + rdu) + dp
    dFc(3) = vv*(dUc(3,CC) + rdv) + dp
    dFc(4) = uu*(dUc(4,CC) + dp) + HH*rdu

    dGc(1) = dUc(3,CC)
    dGc(2) = uu*dUc(3,CC) + vv*rdu
    dGc(3) = vv*(dUc(3,CC) + rdv) + dp
    dGc(4) = vv*(dUc(4,CC) + dp) + HH*rdv

c** shock smoothing
    vol(CC) = vol(CC)/sqrt(2.*areaC(CC))
    CON = -eps1coef*min(0.5,abs(vol(CC)))*vol(CC)*vv2
    dFc(2) = dFc(2) + CON
    dGc(3) = dGc(3) + CON

```

```

coef = 0.25/deltC(CC)

dU(1,N1) = dU(1,N1) + coef*dUc(1,CC) - 0.25*
*      (dFc(1)*dy42 - dGc(1)*dx42)
dU(1,N2) = dU(1,N2) + coef*dUc(1,CC) - 0.25*
*      (dFc(1)*dy13 - dGc(1)*dx13)
dU(1,N3) = dU(1,N3) + coef*dUc(1,CC) + 0.25*
*      (dFc(1)*dy42 - dGc(1)*dx42)
dU(1,N4) = dU(1,N4) + coef*dUc(1,CC) + 0.25*
*      (dFc(1)*dy13 - dGc(1)*dx13)

dU(2,N1) = dU(2,N1) + coef*dUc(2,CC) - 0.25*
*      (dFc(2)*dy42 - dGc(2)*dx42)
dU(2,N2) = dU(2,N2) + coef*dUc(2,CC) - 0.25*
*      (dFc(2)*dy13 - dGc(2)*dx13)
dU(2,N3) = dU(2,N3) + coef*dUc(2,CC) + 0.25*
*      (dFc(2)*dy42 - dGc(2)*dx42)
dU(2,N4) = dU(2,N4) + coef*dUc(2,CC) + 0.25*
*      (dFc(2)*dy13 - dGc(2)*dx13)

dU(3,N1) = dU(3,N1) + coef*dUc(3,CC) - 0.25*
*      (dFc(3)*dy42 - dGc(3)*dx42)
dU(3,N2) = dU(3,N2) + coef*dUc(3,CC) - 0.25*
*      (dFc(3)*dy13 - dGc(3)*dx13)
dU(3,N3) = dU(3,N3) + coef*dUc(3,CC) + 0.25*
*      (dFc(3)*dy42 - dGc(3)*dx42)
dU(3,N4) = dU(3,N4) + coef*dUc(3,CC) + 0.25*
*      (dFc(3)*dy13 - dGc(3)*dx13)

dU(4,N1) = dU(4,N1) + coef*dUc(4,CC) - 0.25*
*      (dFc(4)*dy42 - dGc(4)*dx42)
dU(4,N2) = dU(4,N2) + coef*dUc(4,CC) - 0.25*
*      (dFc(4)*dy13 - dGc(4)*dx13)
dU(4,N3) = dU(4,N3) + coef*dUc(4,CC) + 0.25*
*      (dFc(4)*dy42 - dGc(4)*dx42)
dU(4,N4) = dU(4,N4) + coef*dUc(4,CC) + 0.25*
*      (dFc(4)*dy13 - dGc(4)*dx13)

enddo
enddo

EF2 = 0

do col = 1, E2colormax
  EF1 = EF2 + 1
  EF2 = EF1 - 1 + NEcolor(col)
  do EF = EF1, EF2

    FF = eface(EF)
    CC = face(FF,1)
    N1 = face(FF,3)
    N2 = face(FF,4)

    dy21 = y(N2) - y(N1)
    dx21 = x(N2) - x(N1)

    uu = Uc(2,CC)/Uc(1,CC)
    vv = Uc(3,CC)/Uc(1,CC)
    ww2 = 0.5*(uu**2 + vv**2)

    dp = gam1*(dUc(4,CC) - uu*dUc(2,CC) - vv*dUc(3,CC)
*      + ww2*dUc(1,CC))

    dFc(2) = dp
    dGc(3) = dp

    dU(2,N1) = dU(2,N1) - 0.25*dFc(2)*dy21
    dU(2,N2) = dU(2,N2) - 0.25*dFc(2)*dy21

    dU(3,N1) = dU(3,N1) + 0.25*dGc(3)*dx21
    dU(3,N2) = dU(3,N2) + 0.25*dGc(3)*dx21

  enddo
enddo

```

```

        enddo

        do NN = 1, Nmax
            dU(1,NN) = dU(1,NN)*deltN(NN)
            dU(2,NN) = dU(2,NN)*deltN(NN)
            dU(3,NN) = dU(3,NN)*deltN(NN)
            dU(4,NN) = dU(4,NN)*deltN(NN)
        enddo

        return
    end

C*****
C*
C*      subroutine to calculate forth difference smoothing
C*
C*****

    subroutine smooth
    implicit none

    include 'QUAD.INC'
    integer N1, N2, N3, N4           !nodes at corners of cell
    integer NN, CC1, CC2, col, CC    !pointers
    integer FF, EF, EF1, EF2        !edge nodes
    integer P1, P2, PN              !periodic nodes
    real dx31, dx24                 !change in x
    real dx12, dx23, dx34, dx41
    real dy31, dy24                 !change in y
    real dy12, dy23, dy34, dy41
    real dxC1, dxC2, dxC3, dxC4     !change in x in cell
    real dyC1, dyC2, dyC3, dyC4     !change in y in cell
    real del12, del13, del14
    real del23, del24, del34
    real coef12, coef13, coef14
    real coef23, coef24, coef34
    real del2(4,Maxnodes)           !gradient at node
    real coef                       !coefficient

    do NN = 1, Nmax
        del2(1,NN) = 0.
        del2(2,NN) = 0.
        del2(3,NN) = 0.
        del2(4,NN) = 0.
    enddo

    CC2 = 0

    do col = 1, Ccolormax
        CC1 = CC2 + 1
        CC2 = CC1 - 1 + NCcolor(col)

CVD$      NODEPCHK
        do CC = CC1, CC2
            N1 = cell(CC,6)
            N2 = cell(CC,6)
            N3 = cell(CC,7)
            N4 = cell(CC,8)

            dy24 = y(N2) - y(N4)
            dy31 = y(N3) - y(N1)
            dy12 = y(N1) - y(N2)
            dy23 = y(N2) - y(N3)
            dy34 = y(N3) - y(N4)
            dy41 = y(N4) - y(N1)

            dx24 = x(N2) - x(N4)
            dx31 = x(N3) - x(N1)
            dx12 = x(N1) - x(N2)

```

```

dx23 = x(N2) - x(N3)
dx34 = x(N3) - x(N4)
dx41 = x(N4) - x(N1)

coef = ((max(0.,min(1.,(1.+10.*vol(CC))))-1.)*eps1coef + 1.)
      *(0.5/abs(-dx12*dy41 + dy12*dx41))

dxC1 = U(1,N1)*dy24 + U(1,N2)*dy41 + U(1,N4)*dy12
dxC2 = U(2,N1)*dy24 + U(2,N2)*dy41 + U(2,N4)*dy12
dxC3 = U(3,N1)*dy24 + U(3,N2)*dy41 + U(3,N4)*dy12
dxC4 = U(4,N1)*dy24 + U(4,N2)*dy41 + U(4,N4)*dy12

dyC1 = U(1,N1)*dx24 + U(1,N2)*dx41 + U(1,N4)*dx12
dyC2 = U(2,N1)*dx24 + U(2,N2)*dx41 + U(2,N4)*dx12
dyC3 = U(3,N1)*dx24 + U(3,N2)*dx41 + U(3,N4)*dx12
dyC4 = U(4,N1)*dx24 + U(4,N2)*dx41 + U(4,N4)*dx12

del2(1,N1) = del2(1,N1) + coef*(dxC1*dy24 + dyC1*dx24)
del2(2,N1) = del2(2,N1) + coef*(dxC2*dy24 + dyC2*dx24)
del2(3,N1) = del2(3,N1) + coef*(dxC3*dy24 + dyC3*dx24)
del2(4,N1) = del2(4,N1) + coef*(dxC4*dy24 + dyC4*dx24)

coef = ((max(0.,min(1.,(1.+10.*vol(CC))))-1.)*eps1coef + 1.)
      *(0.5/abs(-dx12*dy23 + dy12*dx23))

dxC1 = U(1,N1)*dy23 + U(1,N2)*dy31 + U(1,N3)*dy12
dxC2 = U(2,N1)*dy23 + U(2,N2)*dy31 + U(2,N3)*dy12
dxC3 = U(3,N1)*dy23 + U(3,N2)*dy31 + U(3,N3)*dy12
dxC4 = U(4,N1)*dy23 + U(4,N2)*dy31 + U(4,N3)*dy12

dyC1 = U(1,N1)*dx23 + U(1,N2)*dx31 + U(1,N3)*dx12
dyC2 = U(2,N1)*dx23 + U(2,N2)*dx31 + U(2,N3)*dx12
dyC3 = U(3,N1)*dx23 + U(3,N2)*dx31 + U(3,N3)*dx12
dyC4 = U(4,N1)*dx23 + U(4,N2)*dx31 + U(4,N3)*dx12

del2(1,N2) = del2(1,N2) + coef*(dxC1*dy31 + dyC1*dx31)
del2(2,N2) = del2(2,N2) + coef*(dxC2*dy31 + dyC2*dx31)
del2(3,N2) = del2(3,N2) + coef*(dxC3*dy31 + dyC3*dx31)
del2(4,N2) = del2(4,N2) + coef*(dxC4*dy31 + dyC4*dx31)

coef = ((max(0.,min(1.,(1.+10.*vol(CC))))-1.)*eps1coef + 1.)
      *(0.5/abs(-dx23*dy34 + dy23*dx34))

dxC1 = U(1,N4)*dy23 + U(1,N2)*dy34 - U(1,N3)*dy24
dxC2 = U(2,N4)*dy23 + U(2,N2)*dy34 - U(2,N3)*dy24
dxC3 = U(3,N4)*dy23 + U(3,N2)*dy34 - U(3,N3)*dy24
dxC4 = U(4,N4)*dy23 + U(4,N2)*dy34 - U(4,N3)*dy24

dyC1 = U(1,N4)*dx23 + U(1,N2)*dx34 - U(1,N3)*dx24
dyC2 = U(2,N4)*dx23 + U(2,N2)*dx34 - U(2,N3)*dx24
dyC3 = U(3,N4)*dx23 + U(3,N2)*dx34 - U(3,N3)*dx24
dyC4 = U(4,N4)*dx23 + U(4,N2)*dx34 - U(4,N3)*dx24

del2(1,N3) = del2(1,N3) - coef*(dxC1*dy24 + dyC1*dx24)
del2(2,N3) = del2(2,N3) - coef*(dxC2*dy24 + dyC2*dx24)
del2(3,N3) = del2(3,N3) - coef*(dxC3*dy24 + dyC3*dx24)
del2(4,N3) = del2(4,N3) - coef*(dxC4*dy24 + dyC4*dx24)

coef = ((max(0.,min(1.,(1.+10.*vol(CC))))-1.)*eps1coef + 1.)
      *(0.5/abs(-dx34*dy41 + dy34*dx41))

dxC1 = U(1,N1)*dy34 + U(1,N3)*dy41 - U(1,N4)*dy31
dxC2 = U(2,N1)*dy34 + U(2,N3)*dy41 - U(2,N4)*dy31
dxC3 = U(3,N1)*dy34 + U(3,N3)*dy41 - U(3,N4)*dy31
dxC4 = U(4,N1)*dy34 + U(4,N3)*dy41 - U(4,N4)*dy31

dyC1 = U(1,N1)*dx34 + U(1,N3)*dx41 - U(1,N4)*dx31
dyC2 = U(2,N1)*dx34 + U(2,N3)*dx41 - U(2,N4)*dx31
dyC3 = U(3,N1)*dx34 + U(3,N3)*dx41 - U(3,N4)*dx31
dyC4 = U(4,N1)*dx34 + U(4,N3)*dx41 - U(4,N4)*dx31

del2(1,N4) = del2(1,N4) - coef*(dxC1*dy31 + dyC1*dx31)

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```

del2(2,N4) = del2(2,N4) - coef*(dxC2*dy31 + dyC2*dx31)
del2(3,N4) = del2(3,N4) - coef*(dxC3*dy31 + dyC3*dx31)
del2(4,N4) = del2(4,N4) - coef*(dxC4*dy31 + dyC4*dx31)

enddo
enddo

EF2 = 0

do col = 1, Elcolormax
  EF1 = EF2 + 1
  EF2 = EF1 - 1 + NEcolor(col)
CVD$ NODEPCHK
do EF = EF1, EF2
  FF = eface(EF)
  CC = face(FF,1)
  N1 = face(FF,3)
  N2 = face(FF,4)
  N3 = face(FF,6)
  N4 = face(FF,5)

  dy24 = y(N2) - y(N4)
  dy31 = y(N3) - y(N1)
  dy12 = y(N1) - y(N2)
  dy23 = y(N2) - y(N3)
  dy41 = y(N4) - y(N1)

  dx24 = x(N2) - x(N4)
  dx31 = x(N3) - x(N1)
  dx12 = x(N1) - x(N2)
  dx23 = x(N2) - x(N3)
  dx41 = x(N4) - x(N1)

  coef = ((max(0.,min(1.,(1.+10.*vol(CC))))-1.)*eps1coef +1.)
  & *(0.5/abs(-dx12*dy31 + dy12*dx31))

  dxC1 = U(1,N1)*dy23 + U(1,N2)*dy31 + U(1,N3)*dy12
  dxC2 = U(2,N1)*dy23 + U(2,N2)*dy31 + U(2,N3)*dy12
  dxC3 = U(3,N1)*dy23 + U(3,N2)*dy31 + U(3,N3)*dy12
  dxC4 = U(4,N1)*dy23 + U(4,N2)*dy31 + U(4,N3)*dy12

  dyC1 = U(1,N1)*dx23 + U(1,N2)*dx31 + U(1,N3)*dx12
  dyC2 = U(2,N1)*dx23 + U(2,N2)*dx31 + U(2,N3)*dx12
  dyC3 = U(3,N1)*dx23 + U(3,N2)*dx31 + U(3,N3)*dx12
  dyC4 = U(4,N1)*dx23 + U(4,N2)*dx31 + U(4,N3)*dx12

  del2(1,N1) = del2(1,N1) + coef*(dxC1*dy12 + dyC1*dx12)
  del2(2,N1) = del2(2,N1) + coef*(dxC2*dy12 + dyC2*dx12)
  del2(3,N1) = del2(3,N1) + coef*(dxC3*dy12 + dyC3*dx12)
  del2(4,N1) = del2(4,N1) + coef*(dxC4*dy12 + dyC4*dx12)

  coef = ((max(0.,min(1.,(1.+10.*vol(CC))))-1.)*eps1coef +1.)
  & *(0.5/abs(-dx12*dy24 + dy12*dx24))

  dxC1 = U(1,N1)*dy24 + U(1,N2)*dy41 + U(1,N4)*dy12
  dxC2 = U(2,N1)*dy24 + U(2,N2)*dy41 + U(2,N4)*dy12
  dxC3 = U(3,N1)*dy24 + U(3,N2)*dy41 + U(3,N4)*dy12
  dxC4 = U(4,N1)*dy24 + U(4,N2)*dy41 + U(4,N4)*dy12

  dyC1 = U(1,N1)*dx24 + U(1,N2)*dx41 + U(1,N4)*dx12
  dyC2 = U(2,N1)*dx24 + U(2,N2)*dx41 + U(2,N4)*dx12
  dyC3 = U(3,N1)*dx24 + U(3,N2)*dx41 + U(3,N4)*dx12
  dyC4 = U(4,N1)*dx24 + U(4,N2)*dx41 + U(4,N4)*dx12

  del2(1,N2) = del2(1,N2) + coef*(dxC1*dy12 + dyC1*dx12)
  del2(2,N2) = del2(2,N2) + coef*(dxC2*dy12 + dyC2*dx12)
  del2(3,N2) = del2(3,N2) + coef*(dxC3*dy12 + dyC3*dx12)
  del2(4,N2) = del2(4,N2) + coef*(dxC4*dy12 + dyC4*dx12)

enddo
enddo

```

```

CVD$  NODEPCHK
      do PN = 1, Pmax
        P1 = pnode(PN,1)
        P2 = pnode(PN,2)
        del2(1,P1) = del2(1,P1) + del2(1,P2)
        del2(2,P1) = del2(2,P1) + del2(2,P2)
        del2(3,P1) = del2(3,P1) + del2(3,P2)
        del2(4,P1) = del2(4,P1) + del2(4,P2)
        del2(1,P2) = del2(1,P1)
        del2(2,P2) = del2(2,P1)
        del2(3,P2) = del2(3,P1)
        del2(4,P2) = del2(4,P1)
      enddo

      CC2 = 0

      do col = 1, Ccolormax
        CC1 = CC2 + 1
        CC2 = CC1 - 1 + NCcolor(col)
      enddo

CVD$  NODEPCHK
      do CC = CC1, CC2
        N1 = cell(CC,5)
        N2 = cell(CC,6)
        N3 = cell(CC,7)
        N4 = cell(CC,8)

        coef12 = eps2/deltC(CC)
        coef13 = eps2/deltC(CC)
        coef14 = eps2/deltC(CC)
        coef23 = eps2/deltC(CC)
        coef24 = eps2/deltC(CC)
        coef34 = eps2/deltC(CC)

        del12 = coef12*(del2(1,N1) - del2(1,N2))
        del13 = coef13*(del2(1,N1) - del2(1,N3))
        del14 = coef14*(del2(1,N1) - del2(1,N4))
        del23 = coef23*(del2(1,N2) - del2(1,N3))
        del24 = coef24*(del2(1,N2) - del2(1,N4))
        del34 = coef34*(del2(1,N3) - del2(1,N4))

        dU(1,N1) = dU(1,N1) + (-del12 - del13 - del14)*deltN(N1)
        dU(1,N2) = dU(1,N2) + ( del12 - del23 - del24)*deltN(N2)
        dU(1,N3) = dU(1,N3) + ( del13 + del23 - del34)*deltN(N3)
        dU(1,N4) = dU(1,N4) + ( del14 + del24 + del34)*deltN(N4)

        del12 = coef12*(del2(2,N1) - del2(2,N2))
        del13 = coef13*(del2(2,N1) - del2(2,N3))
        del14 = coef14*(del2(2,N1) - del2(2,N4))
        del23 = coef23*(del2(2,N2) - del2(2,N3))
        del24 = coef24*(del2(2,N2) - del2(2,N4))
        del34 = coef34*(del2(2,N3) - del2(2,N4))

        dU(2,N1) = dU(2,N1) + (-del12 - del13 - del14)*deltN(N1)
        dU(2,N2) = dU(2,N2) + ( del12 - del23 - del24)*deltN(N2)
        dU(2,N3) = dU(2,N3) + ( del13 + del23 - del34)*deltN(N3)
        dU(2,N4) = dU(2,N4) + ( del14 + del24 + del34)*deltN(N4)

        del12 = coef12*(del2(3,N1) - del2(3,N2))
        del13 = coef13*(del2(3,N1) - del2(3,N3))
        del14 = coef14*(del2(3,N1) - del2(3,N4))
        del23 = coef23*(del2(3,N2) - del2(3,N3))
        del24 = coef24*(del2(3,N2) - del2(3,N4))
        del34 = coef34*(del2(3,N3) - del2(3,N4))

        dU(3,N1) = dU(3,N1) + (-del12 - del13 - del14)*deltN(N1)
        dU(3,N2) = dU(3,N2) + ( del12 - del23 - del24)*deltN(N2)
        dU(3,N3) = dU(3,N3) + ( del13 + del23 - del34)*deltN(N3)
        dU(3,N4) = dU(3,N4) + ( del14 + del24 + del34)*deltN(N4)

        del12 = coef12*(del2(4,N1) - del2(4,N2))

```

```

        del13 = coef13*(del2(4,N1) - del2(4,N3))
        del14 = coef14*(del2(4,N1) - del2(4,N4))
        del23 = coef23*(del2(4,N2) - del2(4,N3))
        del24 = coef24*(del2(4,N2) - del2(4,N4))
        del34 = coef34*(del2(4,N3) - del2(4,N4))

        dU(4,N1) = dU(4,N1) + (-del12 - del13 - del14)*deltN(N1)
        dU(4,N2) = dU(4,N2) + ( del12 - del23 - del24)*deltN(N2)
        dU(4,N3) = dU(4,N3) + ( del13 + del23 - del34)*deltN(N3)
        dU(4,N4) = dU(4,N4) + ( del14 + del24 + del34)*deltN(N4)

        enddo
    enddo

    return
end

C*****
C*
C*   this subroutine accounts for periodic nodes
C*
C*****

        subroutine bperiodic
        implicit none

        include 'QUAD.INC'
        integer PN          !pointer
        integer P1, P2      !periodic nodes

CVD$  NODEPCHK
        do PN = 1, Pmax
            P1 = pnode(PN,1)
            P2 = pnode(PN,2)
            dU(1,P1) = dU(1,P1) + dU(1,P2)
            dU(2,P1) = dU(2,P1) + dU(2,P2)
            dU(3,P1) = dU(3,P1) + dU(3,P2)
            dU(4,P1) = dU(4,P1) + dU(4,P2)
            dU(1,P2) = dU(1,P1)
            dU(2,P2) = dU(2,P1)
            dU(3,P2) = dU(3,P1)
            dU(4,P2) = dU(4,P1)
        enddo

        return
    end

C*****
C*
C*   adjust inlet state vector for boundary condition
C*
C*****

        subroutine binlet
        implicit none

        include 'QUAD.INC'
        real rrinl, uuinl, vvlinl, ppinl !average values at inlet
        real ww2inl, aainl
        real rOinf, aOinf                !stag. density and speed of sound
        real HOpres                       !stagnation enthalpy and pressure
        real spres                        !entropy
        real coef
        real aaa1, aaa2, aaa3, aaa4      !coef. of inverse of matrix

```

```

real bbb1, bbb2, bbb3, bbb4
real ccc1, ccc2, ccc3, ccc4
real ddd1, ddd2, ddd3, ddd4
real ww2, uu, vv, pp, rr, aa      !values at next interior node
real dHO, ds, dtan, dw4
real drr, duu, dvv, dpp
integer IN, NN                    !boundary face and its nodes

c** average values at inlet
rrin1 = U(1,innode(Imax/2))
uuin1 = U(2,innode(Imax/2))/rrin1
vvin1 = U(3,innode(Imax/2))/rrin1
ww2in1 = uuin1**2 + vvin1**2
ppin1 = gam1*(U(4,innode(Imax/2))
&      - 0.5*rrin1*ww2in1)
aain1 = sqrt(gam*ppin1/rrin1)

c* subsonic inlet nodes
if ((sqrt(ww2in1)/aain1).lt.1. .or. pitch.ne.0.) then
c** prescribed values
HOpres = 1./gam1
spres = 0.

c** coefficient of matrix
coef = 1./(uuin1*(aain1+uuin1) + vvin1**2)
aaa1 = (rrin1*uuin1/aain1)*coef
aaa2 = -(ppin1/aain1)*(uuin1*gam/gam1 + ww2in1/aain1)*coef
aaa3 = -(rrin1*vvin1/aain1)*coef
aaa4 = (ww2in1/aain1**2)*coef
bbb1 = uuin1*coef
bbb2 = -(uuin1*ppin1/(gam1*rrin1))*coef
bbb3 = -vvin1*coef
bbb4 = -(uuin1/rrin1)*coef
ccc1 = vvin1*coef
ccc2 = -(vvin1*ppin1/(gam1*rrin1))*coef
ccc3 = (aain1 + uuin1)*coef
ccc4 = -(vvin1/rrin1)*coef
ddd1 = rrin1*aain1*uuin1*coef
ddd2 = -(ppin1*aain1*uuin1/gam1)*coef
ddd3 = -rrin1*aain1*vvin1*coef
ddd4 = ww2in1*coef

CVD$ NODEPCHK
do IN = 1, Imax
  NN = innode(IN)
  rr = U(1,NN)
  uu = U(2,NN)/rr
  vv = U(3,NN)/rr
  ww2 = uu**2 + vv**2
  pp = gam1*(U(4,NN) - 0.5*rr*ww2)
  aa = sqrt(gam*pp/rr)
  dHO = HOpres - ((gam/gam1)*pp/rr + 0.5*ww2)
  ds = spres - (log(gam*pp) - gam*log(rr))
  dtan = (Sin1 - vv/uu)*uuin1**2
  dw4 = dU(1,NN)*(aa*uu + gam1*ww2)
&      - dU(2,NN)*(aa + gam1*uu)
&      - dU(3,NN)*gam1*vv
&      + dU(4,NN)*gam1
  drr = aaa1*dHO + aaa2*ds + aaa3*dtan + aaa4*dw4
  duu = bbb1*dHO + bbb2*ds + bbb3*dtan + bbb4*dw4
  dvv = ccc1*dHO + ccc2*ds + ccc3*dtan + ccc4*dw4
  dpp = ddd1*dHO + ddd2*ds + ddd3*dtan + ddd4*dw4

  dU(1,NN) = drr
  dU(2,NN) = rr*duu + uu*drr
  dU(3,NN) = rr*dvv + vv*drr
  dU(4,NN) = dpp/gam1 + 0.5*ww2*drr + rr*(uu*duu + vv*dvv)
enddo

c* supersonic inlet nodes
else if ((sqrt(ww2in1)/aain1).ge.1.) then
c* state vector components far from body

```

```

      r0inf = 1.
      a0inf = 1.

CVD$  NODEPCHK
      do IN = 1, Imax
        NN = innode(IN)
        rr = r0inf*(1.0+0.5*gam1*Minl**2)**(-1./gam1)
        aa = a0inf*(1.0+0.5*gam1*Minl**2)**(-0.5)
        uu = Minl*aa
        pp = rr*aa**2/gam

        dU(1,NN) = rr - U(1,NN)
        dU(2,NN) = rr*uu - U(2,NN)
        dU(3,NN) = - U(3,NN)
        dU(4,NN) = pp/gam1 + .5*(uu**2)*rr - U(4,NN)
      enddo
    endif

    return
  end

C*****
C*
C*      adjust      clst state vector for boundary condition
C*
C*****

      subroutine boutlet
      implicit none

      include 'QUAD.INC'
      real rrout, uuout, vvout, ppout !average values at outlet
      real aaout, ww2out
      real aaa1, aaa4                !coef. of inverse of matrix
      real bbb3, bbb4
      real ccc2
      real ddd4
      real ww2, uu, vv, pp, rr, aa   !values at next interior node
      real dw1, dw2, dw3, dp
      real drr, duu, dvv, dpp
      integer ON, NN                 !boundary face and its nodes

      rrout = U(1,outnode(Onmax/2))
      uuout = U(2,outnode(Onmax/2))/rrout
      vvout = U(3,outnode(Onmax/2))/rrout
      ww2out = uuout**2 + vvout**2
      ppout = gam1*(U(4,outnode(Onmax/2))
      *      - 0.5*rrout*ww2out)
      aaout = sqrt(gam*ppout/rrout)

      aaa1 = -1./aaout**2
      aaa4 = 1./aaout**2
      bbb3 = .1/(rrout*aaout)
      bbb4 = -1./(rrout*aaout)
      ccc2 = .1/(rrout*aaout)
      ddd4 = 1.

C* set boundary values of state vector for nodes
CVD$  NODEPCHK
      do ON = 1, Onmax
        NN = outnode(ON)
        rr = U(1,NN)
        uu = U(2,NN)/rr
        vv = U(3,NN)/rr
        ww2 = uu**2 + vv**2
        pp = gam1*(U(4,NN) - 0.5*rr*ww2)
        aa = sqrt(gam*pp/rr)
        if (sqrt(ww2) .lt. aa) then
          dw1 = dU(1,NN)*(0.5*ww2*gam1 - aa**2)

```

```

&      - dU(2,NN)*gam1*uu
&      - dU(3,NN)*gam1*vv
&      + dU(4,NN)*gam1
dw2 = - dU(1,NN)*aa*vv
&      + dU(3,NN)*aa
dw3 = dU(1,NN)*(0.5*ww2*gam1 - aa*uu)
&      - dU(2,NN)*(gam1*uu - aa)
&      - dU(3,NN)*gam1*vv
&      + dU(4,NN)*gam1
dp = pout - pp
drr = aaal*dw1 + aaa4*dp
duu = bbb3*dw3 + bbb4*dp
dvv = ccc2*dw2
dpp = ddd4*dp
dU(1,NN) = drr
dU(2,NN) = rr*duu + uu*drr
dU(3,NN) = rr*dvv + vv*drr
dU(4,NN) = dpp/gam1 + 0.5*ww2*drr + rr*(uu*duu + vv*dvv)

endif
enddo

return
end

C*****
C*
C*      this subroutine changes the momentum change to make
C*      flow tangent to wall
C*
C*****

subroutine tangent
implicit none

include 'QUAD.INC'
integer EN          !pointer
integer NN          !node on wall
real drwn           !change in momentum normal to wall

CVD$ NODEPCHK
do EN = 1, bnode
  NN = enode(EN)
  drwn = -(U(2,NN) + dU(2,NN))*senode(EN) +
&        (U(3,NN) + dU(3,NN))*cenode(EN)
  dU(2,NN) = dU(2,NN) + drwn*senode(EN)
  dU(3,NN) = dU(3,NN) - drwn*cenode(EN)
enddo

return
end

```

A.2.4 Jameson Scheme

```

C*****
C*
C*      main program for quadrilateral Jameson scheme
C*
C*****

program quadrilateral
implicit none

```

```

include 'QUAD.INC'
integer Niter                      !number of iterations
real maxchange                    !max change in state vector
integer maxnode, maxeqn           !where max change occurs
integer CC                        !pointer

c* read in data from file
call gridio(1)
call flowio(1)
call input

durms = 999.
Niter = 0

c* start history file from the top
open(unit=35,status='unknown',form='formatted')
write(35,2) Minl
close(unit=35)
2   format('inlet Mach number = ',f5.3)

c* loop until converged
do while ((Niter.lt.Maxiter) .and. (durms.gt.2.e-7))
  Niter = Niter + 1

c* calculate value at next time step
call update(maxchange, maxnode, maxeqn)

  if (mod(Niter,10).eq.0 .or. Niter.lt.10) then
    call flowio(0)
    open(unit=50, status='unknown', form='unformatted')
    write(50) Cmax, (vol(CC),CC=1,Cmax)
    close(unit=50)

10   open(unit=35, status='old',access='append',err=10)
    write(35,1) Niter, durms, maxchange, x(maxnode),
    &          y(maxnode), maxeqn
    close(unit=35)
    write(6,1) Niter, durms, maxchange, x(maxnode),
    &          y(maxnode), maxeqn
    &          endif

1   format('Niter=',i4,' rms=',f9.7,' max=',f9.7,' x=',f6.3,
    &      ' y=',f6.3,' eqn=',i1)

enddo

c* write out data to file
call flowio(0)

stop
end

c*****
c*
c*   update state vectors at next time step
c*
c*****

subroutine update(maxchange, maxnode, maxeqn)

include 'QUAD.INC'
integer NN, 1                      !pointer
real UO(4,Maxnodes)              !starting values of state vector
real alpha1, alpha2              !coefficients
real alpha3, alpha4
real maxchange                    !max change in state vector
integer maxnode, maxeqn          !where max change occurs

```

```

c* set coefficient values
  alpha1 = 0.25
  alpha2 = 1./3.
  alpha3 = 0.5
  alpha4 = 1.

c* find timestep at each node
  call timestep

c* advance to next time step in four steps

c* step one
  call calcflux
  call dissipation
  do NN = 1, Nmax
    UO(1,NN) = U(1,NN)
    UO(2,NN) = U(2,NN)
    UO(3,NN) = U(3,NN)
    UO(4,NN) = U(4,NN)
    dU(1,NN) = U(1,NN)
    dU(2,NN) = U(2,NN)
    dU(3,NN) = U(3,NN)
    dU(4,NN) = U(4,NN)
    U(1,NN) = UO(1,NN) - alpha1*deltN(NN)
    &      *(flux(1,NN)-dis(1,NN))
    U(2,NN) = UO(2,NN) - alpha1*deltN(NN)
    &      *(flux(2,NN)-dis(2,NN))
    U(3,NN) = UO(3,NN) - alpha1*deltN(NN)
    &      *(flux(3,NN)-dis(3,NN))
    U(4,NN) = UO(4,NN) - alpha1*deltN(NN)
    &      *(flux(4,NN)-dis(4,NN))
    dU(1,NN) = U(1,NN) - dU(1,NN)
    dU(2,NN) = U(2,NN) - dU(2,NN)
    dU(3,NN) = U(3,NN) - dU(3,NN)
    dU(4,NN) = U(4,NN) - dU(4,NN)
  enddo
  call binlet
  call boutlet
  call tangent

c* step two
  call calcflux
  call dissipation
  do NN = 1, Nmax
    dU(1,NN) = U(1,NN)
    dU(2,NN) = U(2,NN)
    dU(3,NN) = U(3,NN)
    dU(4,NN) = U(4,NN)
    U(1,NN) = UO(1,NN) - alpha2*deltN(NN)
    &      *(flux(1,NN)-dis(1,NN))
    U(2,NN) = UO(2,NN) - alpha2*deltN(NN)
    &      *(flux(2,NN)-dis(2,NN))
    U(3,NN) = UO(3,NN) - alpha2*deltN(NN)
    &      *(flux(3,NN)-dis(3,NN))
    U(4,NN) = UO(4,NN) - alpha2*deltN(NN)
    &      *(flux(4,NN)-dis(4,NN))
    dU(1,NN) = U(1,NN) - dU(1,NN)
    dU(2,NN) = U(2,NN) - dU(2,NN)
    dU(3,NN) = U(3,NN) - dU(3,NN)
    dU(4,NN) = U(4,NN) - dU(4,NN)
  enddo
  call binlet
  call boutlet
  call tangent

c* step three
  call calcflux
  do NN = 1, Nmax
    dU(1,NN) = U(1,NN)
    dU(2,NN) = U(2,NN)
    dU(3,NN) = U(3,NN)

```



```

        dU(4,NN) = U(4,NN)
        U(1,NN) = UO(1,NN) - alpha3*deltN(NN)
        * (flux(1,NN)-dis(1,NN))
    &
        U(2,NN) = UO(2,NN) - alpha3*deltN(NN)
        * (flux(2,NN)-dis(2,NN))
    &
        U(3,NN) = UO(3,NN) - alpha3*deltN(NN)
        * (flux(3,NN)-dis(3,NN))
    &
        U(4,NN) = UO(4,NN) - alpha3*deltN(NN)
        * (flux(4,NN)-dis(4,NN))
    &
        dU(1,NN) = U(1,NN) - dU(1,NN)
        dU(2,NN) = U(2,NN) - dU(2,NN)
        dU(3,NN) = U(3,NN) - dU(3,NN)
        dU(4,NN) = U(4,NN) - dU(4,NN)
    enddo
    call binlet
    call boutlet
    call tangent

c* step four
    call calcflux
    do NN = 1, Nmax
        dU(1,NN) = U(1,NN)
        dU(2,NN) = U(2,NN)
        dU(3,NN) = U(3,NN)
        dU(4,NN) = U(4,NN)
        U(1,NN) = UO(1,NN) - alpha4*deltN(NN)
        * (flux(1,NN)-dis(1,NN))
    &
        U(2,NN) = UO(2,NN) - alpha4*deltN(NN)
        * (flux(2,NN)-dis(2,NN))
    &
        U(3,NN) = UO(3,NN) - alpha4*deltN(NN)
        * (flux(3,NN)-dis(3,NN))
    &
        U(4,NN) = UO(4,NN) - alpha4*deltN(NN)
        * (flux(4,NN)-dis(4,NN))
    &
        dU(1,NN) = U(1,NN) - U(1,NN)
        dU(2,NN) = U(2,NN) - U(2,NN)
        dU(3,NN) = U(3,NN) - U(3,NN)
        dU(4,NN) = U(4,NN) - U(4,NN)
    enddo
    call binlet
    call boutlet
    call tangent

c* find root mean square difference in state vector
    durms = 0.0
    maxchange = 0.

    do i = 1, 4
        do NN = 1, Nmax
            durms = durms + (U(1,NN) - UO(1,NN))**2
            if (abs(U(1,NN)-UO(1,NN)).gt.abs(maxchange)) then
                maxchange = U(1,NN) - UO(1,NN)
                maxnode = NN
                maxeqn = 1
            endif
        enddo
    enddo

    durms = sqrt(durms/(4.*Nmax))

    return
end

C*****
C*
C*      calculate time step for nodes
C*
C*****

```

```

subroutine timestep
implicit none

include 'QUAD.INC'
integer CC, NN                                !pointer
integer N1, N2, N3, N4                       !nodes related to face
integer PN, P1, P2                           !periodic nodes
real dx1, dy1, d1
real dxm, dym, dm
real uu, vv, aa, rr, vv2, pp                !values at node
real time1, time2

c* zero out delt
do NN = 1, Nmax
    deltN(NN) = 0.
enddo

c* find time step for each cell
do CC = 1, Cmax
    N1 = cell(CC,5)
    N2 = cell(CC,6)
    N3 = cell(CC,7)
    N4 = cell(CC,8)
    dx1 = .5*(x(N3) + x(N2) - x(N4) - x(N1))
    dy1 = .5*(y(N3) + y(N2) - y(N4) - y(N1))
    d1 = sqrt(dx1**2 + dy1**2)
    dxm = .5*(x(N4) + x(N3) - x(N1) - x(N2))
    dym = .5*(y(N4) + y(N3) - y(N1) - y(N2))
    dm = sqrt(dxm**2 + dym**2)
    rr = 0.25*(U(1,N1) + U(1,N2) + U(1,N3) + U(1,N4))
    uu = 0.25*(U(2,N1) + U(2,N2) + U(2,N3) + U(2,N4))/rr
    vv = 0.25*(U(3,N1) + U(3,N2) + U(3,N3) + U(3,N4))/rr
    pp = gam1*(0.25*(U(4,N1) + U(4,N2) + U(4,N3) + U(4,N4))
    & - 0.5*rr*(uu**2 + vv**2))
    aa = sqrt(gam*pp/rr)

    time1 = CFL/(abs(uu*dy1-vv*dx1) + aa*d1)
    time2 = CFL/(abs(uu*dym-vv*dxm) + aa*dm)

    deltC(CC) = min(time1,time2)

    deltN(N1) = deltN(N1) + 1./deltC(CC)
    deltN(N2) = deltN(N2) + 1./deltC(CC)
    deltN(N3) = deltN(N3) + 1./deltC(CC)
    deltN(N4) = deltN(N4) + 1./deltC(CC)
enddo

c** find time step at periodic nodes
CVD$ NODEPCHK
do PN = 1, Pmax
    P1 = pnode(PN,1)
    P2 = pnode(PN,2)
    deltN(P1) = deltN(P1) + deltN(P2)
    deltN(P2) = deltN(P1)
enddo

c** delt is actually delt/areaH
do NN = 1, Nmax
    deltN(NN) = 1./deltN(NN)
enddo

return
end

```

```

c*                                     *
c*      calculate flux vector values at nodes                                     *
c*                                     *
c*****
      subroutine calcflux
      implicit none

      include 'QUAD.INC'
      real delflux
      integer CC, NN
      integer N1, N2, N3, N4          !pointers
                                      !nodes around edge
      real WW
      real dy31, dy24, dx31, dx24    !change in x and y
      real dF24, dF13, dG24, dG13    !change in F and G

      do NN = 1, Nmax
c*   set flux to zero
          flux(1,NN) = 0.
          flux(2,NN) = 0.
          flux(3,NN) = 0.
          flux(4,NN) = 0.

c*   calculate f and g at all nodes
          WW = 0.5*(U(2,NN)**2 + U(3,NN)**2)/U(1,NN)
          F(1,NN) = U(2,NN)
          G(2,NN) = U(2,NN)*U(3,NN)/U(1,NN)
          F(2,NN) = U(2,NN)**2/U(1,NN) + gam1*(U(4,NN) - WW)
          F(3,NN) = U(2,NN)*U(3,NN)/U(1,NN)
          F(4,NN) = (U(2,NN)/U(1,NN))*(gam*U(4,NN) - gam1*WW)

          G(1,NN) = U(3,NN)
          G(2,NN) = U(2,NN)*U(3,NN)/U(1,NN)
          G(3,NN) = U(3,NN)**2/U(1,NN) + gam1*(U(4,NN) - WW)
          G(4,NN) = (U(3,NN)/U(1,NN))*(gam*U(4,NN) - gam1*WW)
      enddo

c*   find change in state vector for cell
      do CC = 1, Cmax
          N1 = cell(CC,5)
          N2 = cell(CC,6)
          N3 = cell(CC,7)
          N4 = cell(CC,8)

          dy31 = y(N3) - y(N1)
          dy24 = y(N2) - y(N4)
          dx31 = x(N3) - x(N1)
          dx24 = x(N2) - x(N4)

          dF24 = F(1,N2) - F(1,N4)
          dF13 = F(1,N1) - F(1,N3)
          dG24 = G(1,N2) - G(1,N4)
          dG13 = G(1,N1) - G(1,N3)

          delflux = 0.5*(-dF24*dy31 + dG24*dx31 -
      *      dF13*dy24 + dG13*dx24)

          flux(1,N1) = flux(1,N1) - delflux
          flux(1,N2) = flux(1,N2) - delflux
          flux(1,N3) = flux(1,N3) - delflux
          flux(1,N4) = flux(1,N4) - delflux

          dF24 = F(2,N2) - F(2,N4)
          dF13 = F(2,N1) - F(2,N3)
          dG24 = G(2,N2) - G(2,N4)
          dG13 = G(2,N1) - G(2,N3)

          delflux = 0.5*(-dF24*dy31 + dG24*dx31 -
      *      dF13*dy24 + dG13*dx24)

          flux(2,N1) = flux(2,N1) - delflux
          flux(2,N2) = flux(2,N2) - delflux
          flux(2,N3) = flux(2,N3) - delflux

```

```

        flux(2,N4) = flux(2,N4) - delflux

        dF24 = F(3,N2) - F(3,N4)
        dF13 = F(3,N1) - F(3,N3)
        dG24 = G(3,N2) - G(3,N4)
        dG13 = G(3,N1) - G(3,N3)

        delflux = 0.5*(-dF24*dy31 + dG24*dx31 -
*               dF13*dy24 + dG13*dx24)

        flux(3,N1) = flux(3,N1) - delflux
        flux(3,N2) = flux(3,N2) - delflux
        flux(3,N3) = flux(3,N3) - delflux
        flux(3,N4) = flux(3,N4) - delflux

        dF24 = F(4,N2) - F(4,N4)
        dF13 = F(4,N1) - F(4,N3)
        dG24 = G(4,N2) - G(4,N4)
        dG13 = G(4,N1) - G(4,N3)

        delflux = 0.5*(-dF24*dy31 + dG24*dx31 -
*               dF13*dy24 + dG13*dx24)

        flux(4,N1) = flux(4,N1) - delflux
        flux(4,N2) = flux(4,N2) - delflux
        flux(4,N3) = flux(4,N3) - delflux
        flux(4,N4) = flux(4,N4) - delflux

    enddo

c*  implement wall boundary conditions
    call bwall

    return
end

c*****
c*
c*      this subroutine applies the wall boundary condition
c*
c*****
c*****
c*****

subroutine bwall
implicit none

include 'QUAD.INC'
real delflux
integer PN, EF, FF, CC !pointer
integer N1, N2         !nodes on edge
integer P1, P2
real coef
real dy12, dx12        !change in x and y
real pres1, pres2      !pressure
integer EF1, EF2, col  !coloring pointers

c*  upper and lower boundary
    EF2 = 0

    do col = 1, Eicolormax
        EF1 = EF2 + 1
        EF2 = EF1 - 1 + NEcolor(col)
        do EF = EF1, EF2

            FF = eface(EF)
            CC = face(FF,1)
            N1 = face(FF,3)
            N2 = face(FF,4)

```

```

dy12 = y(N1) - y(N2)
dx12 = x(N1) - x(N2)

pres1 = gam1*(U(4,N1) - 0.5*(U(2,N1)**2 + U(3,N1)**2)/
      U(1,N1))
pres2 = gam1*(U(4,N2) - 0.5*(U(2,N2)**2 + U(3,N2)**2)/
      U(1,N2))

delflux = 0.5*(-(F(1,N2) + F(1,N1))*dy12 +
      (G(1,N2) + G(1,N1))*dx12)

flux(1,N1) = flux(1,N1) - delflux
flux(1,N2) = flux(1,N2) - delflux

delflux = 0.5*(-(F(2,N2) + F(2,N1) - pres1 - pres2)*dy12 +
      (G(2,N2) + G(2,N1))*dx12)

flux(2,N1) = flux(2,N1) - delflux
flux(2,N2) = flux(2,N2) - delflux

delflux = 0.5*(-(F(3,N2) + F(3,N1))*dy12 +
      (G(3,N2) + G(3,N1) - pres1 - pres2)*dx12)

flux(3,N1) = flux(3,N1) - delflux
flux(3,N2) = flux(3,N2) - delflux

delflux = 0.5*(-(F(4,N2) + F(4,N1))*dy12 +
      (G(4,N2) + G(4,N1))*dx12)

flux(4,N1) = flux(4,N1) - delflux
flux(4,N2) = flux(4,N2) - delflux

enddo
enddo

c* account for periodic nodes
CVD$ NODEPCHK
do PN = 1, Pmax
  P1 = pnode(PN,1)
  P2 = pnode(PN,2)
  flux(1,P1) = flux(1,P1) + flux(1,P2)
  flux(2,P1) = flux(2,P1) + flux(2,P2)
  flux(3,P1) = flux(3,P1) + flux(3,P2)
  flux(4,P1) = flux(4,P1) + flux(4,P2)
  flux(1,P2) = flux(1,P1)
  flux(2,P2) = flux(2,P1)
  flux(3,P2) = flux(3,P1)
  flux(4,P2) = flux(4,P1)
enddo

return
end

c*****
c*
c* calculate the dissipation at each of the nodes for
c* the current values of the state vector
c*
c*****

subroutine dissipation
implicit none

include 'QUAD.INC'
integer N1, N2, N3, N4      !nodes at end of face
real pres1, pres2, pres3    !pressure at nodes
real pres4, pres            !
integer EF, FF, CC, NN      !pointers

```

```

integer EF1, EF2
integer PM, P1, P2                                !periodic nodes
real del2(4,Maxnodes)                             !second order changes
real eps1(Maxnodes)                               !dissipation coefficients
real del12, del13, del14
real del23, del24, del34
real coef12, coef13, coef14
real coef23, coef24, coef34
real coef1122, coef1133, coef1144
real coef2233, coef2244, coef3344
real coef
real dx31, dx24                                !change in x
real dx12, dx23, dx34, dx41
real dy31, dy24                                !change in y
real dy12, dy23, dy34, dy41
real dxC1, dxC2, dxC3, dxC4                    !change in x in cell
real dyC1, dyC2, dyC3, dyC4                    !change in y in cell
integer CC1, CC2, col                            !color pointers

do NM = 1, Nmax
  del2(1,NM) = 0.
  del2(2,NM) = 0.
  del2(3,NM) = 0.
  del2(4,NM) = 0.

  dis(1,NM) = 0.
  dis(2,NM) = 0.
  dis(3,NM) = 0.
  dis(4,NM) = 0.

  eps1(NM) = 0.
enddo

c** Mavriplis smoothing
if (sigE .eq. 0.) then

  CC2 = 0

  do col = 1, Ccolormax
    CC1 = CC2 + 1
    CC2 = CC1 - 1 + NCcolor(col)

CVD$ NODEPCHK
    do CC = CC1, CC2
      N1 = cell(CC,5)
      N2 = cell(CC,6)
      N3 = cell(CC,7)
      N4 = cell(CC,8)

      del2(1,N1) = del2(1,N1) +
&               (-3.*U(1,N1) + U(1,N2) +
&               U(1,N3) + U(1,N4))
      del2(1,N2) = del2(1,N2) +
&               (U(1,N1) - 3.*U(1,N2) +
&               U(1,N3) + U(1,N4))
      del2(1,N3) = del2(1,N3) +
&               (U(1,N1) + U(1,N2) -
&               3.*U(1,N3) + U(1,N4))
      del2(1,N4) = del2(1,N4) +
&               (U(1,N1) + U(1,N2) +
&               U(1,N3) - 3.*U(1,N4))

      del2(2,N1) = del2(2,N1) +
&               (-3.*U(2,N1) + U(2,N2) +
&               U(2,N3) + U(2,N4))
      del2(2,N2) = del2(2,N2) +
&               (U(2,N1) - 3.*U(2,N2) +
&               U(2,N3) + U(2,N4))
      del2(2,N3) = del2(2,N3) +
&               (U(2,N1) + U(2,N2) -
&               3.*U(2,N3) + U(2,N4))
      del2(2,N4) = del2(2,N4) +

```

```

&          (U(2,N1) + U(2,N2) +
&          U(2,N3) - 3.*U(2,N4))

      del12(3,N1) = del12(3,N1) +
&          (-3.*U(3,N1) + U(3,N2) +
&          U(3,N3) + U(3,N4))
      del12(3,N2) = del12(3,N2) +
&          (U(3,N1) - 3.*U(3,N2) +
&          U(3,N3) + U(3,N4))
      del12(3,N3) = del12(3,N3) +
&          (U(3,N1) + U(3,N2) -
&          3.*U(3,N3) + U(3,N4))
      del12(3,N4) = del12(3,N4) +
&          (U(3,N1) + U(3,N2) +
&          U(3,N3) - 3.*U(3,N4))

      del12(4,N1) = del12(4,N1) +
&          (-3.*U(4,N1) + U(4,N2) +
&          U(4,N3) + U(4,N4))
      del12(4,N2) = del12(4,N2) +
&          (U(4,N1) - 3.*U(4,N2) +
&          U(4,N3) + U(4,N4))
      del12(4,N3) = del12(4,N3) +
&          (U(4,N1) + U(4,N2) -
&          3.*U(4,N3) + U(4,N4))
      del12(4,N4) = del12(4,N4) +
&          (U(4,N1) + U(4,N2) +
&          U(4,N3) - 3.*U(4,N4))

      enddo
    enddo

CVD$ NODEPCHK
do PN = 1, Pmax
  P1 = pnode(PN,1)
  P2 = pnode(PN,2)
  del12(1,P1) = del12(1,P1) + del12(1,P2)
  del12(2,P1) = del12(2,P1) + del12(2,P2)
  del12(3,P1) = del12(3,P1) + del12(3,P2)
  del12(4,P1) = del12(4,P1) + del12(4,P2)
  del12(1,P2) = del12(1,P1)
  del12(2,P2) = del12(2,P1)
  del12(3,P2) = del12(3,P1)
  del12(4,P2) = del12(4,P1)
enddo

c** N1 scheme method of smoothing
  else if (sigE .eq. 1.) then

    CC2 = 0

    do col = 1, Ccolormax
      CC1 = CC2 + 1
      CC2 = CC1 - 1 + NCcolor(col)

CVD$ NODEPCHK
do CC = CC1, CC2
  N1 = cell(CC,5)
  N2 = cell(CC,6)
  N3 = cell(CC,7)
  N4 = cell(CC,8)

  dy24 = y(N2) - y(N4)
  dy31 = y(N3) - y(N1)
  dy12 = y(N1) - y(N2)
  dy23 = y(N2) - y(N3)
  dy34 = y(N3) - y(N4)
  dy41 = y(N4) - y(N1)

  dx24 = x(N2) - x(N4)
  dx31 = x(N3) - x(N1)
  dx12 = x(N1) - x(N2)

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```

dx23 = x(N2) - x(N3)
dx34 = x(N3) - x(N4)
dx41 = x(N4) - x(N1)

coef = 1./abs(-dx12*dy41 + dy12*dx41)

dxC1 = U(1,N1)*dy24 + U(1,N2)*dy41 + U(1,N4)*dy12
dxC2 = U(2,N1)*dy24 + U(2,N2)*dy41 + U(2,N4)*dy12
dxC3 = U(3,N1)*dy24 + U(3,N2)*dy41 + U(3,N4)*dy12
dxC4 = U(4,N1)*dy24 + U(4,N2)*dy41 + U(4,N4)*dy12

dyC1 = U(1,N1)*dx24 + U(1,N2)*dx41 + U(1,N4)*dx12
dyC2 = U(2,N1)*dx24 + U(2,N2)*dx41 + U(2,N4)*dx12
dyC3 = U(3,N1)*dx24 + U(3,N2)*dx41 + U(3,N4)*dx12
dyC4 = U(4,N1)*dx24 + U(4,N2)*dx41 + U(4,N4)*dx12

del2(1,N1) = del2(1,N1) - 0.5*coef*(dxC1*dy24 + dyC1*dx24)
del2(2,N1) = del2(2,N1) - 0.5*coef*(dxC2*dy24 + dyC2*dx24)
del2(3,N1) = del2(3,N1) - 0.5*coef*(dxC3*dy24 + dyC3*dx24)
del2(4,N1) = del2(4,N1) - 0.5*coef*(dxC4*dy24 + dyC4*dx24)

coef = 1./abs(-dx12*dy23 + dy12*dx23)

dxC1 = U(1,N1)*dy23 + U(1,N2)*dy31 + U(1,N3)*dy12
dxC2 = U(2,N1)*dy23 + U(2,N2)*dy31 + U(2,N3)*dy12
dxC3 = U(3,N1)*dy23 + U(3,N2)*dy31 + U(3,N3)*dy12
dxC4 = U(4,N1)*dy23 + U(4,N2)*dy31 + U(4,N3)*dy12

dyC1 = U(1,N1)*dx23 + U(1,N2)*dx31 + U(1,N3)*dx12
dyC2 = U(2,N1)*dx23 + U(2,N2)*dx31 + U(2,N3)*dx12
dyC3 = U(3,N1)*dx23 + U(3,N2)*dx31 + U(3,N3)*dx12
dyC4 = U(4,N1)*dx23 + U(4,N2)*dx31 + U(4,N3)*dx12

del2(1,N2) = del2(1,N2) - 0.5*coef*(dxC1*dy31 + dyC1*dx31)
del2(2,N2) = del2(2,N2) - 0.5*coef*(dxC2*dy31 + dyC2*dx31)
del2(3,N2) = del2(3,N2) - 0.5*coef*(dxC3*dy31 + dyC3*dx31)
del2(4,N2) = del2(4,N2) - 0.5*coef*(dxC4*dy31 + dyC4*dx31)

coef = 1./abs(-dx23*dy34 + dy23*dx34)

dxC1 = U(1,N4)*dy23 + U(1,N2)*dy34 - U(1,N3)*dy24
dxC2 = U(2,N4)*dy23 + U(2,N2)*dy34 - U(2,N3)*dy24
dxC3 = U(3,N4)*dy23 + U(3,N2)*dy34 - U(3,N3)*dy24
dxC4 = U(4,N4)*dy23 + U(4,N2)*dy34 - U(4,N3)*dy24

dyC1 = U(1,N4)*dx23 + U(1,N2)*dx34 - U(1,N3)*dx24
dyC2 = U(2,N4)*dx23 + U(2,N2)*dx34 - U(2,N3)*dx24
dyC3 = U(3,N4)*dx23 + U(3,N2)*dx34 - U(3,N3)*dx24
dyC4 = U(4,N4)*dx23 + U(4,N2)*dx34 - U(4,N3)*dx24

del2(1,N3) = del2(1,N3) + 0.5*coef*(dxC1*dy24 + dyC1*dx24)
del2(2,N3) = del2(2,N3) + 0.5*coef*(dxC2*dy24 + dyC2*dx24)
del2(3,N3) = del2(3,N3) + 0.5*coef*(dxC3*dy24 + dyC3*dx24)
del2(4,N3) = del2(4,N3) + 0.5*coef*(dxC4*dy24 + dyC4*dx24)

coef = 1./abs(-dx34*dy41 + dy34*dx41)

dxC1 = U(1,N1)*dy34 + U(1,N3)*dy41 - U(1,N4)*dy31
dxC2 = U(2,N1)*dy34 + U(2,N3)*dy41 - U(2,N4)*dy31
dxC3 = U(3,N1)*dy34 + U(3,N3)*dy41 - U(3,N4)*dy31
dxC4 = U(4,N1)*dy34 + U(4,N3)*dy41 - U(4,N4)*dy31

dyC1 = U(1,N1)*dx34 + U(1,N3)*dx41 - U(1,N4)*dx31
dyC2 = U(2,N1)*dx34 + U(2,N3)*dx41 - U(2,N4)*dx31
dyC3 = U(3,N1)*dx34 + U(3,N3)*dx41 - U(3,N4)*dx31
dyC4 = U(4,N1)*dx34 + U(4,N3)*dx41 - U(4,N4)*dx31

del2(1,N4) = del2(1,N4) + 0.5*coef*(dxC1*dy31 + dyC1*dx31)
del2(2,N4) = del2(2,N4) + 0.5*coef*(dxC2*dy31 + dyC2*dx31)
del2(3,N4) = del2(3,N4) + 0.5*coef*(dxC3*dy31 + dyC3*dx31)
del2(4,N4) = del2(4,N4) + 0.5*coef*(dxC4*dy31 + dyC4*dx31)

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        enddo
    enddo

    EF2 = 0

    do col = 1, Eicolormax
        EF1 = EF2 + 1
        EF2 = EF1 - 1 + NEcolor(col)

        do EF = EF1, EF2
            FF = eface(EF)
            CC = face(FF,1)
            N1 = face(FF,3)
            N2 = face(FF,4)
            N3 = face(FF,6)
            N4 = face(FF,5)

            dy24 = y(N2) - y(N4)
            dy31 = y(N3) - y(N1)
            dy12 = y(N1) - y(N2)
            dy23 = y(N2) - y(N3)
            dy41 = y(N4) - y(N1)

            dx24 = x(N2) - x(N4)
            dx31 = x(N3) - x(N1)
            dx12 = x(N1) - x(N2)
            dx23 = x(N2) - x(N3)
            dx41 = x(N4) - x(N1)

            coef = 1./abs(-dx12*dy41 + dy12*dx41)

            dxC1 = U(1,N1)*dy24 + U(1,N2)*dy41 + U(1,N4)*dy12
            dxC2 = U(2,N1)*dy24 + U(2,N2)*dy41 + U(2,N4)*dy12
            dxC3 = U(3,N1)*dy24 + U(3,N2)*dy41 + U(3,N4)*dy12
            dxC4 = U(4,N1)*dy24 + U(4,N2)*dy41 + U(4,N4)*dy12

            dyC1 = U(1,N1)*dx24 + U(1,N2)*dx41 + U(1,N4)*dx12
            dyC2 = U(2,N1)*dx24 + U(2,N2)*dx41 + U(2,N4)*dx12
            dyC3 = U(3,N1)*dx24 + U(3,N2)*dx41 + U(3,N4)*dx12
            dyC4 = U(4,N1)*dx24 + U(4,N2)*dx41 + U(4,N4)*dx12

            del2(1,N1) = del2(1,N1) + 0.5*coef*(dxC1*dy12 + dyC1*dx12)
            del2(2,N1) = del2(2,N1) + 0.5*coef*(dxC2*dy12 + dyC2*dx12)
            del2(3,N1) = del2(3,N1) + 0.5*coef*(dxC3*dy12 + dyC3*dx12)
            del2(4,N1) = del2(4,N1) + 0.5*coef*(dxC4*dy12 + dyC4*dx12)

            coef = 1./abs(-dx12*dy23 + dy12*dx23)

            dxC1 = U(1,N1)*dy23 + U(1,N2)*dy31 + U(1,N3)*dy12
            dxC2 = U(2,N1)*dy23 + U(2,N2)*dy31 + U(2,N3)*dy12
            dxC3 = U(3,N1)*dy23 + U(3,N2)*dy31 + U(3,N3)*dy12
            dxC4 = U(4,N1)*dy23 + U(4,N2)*dy31 + U(4,N3)*dy12

            dyC1 = U(1,N1)*dx23 + U(1,N2)*dx31 + U(1,N3)*dx12
            dyC2 = U(2,N1)*dx23 + U(2,N2)*dx31 + U(2,N3)*dx12
            dyC3 = U(3,N1)*dx23 + U(3,N2)*dx31 + U(3,N3)*dx12
            dyC4 = U(4,N1)*dx23 + U(4,N2)*dx31 + U(4,N3)*dx12

            del2(1,N2) = del2(1,N2) + 0.5*coef*(dxC1*dy12 + dyC1*dx12)
            del2(2,N2) = del2(2,N2) + 0.5*coef*(dxC2*dy12 + dyC2*dx12)
            del2(3,N2) = del2(3,N2) + 0.5*coef*(dxC3*dy12 + dyC3*dx12)
            del2(4,N2) = del2(4,N2) + 0.5*coef*(dxC4*dy12 + dyC4*dx12)

        enddo
    enddo

CVD$ NODEPCHK
    do PN = 1, Pmax
        P1 = pnode(PN,1)
        P2 = pnode(PN,2)
        del2(1,P1) = del2(1,P1) + del2(1,P2)
        del2(2,P1) = del2(2,P1) + del2(2,P2)
    enddo

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```

        del2(3,P1) = del2(3,P1) + del2(3,P2)
        del2(4,P1) = del2(4,P1) + del2(4,P2)
        del2(1,P2) = del2(1,P1)
        del2(2,P2) = del2(2,P1)
        del2(3,P2) = del2(3,P1)
        del2(4,P2) = del2(4,P1)
    enddo

endif

c* undivided Laplacian of pressure for eps1 coefficient
CC2 = 0

do col = 1, Ccolormax
    CC1 = CC2 + 1
    CC2 = CC1 - 1 + NCcolor(col)

cCVD$ NODEPCHK
    do CC = CC1, CC2
        N1 = cell(CC,5)
        N2 = cell(CC,6)
        N3 = cell(CC,7)
        N4 = cell(CC,8)

        pres1 = gam1*(U(4,N1) - 0.5*(U(2,N1)**2 +
&          U(3,N1)**2)/U(1,N1))
        pres2 = gam1*(U(4,N2) - 0.5*(U(2,N2)**2 +
&          U(3,N2)**2)/U(1,N2))
        pres3 = gam1*(U(4,N3) - 0.5*(U(2,N3)**2 +
&          U(3,N3)**2)/U(1,N3))
        pres4 = gam1*(U(4,N4) - 0.5*(U(2,N4)**2 +
&          U(3,N4)**2)/U(1,N4))

        eps1(N1) = eps1(N1) - 3.*pres1 + pres2 + pres3 + pres4
        eps1(N2) = eps1(N2) + pres1 - 3.*pres2 + pres3 + pres4
        eps1(N3) = eps1(N3) + pres1 + pres2 - 3.*pres3 + pres4
        eps1(N4) = eps1(N4) + pres1 + pres2 + pres3 - 3.*pres4
    enddo
enddo

CVD$ NODEPCHK
do PN = 1, Pmax
    Pi = pnode(PN,1)
    P2 = pnode(PN,2)
    eps1(P1) = eps1(P1) + eps1(P2)
    eps1(P2) = eps1(P1)
enddo

do NN = 1, Nmax
    pres = gam1*(U(4,NN) - 0.5*(U(2,NN)**2 +
&          U(3,NN)**2)/U(1,NN))
    eps1(NN) = abs(eps1(NN)/pres)
enddo

CC2 = 0

do col = 1, Ccolormax
    CC1 = CC2 + 1
    CC2 = CC1 - 1 + NCcolor(col)

CVD$ NODEPCHK
    do CC = CC1, CC2
        N1 = cell(CC,5)
        N2 = cell(CC,6)
        N3 = cell(CC,7)
        N4 = cell(CC,8)

        coef12 = 0.5*eps1coef*(eps1(N1) + eps1(N2))
        coef13 = 0.5*eps1coef*(eps1(N1) + eps1(N3))
        coef14 = 0.5*eps1coef*(eps1(N1) + eps1(N4))
        coef23 = 0.5*eps1coef*(eps1(N2) + eps1(N3))
        coef24 = 0.5*eps1coef*(eps1(N2) + eps1(N4))
    enddo
enddo

```

```

coef34 = 0.5*apsicoef*(eps1(N3) + eps1(N4))

coef1122 = max(0.,eps2-coef12)/deltC(CC)
coef1133 = max(0.,eps2-coef13)/deltC(CC)
coef1144 = max(0.,eps2-coef14)/deltC(CC)
coef2233 = max(0.,eps2-coef23)/deltC(CC)
coef2244 = max(0.,eps2-coef24)/deltC(CC)
coef3344 = max(0.,eps2-coef34)/deltC(CC)

del12 = coef12*(U(1,N1) - U(1,N2)) -
&      coef1122*(del2(1,N1) - del2(1,N2))
del13 = coef13*(U(1,N1) - U(1,N3)) -
&      coef1133*(del2(1,N1) - del2(1,N3))
del14 = coef14*(U(1,N1) - U(1,N4)) -
&      coef1144*(del2(1,N1) - del2(1,N4))
del23 = coef23*(U(1,N2) - U(1,N3)) -
&      coef2233*(del2(1,N2) - del2(1,N3))
del24 = coef24*(U(1,N2) - U(1,N4)) -
&      coef2244*(del2(1,N2) - del2(1,N4))
del34 = coef34*(U(1,N3) - U(1,N4)) -
&      coef3344*(del2(1,N3) - del2(1,N4))

dis(1,N1) = dis(1,N1) - del12 - del13 - del14
dis(1,N2) = dis(1,N2) + del12 - del23 - del24
dis(1,N3) = dis(1,N3) + del13 + del23 - del34
dis(1,N4) = dis(1,N4) + del14 + del24 + del34

del12 = coef12*(U(2,N1) - U(2,N2)) -
&      coef1122*(del2(2,N1) - del2(2,N2))
del13 = coef13*(U(2,N1) - U(2,N3)) -
&      coef1133*(del2(2,N1) - del2(2,N3))
del14 = coef14*(U(2,N1) - U(2,N4)) -
&      coef1144*(del2(2,N1) - del2(2,N4))
del23 = coef23*(U(2,N2) - U(2,N3)) -
&      coef2233*(del2(2,N2) - del2(2,N3))
del24 = coef24*(U(2,N2) - U(2,N4)) -
&      coef2244*(del2(2,N2) - del2(2,N4))
del34 = coef34*(U(2,N3) - U(2,N4)) -
&      coef3344*(del2(2,N3) - del2(2,N4))

dis(2,N1) = dis(2,N1) - del12 - del13 - del14
dis(2,N2) = dis(2,N2) + del12 - del23 - del24
dis(2,N3) = dis(2,N3) + del13 + del23 - del34
dis(2,N4) = dis(2,N4) + del14 + del24 + del34

del12 = coef12*(U(3,N1) - U(3,N2)) -
&      coef1122*(del2(3,N1) - del2(3,N2))
del13 = coef13*(U(3,N1) - U(3,N3)) -
&      coef1133*(del2(3,N1) - del2(3,N3))
del14 = coef14*(U(3,N1) - U(3,N4)) -
&      coef1144*(del2(3,N1) - del2(3,N4))
del23 = coef23*(U(3,N2) - U(3,N3)) -
&      coef2233*(del2(3,N2) - del2(3,N3))
del24 = coef24*(U(3,N2) - U(3,N4)) -
&      coef2244*(del2(3,N2) - del2(3,N4))
del34 = coef34*(U(3,N3) - U(3,N4)) -
&      coef3344*(del2(3,N3) - del2(3,N4))

dis(3,N1) = dis(3,N1) - del12 - del13 - del14
dis(3,N2) = dis(3,N2) + del12 - del23 - del24
dis(3,N3) = dis(3,N3) + del13 + del23 - del34
dis(3,N4) = dis(3,N4) + del14 + del24 + del34

del12 = coef12*(U(4,N1) - U(4,N2)) -
&      coef1122*(del2(4,N1) - del2(4,N2))
del13 = coef13*(U(4,N1) - U(4,N3)) -
&      coef1133*(del2(4,N1) - del2(4,N3))
del14 = coef14*(U(4,N1) - U(4,N4)) -
&      coef1144*(del2(4,N1) - del2(4,N4))
del23 = coef23*(U(4,N2) - U(4,N3)) -
&      coef2233*(del2(4,N2) - del2(4,N3))
del24 = coef24*(U(4,N2) - U(4,N4)) -

```

```

&      coef2244*(del2(4,N2) - del2(4,N4))
del34 = coef34*(U(4,N3) - U(4,N4)) -
&      coef3344*(del2(4,N3) - del2(4,N4))

dis(4,N1) = dis(4,N1) - del12 - del13 - del14
dis(4,N2) = dis(4,N2) + del12 - del13 - del14
dis(4,N3) = dis(4,N3) + del13 + del13 - del14
dis(4,N4) = dis(4,N4) + del14 + del14 + del14

enddo
enddo

CVD$ NODEPCHK
do PN = 1, Pmax
  P1 = pnode(PN,1)
  P2 = pnode(PN,2)
  dis(1,P1) = dis(1,P1) + dis(1,P2)
  dis(2,P1) = dis(2,P1) + dis(2,P2)
  dis(3,P1) = dis(3,P1) + dis(3,P2)
  dis(4,P1) = dis(4,P1) + dis(4,P2)
  dis(1,P2) = dis(1,P1)
  dis(2,P2) = dis(2,P1)
  dis(3,P2) = dis(3,P1)
  dis(4,P2) = dis(4,P1)
enddo

return
end

C*****
c*
c*      adjust inlet state vector for boundary condition
c*
C*****

subroutine binlet
implicit none

include 'QUAD.INC'
real rrinl, uuinl, vvinnl, ppinnl !average values at inlet
real ww2inl, aainl
real r0inf, a0inf                !stag. density and speed of sound
real HOpres                      !stagnation enthalpy and pressure
real spres                       !entropy
real coef
real aaa1, aaa2, aaa3, aaa4      !coef. of inverse of matrix
real bbb1, bbb2, bbb3, bbb4
real ccc1, ccc2, ccc3, ccc4
real ddd1, ddd2, ddd3, ddd4
real ww2, uu, vv, pp, rr, aa    !values at next interior node
real dHO, ds, dtan, dw4
real drr, duu, dvv, dpp
integer IN, NH                   !boundary face and its nodes

c** average values at inlet
rrinl = U(1,innode(Imax/2)) - dU(1,innode(Imax/2))
uuinl = (U(2,innode(Imax/2)) - dU(2,innode(Imax/2)))/rrinl
vvinnl = (U(3,innode(Imax/2)) - dU(3,innode(Imax/2)))/rrinl
ww2inl = uuinl**2 + vvinnl**2
ppinnl = gam1*((U(4,innode(Imax/2)) - dU(4,innode(Imax/2)))
&      - 0.5*rrinl*ww2inl)
aainl = sqrt(gam*ppinnl/rrinl)

c* subsonic inlet nodes
if ((sqrt(ww2inl)/aainl).lt.1. .or. pitch.ne.0.) then
c** prescribed values
  HOpres = 1./gam1

```

```

      spres = 0.

c** coefficient of matrix
      coef = 1./(uuinl*(aa1nl+uuinl) + vv1nl**2)
      aaa1 = (rr1nl+uu1nl/aa1nl)*coef
      aaa2 = -(pp1nl/aa1nl)*(uu1nl*gam/gam1 + vv21nl/aa1nl)*coef
      aaa3 = -(rr1nl+vv1nl/aa1nl)*coef
      aaa4 = (vv21nl/aa1nl**2)*coef
      bbb1 = uu1nl*coef
      bbb2 = -(uu1nl*pp1nl/(gam1+rr1nl))*coef
      bbb3 = -vv1nl*coef
      bbb4 = -(uu1nl/rr1nl)*coef
      ccc1 = vv1nl*coef
      ccc2 = -(vv1nl*pp1nl/(gam1+rr1nl))*coef
      ccc3 = (aa1nl + uu1nl)*coef
      ccc4 = -(vv1nl/rr1nl)*coef
      ddd1 = rr1nl*aa1nl+uu1nl*coef
      ddd2 = -(pp1nl*aa1nl+uu1nl/gam1)*coef
      ddd3 = -rr1nl*aa1nl+vv1nl*coef
      ddd4 = vv21nl*coef

CVD$ NODEPCHK
      do IN = 1, Imax
        NN = innode(IN)
        rr = U(1,NN) - dU(1,NN)
        uu = (U(2,NN) - dU(2,NN))/rr
        vv = (U(3,NN) - dU(3,NN))/rr
        ww2 = uu**2 + vv**2
        pp = gam1*((U(4,NN) - dU(4,NN)) - 0.5*rr+ww2)
        aa = sqrt(gam*pp/rr)
        dHO = HOpres - ((gam/gam1)*pp/rr + 0.5*ww2)
        ds = spres - (log(gam*pp) - gam*log(rr))
        dtan = (S1nl - vv/uu)*uuinl**2
        dw4 = dU(1,NN)*(aa+uu + gam1*ww2)
        &          - dU(2,NN)*(aa + gam1*uu)
        &          - dU(3,NN)*gam1*vv
        &          + dU(4,NN)*gam1
        drr = aaa1*dHO + aaa2*ds + aaa3*dtan + aaa4*dw4
        duu = bbb1*dHO + bbb2*ds + bbb3*dtan + bbb4*dw4
        dvv = ccc1*dHO + ccc2*ds + ccc3*dtan + ccc4*dw4
        dpp = ddd1*dHO + ddd2*ds + ddd3*dtan + ddd4*dw4

        U(1,NN) = U(1,NN) - dU(1,NN) + drr
        U(2,NN) = U(2,NN) - dU(2,NN) + rr*duu + uu*drr
        U(3,NN) = U(3,NN) - dU(3,NN) + rr*dvv + vv*drr
        U(4,NN) = U(4,NN) - dU(4,NN) + dpp/gam1 + 0.5*ww2+drr
        &          + rr*(uu*duu + vv*dvv)
      enddo

c* supersonic inlet nodes
      else if ((sqrt(vv21nl)/aa1nl).ge.1.) then
c* state vector components far from body
        r0inf = 1.
        a0inf = 1.

CVD$ NODEPCHK
        do IN = 1, Imax
          NN = innode(IN)
          rr = r0inf*(1.0+0.5*gam1*Minl**2)**(-1./gam1)
          aa = a0inf*(1.0+0.5*gam1*Minl**2)**(-0.5)
          uu = Minl*aa
          pp = rr*aa**2/gam1

          U(1,NN) = rr
          U(2,NN) = rr*uu
          U(3,NN) = 0.
          U(4,NN) = pp/gam1 + .5*(uu**2)*rr
        enddo
      endif

      return
      end

```

```

c*****
c*
c*      adjust outlet state vector for boundary condition
c*
c*****

      subroutine boutlet
      implicit none

      include 'QUAD.INC'
      real rrout, uuout, vvout, ppout !average values at outlet
      real aaout, ww2out
      real aaa1, aaa4                !coef. of inverse of matrix
      real bbb3, bbb4
      real ccc2
      real ddd4
      real ww2, uu, vv, pp, rr, aa    !values at next interior node
      real dw1, dw2, dw3, dp
      real drr, duu, dvv, dpp
      integer ON, NN                  !boundary face and its nodes

      rrout = U(1,outnode(Onmax/2)) - dU(1,outnode(Onmax/2))
      uuout = (U(2,outnode(Onmax/2)) - dU(2,outnode(Onmax/2)))/rrout
      vvout = (U(3,outnode(Onmax/2)) - dU(3,outnode(Onmax/2)))/rrout
      ww2out = uuout**2 + vvout**2
      ppout = gam1*((U(4,outnode(Onmax/2)) - dU(4,outnode(Onmax/2)))
      & - 0.5*rrout*ww2out)
      aaout = sqrt(gam*ppout/rrout)

      aaa1 = -1./aaout**2
      aaa4 = 1./aaout**2
      bbb3 = .1/(rrout*aaout)
      bbb4 = -1./(rrout*aaout)
      ccc2 = .1/(rrout*aaout)
      ddd4 = 1.

c* set boundary values of state vector for nodes
CVD$ NODEPCHK
      do ON = 1, Onmax
      NN = outnode(ON)
      rr = U(1,NN) - dU(1,NN)
      uu = (U(2,NN) - dU(2,NN))/rr
      vv = (U(3,NN) - dU(3,NN))/rr
      ww2 = uu**2 + vv**2
      pp = gam1*((U(4,NN) - dU(4,NN)) - 0.5*rr*ww2)
      aa = sqrt(gam*pp/rr)
      if (sqrt(ww2) .lt. aa) then
      &      dw1 = dU(1,NN)*(0.5*ww2*gam1 - aa**2)
      &      dw2 = dU(2,NN)*gam1*uu
      &      dw3 = dU(3,NN)*gam1*vv
      &      dw4 = dU(4,NN)*gam1
      &      dw2 = - dU(1,NN)*aa*vv
      &      dw3 = dU(1,NN)*(0.5*ww2*gam1 - aa*uu)
      &      dw4 = dU(2,NN)*(gam1*uu - aa)
      &      dw5 = dU(3,NN)*gam1*vv
      &      dw6 = dU(4,NN)*gam1
      &      dp = pout - pp
      &      drr = aaa1*dw1 + aaa4*dp
      &      duu = bbb3*dw3 + bbb4*dp
      &      dvv = ccc2*dw2
      &      dpp = ddd4*dp
      &      U(1,NN) = U(1,NN) - dU(1,NN) + drr
      &      U(2,NN) = U(2,NN) - dU(2,NN) + rr*duu + uu*drr
      &      U(3,NN) = U(3,NN) - dU(3,NN) + rr*dvv + vv*drr
      &      U(4,NN) = U(4,NN) - dU(4,NN) + dpp/gam1 + 0.5*ww2*drr
      &      + rr*(uu*duu + vv*dvv)
      &
      endif
      enddo

```

```

        enddo

        return
    end

c*****
c*
c*      this subroutine changes the momentum change to make      *
c*      flow tangent to wall                                     *
c*                                                                 *
c*****

        subroutine tangent
        implicit none

        include 'QUAD.INC'
        integer EN                      !pointer
        integer NN                      !node on wall
        real drwn                      !change in momentum normal to wall

CVD$  NODEPCHK
        do EN = 1, bnode
            NN = enode(EN)
            drwn = -U(2,NN)*senode(EN) + U(3,NN)*cenode(EN)
            U(2,NN) = U(2,NN) + drwn*senode(EN)
            U(3,NN) = U(3,NN) - drwn*cenode(EN)
        enddo

        return
    end

```

A.2.5 Plotting Package

```

program plotgen
implicit none

include 'QUAD.INC'

integer i
integer NN, EN, CC
integer ptype                      !type of plot
integer ctype                      !type of contour
integer stype                      !type of surface distributions
integer N1, N2, N3, N4
character*80 TITLE, ITITLE
integer NTITL                      !number of letters in title
external g2pltg, g2pltc           !plotting subroutines
integer indgr
integer a4, a5, a6, a7, a8, a9, a10
real uu, vv, pp, rr, mm, mm2      !from state vectors
real pt, ptinf
real zz, z(Maxnodes)             !contour values
real zmax, zmin                  !max and min of z array
real boundary                    !boundary on contour levels
integer NCONT                    !contour level info
real CBASE, CSTEP
real Cinc                        !contour increment
character*6 NUM                   !same as CSTEP but character
integer NLINE, IOPT(2)           !indicators for line plots
real tau, radius2                !define bump geometry
integer points, npts(2)          !number of points on line
real xline(Maxedges)             !points on line to plot

```

```

      real yline(Maxedges)

c* read data from file
      call gridio(i)
      call flowio(i)

c* initialize GRAFIC
      write(1,1) Min1
1      format('INLET MACH NUMBER = ',F5.3)
      call grinit(5,6,1,TITLE)

      do while (1)

c* prompt user for type of plot
      type*, 'Type of plot'
      type*, ' 0) STOP'
      type*, ' 1) grid'
      type*, ' 2) contour'
      type*, ' 3) surface distribution'
      type*, ' 4) data'
      type 11
11      format(1,' selection = ')
      accept 111, ptype
111     format(I)

      if (ptype.eq.0) then
        stop
      else if (ptype.eq.1) then
        TITLE = ' X Y COMPUTATIONAL GRID'
        indgr = 23
        call gr_control(g2pltg, indgr, TITLE, x, y, Nmax, a4,
          & a5, a6, a7, a8, a9, a10)

      else if (ptype.eq.2) then
c* choose type of contour plot
        do while (1)

          Cinc = 0.

2          type*, 'Type of contour'
          type*, ' 0) TOP LEVEL'
          type*, ' 1) density'
          type*, ' 2) Mach number'
          type*, ' 3) normal velocity'
          type*, ' 4) pressure'
          type*, ' 5) total pressure loss'
          type*, ' 6) entropy'
          type*, ' 7) linear (not related to solution)'
          type*, ' 8) vol'
          type*, ' 9) stagnation enthalpy'
          type*, ' 10) CONTOUR INCREMENT'
          type 22
22      format(1,' selection = ')
          accept 222, ctype
222     format(I)

c* exit from contour loop
          if (ctype.eq.0) goto 999

          if (ctype.eq.10) then
            type 21
            format(1,' CONTOUR INCREMENT = ')
            accept 221, Cinc
221     format(F)
            type*, ' '
            goto 2
          endif

c* set up nodal contour values
          if (ctype.eq.5) then
            ptinf = 1./gam

```



```

endif

zmin = 1.e20
zmax = -1.e20

do NN = 1, Nmax
  rr = U(1,NN)
  uu = U(2,NN)/rr
  vv = U(3,NN)/rr

  if (ctype.eq.1) then
    zz = rr
  else if (ctype.eq.2) then
    pp = gam1*(U(4,NN) - 0.5*rr*(uu**2 + vv**2))
    mm = sqrt(rr*(uu**2 + vv**2)/(gam*pp))
    zz = mm
  else if (ctype.eq.3) then
    zz = sqrt(uu**2 + vv**2)
  else if (ctype.eq.4) then
    pp = gam1*(U(4,NN) - 0.5*rr*(uu**2 + vv**2))
    zz = pp
  else if (ctype.eq.5) then
    pp = gam1*(U(4,NN) - 0.5*rr*(uu**2 + vv**2))
    mm2 = rr*(uu**2 + vv**2)/(gam*pp)
    pt = pp*(1.+5*gam1*mm2)**(gam/gam1)
    zz = (1. - pt/ptinf)*100.
  else if (ctype.eq.6) then
    pp = gam1*(U(4,NN) - 0.5*rr*(uu**2 + vv**2))
    zz = (log(gam*pp) - gam*log(rr))*100.
  else if (ctype.eq.7) then
    zz = (1.+x(NN))*10.
  else if (ctype.eq.9) then
    pp = gam1*(U(4,NN) - 0.5*rr*(uu**2 + vv**2))
    zz = ((gam/gam1)*(pp/rr) + 0.5*(uu**2 + vv**2))*100.
  endif

  zmax = max(zmax,zz)
  zmin = min(zmin,zz)
  z(NN) = zz
enddo

if (ctype.eq.8) then
  open(unit=50, status='unknown', form='unformatted')
  read(50) Cmax, (vol(CC),CC=1,Cmax)
  close(unit=50)
  zmin = 1.e20
  zmax = -1.e20
  do NN = 1, Nmax
    z(NN) = 0.
  enddo
  do CC = 1, Cmax
    N1 = cell(CC,5)
    N2 = cell(CC,6)
    N3 = cell(CC,7)
    N4 = cell(CC,8)
    zz = vol(CC)
    zmax = max(zmax,zz)
    zmin = min(zmin,zz)
    z(N1) = z(N1) + 0.25*zz
    z(N2) = z(N2) + 0.25*zz
    z(N3) = z(N3) + 0.25*zz
    z(N4) = z(N4) + 0.25*zz
  enddo
endif

c* set title and extra variables needed for GRAFIC
if (ctype.eq.1) then
  TITLE = ' X Y DENSITY '
  NTITL = 24
  NCONT = 20
else if (ctype.eq.2) then
  TITLE = ' X Y MACH NUMBER '

```

```

        NTITL = 28
        NCONT = 20
    else if (ctype.eq.3) then
        TITLE = '      X      Y      NORMAL VELOCITY '
        NTITL = 32
        NCONT = 20
    else if (ctype.eq.4) then
        TITLE = '      X      Y      PRESSURE '
        NTITL = 25
        NCONT = 20
    else if (ctype.eq.5) then
        TITLE = '      X      Y      % TOTAL PRESSURE LOSS '
        NTITL = 38
        NCONT = 20
    else if (ctype.eq.6) then
        TITLE = '      X      Y      % ENTROPY '
        NTITL = 26
        NCONT = 20
    else if (ctype.eq.7) then
        TITLE = '      X      Y      LINEAR '
        NTITL = 23
        NCONT = 20
    else if (ctype.eq.8) then
        TITLE = '      X      Y      VOL '
        NTITL = 20
        NCONT = 20
    else if (ctype.eq.9) then
        TITLE = '      X      Y      % STAGNATION ENTHALPY '
        NTITL = 38
        NCONT = 20
    endif

c* find contour levels
    if (Cinc.ne.0.) then
        NCONT = int((zmax-zmin)/Cinc + 2.)
        CSTEP = Cinc
        CBASE = (real(int((zmin/Cinc)-1.)))*Cinc
    elseif (ctype.eq.2) then
        boundary = 0.
        do while (1)
            boundary = boundary + 0.05
            if (boundary.gt.zmin) then
                zmin = boundary - 0.05
                goto 200
            endif
        enddo
200        boundary = 0.
        do while (1)
            boundary = boundary + 0.05
            if (boundary.gt.zmax) then
                zmax = boundary
                goto 210
            endif
        enddo
210        CBASE = zmin
        NCONT = int((zmax-zmin)/0.05)
        CSTEP = 0.05
    else
        call GR_SCALE(zmin, zmax, NCONT-1, CBASE, CSTEP)
    endif

    z(Nmax+1) = NCONT
    z(Nmax+2) = CBASE + 0.01*CSTEP
    z(Nmax+3) = CSTEP

c* finish up title
    TITLE(NTITL+1:NTITL+24) = 'CONTOURS WITH INCREMENT '
    write(NUM,10) CSTEP
10    format(F6.3)
    TITLE(NTITL+25:80) = NUM

```

```

c* plot the contour lines
    indgr = 23
    call gr_control(g2pltc, indgr, TITLE, z, Nmax, x, y,
&                a5, a6, a7, a8, a9, a10)

    enddo

c* plot surface distribution
    else if (ptype.eq.3) then
        do while (i)

            type*, 'Type of surface distribution'
            type*, ' 0) TOP LEVEL'
            type*, ' 1) density'
            type*, ' 2) Mach number'
            type*, ' 3) normal velocity'
            type*, ' 4) surface flow angle'
            type*, ' 5) analytical surface flow angle'
            type*, ' 6) pressure'
            type*, ' 7) entropy'
            type*, ' 8) stagnation enthalpy'
            type*, ' 9) total pressure loss'
            type 33
33          format(9, ' selection = ')
            accept 333, stype
333         format(I)

c* exit from contour loop
            if (stype.eq.0) goto 999

c* set up array with points on line for upper surface
            if (stype.eq.9) then
                ptinf = 1./gam
            endif

            points = 0
            do EN = 1, tnode
                points = points + 1
                NN = enode(EN)
                rr = U(1,NN)
                uu = U(2,NN)/rr
                vv = U(3,NN)/rr

                if (stype.eq.1) then
                    zz = rr
                else if (stype.eq.2) then
                    pp = gam1*(U(4,NN) - 0.5*rr*(uu**2 + vv**2))
                    mm = sqrt(rr*(uu**2 + vv**2)/(gam*pp))
                    zz = mm
                else if (stype.eq.3) then
                    zz = sqrt(uu**2 + vv**2)
                else if (stype.eq.4) then
                    zz = atan(vv/uu)
                else if (stype.eq.5) then
                    zz = atan(senode(EN)/cenode(EN))
                else if (stype.eq.6) then
                    pp = gam1*(U(4,NN) - 0.5*rr*(uu**2 + vv**2))
                    zz = pp
                else if (stype.eq.7) then
                    pp = gam1*(U(4,NN) - 0.5*rr*(uu**2 + vv**2))
                    zz = (log(gam*pp) - gam*log(rr))*100.
                else if (stype.eq.8) then
                    pp = gam1*(U(4,NN) - 0.5*rr*(uu**2 + vv**2))
                    zz = ((gam/gam1)*(pp/rr) + 0.5*(uu**2 + vv**2))*100.
                else if (stype.eq.9) then
                    pp = gam1*(U(4,NN) - 0.5*rr*(uu**2 + vv**2))
                    mm2 = rr*(uu**2 + vv**2)/(gam*pp)
                    pt = pp*(1. + 5*gam1*mm2)**(gam/gam1)
                    zz = (1. - pt/ptinf)*100.
                endif

                xline(points) = x(NN)
            enddo
        enddo
    enddo

```

```

        yline(points) = zz
    enddo

    npts(1) = points

c* set up array with points on line for lower surface
    points = 0
    do EN = tnode+1, bnode
        points = points + 1
        NN = enode(EN)
        rr = U(1,NN)
        uu = U(2,NN)/rr
        vv = U(3,NN)/rr

        if (stype.eq.1) then
            zz = rr
        else if (stype.eq.2) then
            pp = gam1*(U(4,NN) - 0.5*rr*(uu**2 + vv**2))
            mm = sqrt(rr*(uu**2 + vv**2)/(gam*pp))
            zz = mm
        else if (stype.eq.3) then
            zz = sqrt(uu**2 + vv**2)
        else if (stype.eq.4) then
            zz = atan(vv/uu)
        else if (stype.eq.5) then
            zz = atan(enode(EN)/cenode(EN))
        else if (stype.eq.6) then
            pp = gam1*(U(4,NN) - 0.5*rr*(uu**2 + vv**2))
            zz = pp
        else if (stype.eq.7) then
            pp = gam1*(U(4,NN) - 0.5*rr*(uu**2 + vv**2))
            zz = (log(gam*pp) - gam*log(rr))*100.
        else if (stype.eq.8) then
            pp = gam1*(U(4,NN) - 0.5*rr*(uu**2 + vv**2))
            zz = ((gam/gam1)*(pp/rr) + 0.5*(uu**2 + vv**2))*100.
        else if (stype.eq.9) then
            pp = gam1*(U(4,NN) - 0.5*rr*(uu**2 + vv**2))
            mm2 = rr*(uu**2 + vv**2)/(gam*pp)
            pt = pp*(1.+5*gam1*mm2)**(gam/gam1)
            zz = (1. - pt/ptinf)*100.
        endif

        xline(npts(1)+points) = x(NN)
        yline(npts(1)+points) = zz
    enddo

    npts(2) = points

c** set plot indicators
    NLINE = 2
    IOPT(1) = 14
    IOPT(2) = 14

c** set title for plot
    if (stype.eq.1) then
        TITLE = ' X DENSITY DENSITY'
    else if (stype.eq.2) then
        TITLE = ' X MACH NO.MACH NUMBER'
    else if (stype.eq.3) then
        TITLE = ' X VELOCITYNORMAL VELOCITY'
    else if (stype.eq.4) then
        TITLE = ' X ANGLE SURFACE FLOW ANGLE'
    else if (stype.eq.5) then
        TITLE = ' X ANGLE ANALYTICAL SURFACE FLOW ANGLE'
    else if (stype.eq.6) then
        TITLE = ' X PRESSUREPRESSURE'
    else if (stype.eq.7) then
        TITLE = ' X ENTROPY-% ENTROPY'
    else if (stype.eq.8) then
        TITLE = ' X -STAGNATION ENTHALPY-% STAGNATION ENTHALPY'
    else if (stype.eq.9) then

```

```

      TITLE =
      *      ' X TOTAL PRESSURE LOSS' TOTAL PRESSURE LOSS'
      endif

c* plot the line
      indgr = 21
      call gr_line(IOPT,NLINE,TITLE,INDGR,xline,yline,npts)

      enddo
      else if (ptype.eq.4) then
c* calculate mean of total pressure loss
      ptinf = 1./gam
      zz = 0.
      zmax = 0.
      do NN = 1, Nmax
        rr = U(1,NN)
        uu = U(2,NN)/rr
        vv = U(3,NN)/rr
        pp = gam1*(U(4,NN) - 0.5*rr*(uu**2 + vv**2))
        mm2 = rr*(uu**2 + vv**2)/(gam*pp)
        pt = pp*(1.+0.5*gam1*mm2)**(gam/gam1)
        zz = zz + (1. - pt/ptinf)**2
        zmax = max(zmax,abs(1.-pt/ptinf))
      enddo
      zz = sqrt(zz/real(Nmax))

      type*, ' '
      type*, 'inlet Mach number      = ', Minl
      type*, 'rms total pres. loss   = ', zz, ' (',log10(zz),')'
      type*, 'max total pres. loss   = ', zmax, ' (',log10(zmax),')'
      type*, 'cells                  = ', Cmax
      type*, 'faces                  = ', Fmax
      type*, 'nodes                  = ', Nmax
      type*, 'inlet nodes            = ', Imax
      type*, 'outlet nodes           = ', Omax
      type*, 'periodic nodes         = ', Pmax
      type*, 'edge nodes             = ', Emax
      type*, 'face colors            = ', Fcolormax
      type*, 'cell colors            = ', Ccolormax
      type*, 'edge colors            = ', E2colormax
      type*, ' '
999      endif
      enddo

      end

      subroutine g2pltg(ifun, indgr, TITLE, alimits, info_string,
      *      a1, a2, a3, a4, a5, a6, a7, a8, a9, a10)
      implicit none

      include 'QUAD.INC'

      integer FF, EF, NH
      integer indgr
      character*80 TITLE
      integer ifun
      real alimits(4)
      integer a1, a2, a3, a4, a5, a6, a7, a8, a9, a10
      character*80 info_string
      real x1, x2, y1, y2

      if (ifun.eq.0) then
        return
      elseif (ifun.eq.1) then
        call gr_get_limits(x, y, Nmax, alimits)
        if (pitch.ne.0.) then
          alimits(4) = alimits(4) + pitch
        endif
      elseif (ifun.eq.2) then
        info_string = ' '
      end

```

```

elseif (ifun.eq.3) then
  do FF = 1, Fmax
    NN = face(FF,3)
    x1 = x(NN)
    y1 = y(NN)
    NN = face(FF,4)
    x2 = x(NN)
    y2 = y(NN)
    call gr_move(x1, y1, 0)
    call gr_draw(x2, y2, 0)
    if (pitch.ne.0.) then
      y1 = y1 + pitch
      y2 = y2 + pitch
      call gr_move(x1, y1, 0)
      call gr_draw(x2, y2, 0)
    endif
  enddo
endif
return
end

```

```

subroutine g2pltc(ifun, indgr, TITLE, alimits, info_string,
&               z, a2, a3, a4, a5, a6, a7, a8, a9, a10)
  implicit none

  include 'QUAD.INC'

  integer FF, EF, NN, CC, NC      !pointers
  integer node                    !pointer
  integer indgr
  character*80 TITLE
  integer ifun
  real alimits(4)
  real z(Maxnodes)
  integer a2, a3, a4, a5, a6, a7, a8, a9, a10
  character*80 info_string
  real x1, x2, y1, y2, y3, y4
  integer ncont                   !determine contour levels
  real cbase, cstep
  integer KK                      !pointer
  integer N1, N2, N3, N4         !nodes at end of edge face
  real xn(4), yn(4), zn(4)      !x, y, and contour value at nodes
  real xpoint, ypoint           !x, y of pointer
  real value                     !value of contour at pointer
  real cn(2,4)                  !x, y at cell nodes
  integer IIN                    !is pointer in cell?
  real zcont, cont(30)           !contour levels
  real xmax, xmin                !max & min contour values in cell

  if (ifun.eq.0) then
    return
  elseif (ifun.eq.1) then
    call gr_get_limits(x, y, Nmax, alimits)
    if (pitch.ne.0.) then
      alimits(4) = alimits(4) + pitch
    endif
  elseif (ifun.eq.2) then
    c* find contour value at point
    xpoint = alimits(1)
    ypoint = alimits(2)
    c* look for cell (x,y) is in
    do KK = 1, Cmax

```

```

      N1 = cell(KK,6)
      N2 = cell(KK,6)
      N3 = cell(KK,7)
      N4 = cell(KK,8)
      cn(1,1) = x(N1)
      cn(1,2) = x(N2)
      cn(1,3) = x(N3)
      cn(1,4) = x(N4)
      cn(2,1) = y(N1)
      cn(2,2) = y(N2)
      cn(2,3) = y(N3)
      cn(2,4) = y(N4)
      call gr_inside(IIN,cn,4,xpoint,ypoint)
      if (IIN.eq.1) then
c* linear interpolation for value
        zn(1) = z(N1)
        zn(2) = z(N2)
        zn(3) = z(N3)
        zn(4) = z(N4)
        call gr_contour_value(cn,zn,xpoint,ypoint,value)
        goto 30
      endif
      cn(2,1) = cn(2,1) + pitch
      cn(2,2) = cn(2,2) + pitch
      cn(2,3) = cn(2,3) + pitch
      cn(2,4) = cn(2,4) + pitch
      call gr_inside(IIN,cn,4,xpoint,ypoint)
      if (IIN.eq.1) then
c* linear interpolation for value
        zn(1) = z(N1)
        zn(2) = z(N2)
        zn(3) = z(N3)
        zn(4) = z(N4)
        call gr_contour_value(cn,zn,xpoint,ypoint,value)
        goto 30
      endif
    enddo
30    write (INFO_STRING,31) value
31    format(' function value = ',f15.6)

    elseif (ifun.eq.3) then

c* plot boundary
      do EF = 1, Emax
        FF = sface(EF)
        N1 = face(FF,3)
        N2 = face(FF,4)
        call gr_move(x(N1), y(N1), 0)
        call gr_draw(x(N2), y(N2), 0)
      enddo

      if (pitch.ne.0.) then
        do EF = 1, Emax
          FF = sface(EF)
          N1 = face(FF,3)
          N2 = face(FF,4)
          y1 = y(N1) + pitch
          y2 = y(N2) + pitch
          call gr_move(x(N1), y1, 0)
          call gr_draw(x(N2), y2, 0)
        enddo
      endif

c* set up contour levels
      ncont = z(Nmax+1)
      cbase = z(Nmax+2)
      cstep = z(Nmax+3)
      do NC = 1, ncont
        cont(NC) = cbase + (NC-1)*cstep
      enddo

c* loop through cells

```

```

do CC = 1, Cmax
  do NN = 1, 4
    node = cell(CC, NN+4)
    xn(NN) = x(node)
    yn(NN) = y(node)
    zn(NN) = z(node)
  enddo

  zmax = max(zn(1), zn(2), zn(3), zn(4))
  zmin = min(zn(1), zn(2), zn(3), zn(4))

c* find contour crossing on triangle for each contour level
do NC = 1, ncont
  zcont = cont(NC)
  if (zcont.lt.zmin .or. zcont.gt.zmax) then
c*   no need to compute
  else
    call gr_cbox(xn(1), yn(1), zn(1),
      &             xn(2), yn(2), zn(2),
      &             xn(3), yn(3), zn(3),
      &             xn(4), yn(4), zn(4), zcont)
    if (pitch.ne.0.) then
      y1 = yn(1) + pitch
      y2 = yn(2) + pitch
      y3 = yn(3) + pitch
      y4 = yn(4) + pitch
      call gr_cbox(xn(1), y1, zn(1),
        &             xn(2), y2, zn(2),
        &             xn(3), y3, zn(3),
        &             xn(4), y4, zn(4), zcont)
    endif
  endif
enddo
enddo
endif
return
end

```