NASA Grant NAG 2099

Large Eddy Simulation of Flow in Turbine Cascades

Final Progress Report

P.G. Huang
Department of Mechanical Engineering
University of Kentucky
Lexington, Kentucky 40506-0503

Submitted to:
Dr. David Ashpis, Technical Monitor
NASA Glenn Research Center
Cleveland, OH 44135
August 31, 2004

Large Eddy Simulation of Flow in Turbine Cascades

Using LESTool and UNCLE Codes

P.G. Huang

Department of Mechanical Engineering
University of Kentucky
Lexington, Kentucky 40506-0503

Progress Summary

During the period December 23, 1997 and December August 31, 2004, we accomplished the development of 2 CFD codes for DNS/LES/RANS simulation of turbine cascade flows, namely LESTool and UNCLE. LESTool is a structured code making use of 5th order upwind differencing scheme and UNCLE is a second-order-accuracy unstructured code. LESTool has both Dynamic SGS and Sparlart’s DES models and UNCLE makes use of URANS and DES models. The current report provides a description of methodologies used in the codes.
1. Introduction

Flow transition plays an important role in turbomachinery applications. The majority of boundary layer flows in turbomachines involve flow transition under the effects of freestream turbulence, diverse pressure gradients, wide range of Reynolds numbers, flow separation, and unsteady wake-boundary layer interactions.

Prediction of this type of complex flows is an important element in analysis and performance evaluation of gas turbine engine components and ultimately in the design of more efficient jet engines. Especially, in low pressure turbine applications prediction of transition becomes pivotal in terms of efficiency. For low pressure turbines the flow is mostly turbulent at the high Reynolds number conditions encountered at take off and the efficiency is at its design maximum. However, at high altitudes and cruise speeds which correspond to lower Reynolds number conditions, unpredicted losses and substantial drops in efficiency have been observed. These losses are attributed to flow separation on the suction surface of the turbine blades. At low Reynolds numbers, the boundary layers on the airfoil surface have a tendency to remain laminar and hence the flow may separate before it becomes turbulent, causing increase in fuel consumption and drop in efficiency. The impact of such losses is directly felt on the operation costs. It has been estimated that a 1% improvement in the efficiency of a low pressure turbine would result in a saving of $52,000 per year on a typical airliner.

In order to calculate the losses and heat transfer on various components of gas turbine engines, and to be able to improve component efficiencies and reduce losses through better designs, accurate prediction of transitional boundary layers is essential. When one deals with a complex fluid phenomena like a transition, separation and turbulence, several hundred millions grid points are needed to resolve boundary layers and other flow structures correctly. We have started to develop technology to make such large scale simulations not only possible at supercomputing centers like NCSA or NAS but on inexpensive, high-performance clusters of PCs, or "Beowulfs". These clusters are specialized for CFD applications, using the novel approach that the hardware, operating system, and application code are optimized together rather than separately. A Honorable Mention in the Price/Performance Category of the Gordon Bell Prize was awarded for this approach at IEEE/ACM SC2000 Conference on High-Performance Networking and Computing.
Several turbulence test cases have been computed and an overview of the results is given.

2. Code Descriptions

(1) A description of LESTool is give in Appendix I.
(2) A description of UNCLE is given in Appendix II.

3. Publications by the PI

4. Award received by the PI

A Honorable Mention in the Price/Performance Category of the Gordon Bell Prize was awarded for this approach at IEEE/ACM SC2000 Conference on High-Performance Networking and Computing.
Appendix I - LESTool
LES Turbulence Models

1. LES - Filtered Equations

Large-eddy simulation (LES) is based on the definition of a filtering operation: a filtered (or large-scale, or resolved) quantity, denoted by an over bar, is defined as

\[ \bar{f}(x) = \int f(x') G(x, x') dx', \]

where \( D \) is the entire domain and \( G \) is the filter function, which determines the size and structure of the small scales. For compressible flow the large-scale equations are operationally simpler if Favre filtered quantities are used. A Favre filtered variable is defined as \( \bar{f} = \rho f / \rho \). Applying the spatial filter \( G \) to the governing equations leads to

\[ \frac{\partial \bar{\rho}}{\partial t} + \frac{\partial (\bar{\rho} \bar{\mu})}{\partial x_j} = 0 \]

\[ \frac{\partial \bar{\rho} \bar{\mu}}{\partial t} + \frac{\partial (\bar{\rho} \bar{\mu} \bar{u}_j)}{\partial x_j} = -\frac{\partial \bar{\rho}}{\partial x_j} + \frac{\partial \bar{\tau}_{ji}}{\partial x_j} - \bar{\tau}_{ji} + \frac{\partial}{\partial x_j} (\bar{\tau}_{ji} - \bar{\tau}_{ji}) \]

\[ \frac{\partial \bar{\rho} \bar{E}}{\partial t} + \frac{\partial \bar{\rho} \bar{E} \bar{u}_j}{\partial x_j} = -\frac{\partial \bar{\rho}}{\partial x_j} + \frac{\partial \bar{\tau}_{ji}}{\partial x_j} + \bar{\tau}_{ji} \bar{u}_j \]

\[ -\bar{c}_p \frac{\partial q_{SGS}}{\partial x_j} + \frac{1}{2} \frac{\partial}{\partial x_j} (\bar{\tau}_{ji}^S \bar{u}_j) - \frac{\partial}{\partial x_j} D_j \]

\[ -\frac{\partial}{\partial x_j} (\bar{\tau}_{ji} - \bar{\tau}_{ji}) + \frac{\partial}{\partial x_j} (\bar{\tau}_{ji} \bar{u}_j - \bar{\tau}_{ji} \bar{u}_j) \]

where a perfect-gas equation of state is assumed. The filtered stress tensor, \( \bar{\tau}_{ji} \) and \( \bar{\tau}_{ji} \) are given by

\[ \bar{\tau}_{ji} = 2\mu S_{ji} - \frac{2}{3} \delta_{ji} \mu S_{kk} \]

\[ \bar{\tau}_{ji} = 2\tilde{\mu} \bar{S}_{ji} - \frac{2}{3} \delta_{ji} \tilde{\mu} \bar{S}_{kk} \]

Here \( S_{ji} \) is the strain-rate tensor defined by

\[ S_{ji} = \frac{1}{2} \left( \frac{\partial u_i}{\partial x_j} + \frac{\partial u_j}{\partial x_i} \right) \]
The filtered heat fluxes, \( \vec{q}_j = -k \partial \vec{T} / \partial \vec{x}_j \) and \( \vec{q}_j = -\vec{k} \partial \vec{T} / \partial \vec{x}_j \).

All boxed terms must be modeled. The following list defines modeling assumptions for all six terms which have to be modeled.

1. This first boxed term is the divergence of the subgrid-scale (SGS) stresses, \( \tau_{y}^{SGS} = \overline{\rho(\vec{u}_j \vec{u}_j - \overline{\vec{u}_j \vec{u}_j})} \), and has to be modeled. Note that the trace of the SGS stresses in compressible flows cannot be included in the modified pressure and requires separate modeling. The modeling will be described in the next section.

2. This 2\(^{nd}\) boxed term appears because of the nonlinearity of the viscous stresses and is invariably neglected.

3. The SGS heat flux model for \( q_{j}^{SGS} = \overline{\rho(\vec{u}_j \vec{T} - \overline{\vec{u}_j \vec{T}})} \) will be described in the next section.

4. This term, \( D_j = \frac{1}{2}(u_j u_k u_k - \overline{u}_j u_k u_k) \), is similar to the turbulence diffusion that appears in the subgrid-scale kinetic energy equation.

5. This term, \( \partial(\vec{q}_j - \overline{\vec{q}_j}) / \partial \vec{x}_j \), is neglected because of the same reasons as 2.

6. This term, \( \partial(\tau_{\mu i} \vec{u}_i - \overline{\tau_{\mu i} \vec{u}_i}) / \partial \vec{x}_j \), is the viscous SGS work and is neglected.

To summarize the following quantities will be modeled in our implementation:

1. SGS stresses, \( \tau_{y}^{SGS} \).

2. SGS heat flux, \( q_{j}^{SGS} \).

1.2 Dynamic SGS model

For the left-hand side a trace-free Smagorinsky eddy viscosity model is used for \( \tau_{y}^{SGS} \). However the eddy viscosity coefficient will be a function of the instantaneous flow variables. Let

\[
\tau_{y}^{SGS} = -\frac{1}{3} q^2 \delta_y = -2C \overline{\rho} \Delta^2 |\vec{S}| \vec{S}_y^* \]

where \( \vec{S}_y^* \) is the trace-free rate of the strain tensor

\[
\vec{S}_y^* = \vec{S}_y - \frac{\delta_y}{3} \vec{S}_{kk}
\]

and \( |\vec{S}| \) is the norm of \( \vec{S}_y \).
The isotropic part of the SGS Reynolds stress tensor \( q^2 = \tau_{kk} \) has to be modeled separately. It is parameterized by the expression:

\[
q^2 = 2C_i \rho \Delta^2 |\vec{\mathcal{S}}|^2
\]

To compute \( C_i \), the following equation is used:

\[
\rho \vec{u}_i \vec{u}_j \frac{\partial}{\partial \rho} \frac{\partial}{\partial \rho} - 2C_i \rho \Delta^2 \left| \frac{\partial}{\partial \rho} \vec{S} \right|^2 = 2C_i \left( \rho \Delta^2 \left| \vec{S} \right|^2 - \rho \left| \frac{\partial}{\partial \rho} \vec{S} \right|^2 \right)
\]

Germano et al. [5] showed that expressions similar to the ones multiplying \( C_i \) can become zero. Therefore an averaging procedure is needed to make the determination of the SGS coefficients well conditioned. It is assumed that \( C_i \) is independent of the directions in which the flow is homogeneous. Volume averaging leads to

\[
C_i \Delta^2 = \frac{\left\langle L_{kk} \right\rangle}{2 \left( \rho \Delta^2 \right) \left| \frac{\partial}{\partial \rho} \vec{S} \right|^2}
\]

To obtain \( C_i \), we use the following model

\[
T_j - \frac{1}{3} T_{kk} \delta_j = -2C_i \rho \Delta^2 \left| \vec{S} \right|^2
\]

This leads to

\[
L_j = \frac{1}{3} L_{\delta j} \delta_j - 2C_i \rho \Delta^2 \left| \vec{S} \right|^2 + 2C_i \rho \Delta^2 \left| \vec{S} \right|^2 + \frac{1}{3} L_{\delta j} \delta_j + 2C\Delta^2 M_j
\]

with \( M_j \)

\[
M_j = \rho \left| \vec{S} \right|^2 - \rho \left( \frac{\Delta}{\Delta} \right)^2 \left| \vec{S} \right|^2
\]

Contracting of \( M_j \) which was recommended by Lilly is outlined below. Since equation for \( L_j \) represents six independent equations in one unknown, its error can be minimized by applying a least squares approach. Define \( G \) as

\[
G = \left( L_j - \frac{1}{3} \delta_j L_{kk} - 2CM_j \right)^2
\]

By setting \( \partial G / \partial C = 0 \), \( C \) is evaluated as
\[ C \Delta^2 = \frac{1}{2} \left( L_{ij} M_{ij} - \frac{1}{3} L_{kk} M_{kk} \right) \]

where \( \cdot \) denotes a spatial averaging operation.

### 1.3 DES model

The DES turbulence model is based on the blending of RANS and LES to provide a model that can accurately predict unsteady flows without requiring high-precision grids near the walls. A RANS turbulence model is used to simulate the turbulence within the boundary layer, gradually subsuming into the LES model as one moves away from the wall. Further details on this technique can be found in [15], [14] and others. The RANS turbulence model used in LESTool is the one-equation Spalart-Allmaras model,

\[
\begin{align*}
\frac{D \tilde{\nu}}{Dt} &= P_k - L + \frac{1}{\sigma} \left\{ \nabla \cdot \left[ (\nu + \tilde{\nu}) \nabla \tilde{\nu} \right] \right\}, \\
P_k &= c_{b1} \left| \frac{f_{o3} \Omega}{\sigma} + \frac{\tilde{\nu}}{\kappa^2 d^2} \right| \tilde{\nu} + \frac{1}{\sigma} c_{b2} (\nabla \tilde{\nu} \cdot \nabla \tilde{\nu}), \\
L &= c_{w1} f_w \left( \frac{\tilde{\nu}}{d} \right)^2, f_{o1} = \frac{\chi^3}{\chi^3 + c_{o1}}, f_{o2} = \left( 1 + \frac{\chi}{c_{v2}} \right)^{-3}, \\
f_{o3} &= \frac{1}{\chi} (1 - f_{o2}) (1 + f_{o1} \chi), \chi = \frac{\tilde{\nu}}{\nu}, \\
f_w &= g \left( \frac{1 + c_{w3}^6}{g^6 + c_{w3}^6} \right), g = r + c_{w2} (\nu^6 - r), r = \frac{\tilde{\nu}}{\tilde{\kappa}^2 d^2}, \\
\tilde{S} &= \left| f_{o3} \Omega + f_{o2} \frac{\tilde{\nu}}{\kappa^2 d^2} \right|
\end{align*}
\]

with the coefficients given below

\[
\begin{align*}
c_{b1} &= 0.1355, c_{b2} = 0.622, c_{o1} = 7.1, c_{o2} = 5.0, c_{w1} = c_{b1} / \kappa^2 + (1 + c_{b2}) / \sigma, \\
c_{w2} &= 0.3, c_{w3} = 2, \sigma = 2 / 3, \kappa = 0.41, \Omega = |\nabla \times \tilde{u}|,
\end{align*}
\]

and \( d \) is the distance to the nearest wall. Note that this formulation does not include the trip functions. Large values of \( r \) can be truncated at 10 or so. The turbulent viscosity for the momentum equations is found from \( \nu_t = f_{o1} \tilde{\nu} \). For the DES approach the length scale \( d \) is changed into

\[ d' = \min(d, \max(\Delta x, \Delta y, \Delta z)) C_i \]

with \( C_i = 0.65 \). This gives the turbulence model behavior near the wall and LES type behavior away from the wall.

### 2. Numerical method

The compressible Navier-Stokes equations are discretized using an iterative diagonal dominant ADI (DD-ADI) algorithm that can be written in the following form:
The right-hand side is approximated by the newly develop ENO-Pade method [17] for the convective flux and sixth-order central for the diffusive terms. The basic algorithm is the following. First the flux at each cell face is computed as shown below:

\[ F_{i+1/2} = F(U_{i+1/2}^H) - \frac{1}{2} A_{i+1/2}^R (U_{i+1/2}^R - U_{i+1/2}^L) \]

Quantities with the superscript \( H \) are computed using a 8th-order central interpolation method. For the quantities with the superscript \( R \) or \( L \) the standard ENO-interpolation method is used. Then the flux is differentiated using a cell centered Pade formula [10]:

\[
\alpha \phi_{i-1} + \phi_i + \alpha \phi_{i+1} = a \phi_i^{1/2} - \phi_{i-1/2} + b \phi_i^{3/2} - \phi_{i-3/2}
\]

The left-hand side is approximated by a lower-order method. The high order solution can be achieved by performing inner iterations since the order of accuracy is not affected by low order treatment of the left-hand side.

The proposed algorithm has the following form:

\[
(D^L + \Delta t L_x) d(\delta U^1) = D^L \left[ (\delta U^3)^{m-1} - (\delta U^1)^{m-1} \right]
\]

\[
(D^L + \Delta t L_y) d(\delta U^2) = D^L \left[ (\delta U^1)^{m} - (\delta U^2)^{m-1} \right]
\]

\[
(D^L + \Delta t L_z) d(\delta U^3) = D^L \left[ (\delta U^2)^{m} - (\delta U^3)^{m-1} \right]
\]

This algorithm is easily parallelizable because we loop through a three dimensional array plane by plane. The outline of the algorithm is the following:

1. for each \( k \): solve equations in \( i \) and \( j \) planes.
2. for each \( j \): solve equations in \( k \) and \( i \) planes
3. for each \( i \): solve equations \( j \) and \( k \) planes

This means after completion of the above algorithm we have already completed two iterations. From our experience three to four iterations are needed for each time-step to reach the required accuracy.

Fortran 90 was chosen as the programming language for this project because of the new powerful array syntax, the addition of derived types and the module concept. The code has been developed based on an object-oriented methodology to simplify the transition from the Cartesian implementation to an implementation for general coordinates and chimera-type overlapping grids. The new features of Fortran 90 like abstract data types or generic programming support an object-oriented programming style even though it is not designed to be an object-oriented language.
3. Parallelization

3.1 Shared Memory parallel approach

The LESTool code was parallelized in a straightforward manner: The code was compiled with the -mp option to enable parallelism in the compile phase. This was combined with placing OpenMP compiler directives, which instruct the compiler to generate code that will execute in parallel.

These directives were placed at key spots within the code for the greatest parallel efficiency. This resulted in a decomposing of the 3D problem into groups of 1D lines, with each group assigned to a dedicated processor.

The efficiency of the parallel decomposition was enhanced by the use of the first touch policy which is specific to the SGI Origin 2000. This means that memory allocated is physically placed on the processor which touches the memory location first. This means that all large three dimensional blocks of memory where initialized in planes of constant k to get a memory distribution which is favorable for our algorithmic design (see figure 1).

![Figure 1: Partitioning of the data along the k-axis](image)

Since we target shared memory computers there is no explicit data redistribution needed. Instead we split up the k-sweeps among the processors by partitioning work along the y-axis (fig. 2). This is much simpler and easier to implement compared to the distributed memory concept.

![Figure 2: Sweep along the zx-planes](image)
3.2 Distributed memory approach

In order to achieve better portability and performance on a wider range of computational platforms a distributed approach was also implemented. The parallel structure of the distributed version of LESTool is based on splitting the computational grid into sub-blocks, which are then distributed to each processor (figure 3). For a complex geometry, this is a nontrivial exercise where an uneven load-balance across different processors could significantly reduce the computational efficiency of the overall code. The partitioning of the grid into sub-blocks is performed independently of the grid generation, using virtual partitions to define the domains corresponding to each processor. The current algorithm governing the partition process is based on spectral recursive bisection in conjunction with a small database of the computational speeds associated with each node. The splitting is performed as a preprocessing task, generating a file that maps the grid sub-blocks onto the processors. This file is the only difference between performing a serial computation and a parallel computation on a distributed machine.

Figure 3: Communication pattern for the distributed approach
Communications between the grid sub-blocks occurs when the sub-blocks exchange data about the flow variables at the boundaries. As shown in figure 3, the flow variables on the edge of one grid block are communicated to the dummy points of the neighboring grid block, and vice versa. LESTool requires such a communication step after each update, or sub-iteration, of the flow variables. The low-level implementation of the communication between the sub-blocks uses a MPI-based communications system. The communication model is a mailbox algorithm where the data is sent in a non-blocking communication mode as early as possible.

4. Accuracy and Performance of LESTool

4.1 Accuracy of the ENO-Pade method

In a previous proposal a fifth-order upwind scheme has been proposed to simulate flow around a low pressure turbine. This method showed too much numerical dampening so that the transition and separation on the low-pressure turbine couldn't be simulated. In order to develop a code suitable for transition and turbulence a new numerical method - the ENOPade method - was developed [17]. This method combines the stability of the ENO scheme with the high-order accuracy of the Pade method. Several test cases have been computed and the damping of the new scheme is much lower as you can see below from one test case. This case consists of the unsteady advection of a vortex in a uniform flow. Here the capabilities of the different numerical schemes to accurately advect the vortex structures are tested which is critical to LES/DNS simulations.

![Figure 4: Contour lines of vorticity magnitude at three time intervals of ENO-Pade method.](image)

![Figure 5: Swirl velocities of the vortex along y=0 at T=12](image)
4.2 Performance Results

4.2.1 Single processor performance tuning

The key point on cache based architectures to achieve high floating point performance is cache-awareness of the algorithm. This stage of tuning is very important because efficient utilization of the cache architecture assures good overall performance. Single processor performance is the basic step to get good overall performance for the parallel computation.

During the development of the code special care was taken by placing variables which are used concurrently in the calculation close together in the main memory. One major detail is that the first index in the data arrays is used for addressing the different components of global variables, such as conservative variables. This is contrary to a vector architecture, where the index for a variable is normally addressed by the last index in an array. The data is copied into one-dimensional scratch arrays which fit into the second level cache. The main algorithm is performed on these scratch arrays which have a memory layout and access pattern (stride one) optimal for the cache architecture. This measure alone resulted in a floating-point performance of 60 MFLOPS.

To achieve further speedup of the code, the main bottlenecks identified were the tridiagonal and block-tridiagonal matrix solvers. To increase the utilization of memory bandwidth, all arithmetic operations involving the 5x5 block matrices, used in block tridiagonal and periodic block tridiagonal solvers, were unrolled. This resulted in a much longer and less desirable code, but the performance results outweigh this disadvantage. The LESTool code achieves a floating-point performance of about 120 MFLOP/s on a 180 MHz Origin 2000. We consider this a very satisfactory performance.

Single processor run times for a single test case sufficiently large so that it does not fit in the L2 cache of all our CPUs has been chosen. The runtime of the code / time step is reported in table 1. The LosLobos results (733 MHz Pentium III) are much worse than the Athlon results - this is probably related to (2 computations/2 data transfers per cycle). However, we are not absolutely confident that this is the optimal possible performance of LESTool on LosLobos, and as such subsequent results will emphasize the KLAT2 and KFC1 outcomes.

Table 1: Execution time in seconds on different Linux platforms for a single time step for the 64$^3$ test case

<table>
<thead>
<tr>
<th>Compiler</th>
<th>Double Precision</th>
<th>Cost/processor</th>
<th>$/TSpS</th>
</tr>
</thead>
<tbody>
<tr>
<td>700 MHz Athlon Gcc-3.2</td>
<td>37s</td>
<td>$625</td>
<td>3030</td>
</tr>
<tr>
<td>733 MHz Pentium III Egcs-2.9.1.66</td>
<td>123s</td>
<td>$2930</td>
<td>18170</td>
</tr>
<tr>
<td>1.5GHz Athlon MP Gcc-2.96</td>
<td>22s</td>
<td>$750</td>
<td>2240</td>
</tr>
</tbody>
</table>
4.2.2 OpenMP performance tuning

In figure 6, the same test case with the improved fine-grain parallelization is shown. The removal of the barriers gives a much better speedup and provides a better performance of the fine-grain parallelization. The program scales better for larger number of shared processors. However, there still seems to have a limit on the number of the processor used for the fine-grain parallelization. Since the memory is distributed in planes of constant k over the available processors using more than 32 processors is no longer advantageous. By choosing a much larger test case of $256^3$ grid points, as also shown in figure 6 by the dashed line, it can be seen that because of a larger computational load per processor the performance scales much better. For this case, the program performs well up to 64 processors.

![Graph](image)

Figure 6: Speedup with reduced synchronization and orphaned OpenMP directives

4.2.3 Parallel performance

In order to achieve a reasonable throughput the wall time for our computations with LESTool must be at most on the order of 10s / time step. A comparison of the wall times for the three different test cases follows

In figure 7 the wall times are compared for KLAT2 and KFC1. The performance of KFC1 is obviously better than that of KLAT2 because of the higher clock frequency. The clock frequency of KFC1 is double than that of KLAT2 but the speed of LESTool on KFC1 is obviously not twice compared to that on KLAT2. Here, the memory system plays an important role as well--KFC1 has a PC21000 memory system, whereas KLAT2 has a PC100 memory system. Our goal to reach a time of less than 10s / time step is already reached on two processors on KFC1, while for KLAT2 it takes about six processors to come into the same total performance range.
The $128^3$ case is much more demanding. In this case it takes about ten nodes or twenty processors on our dual processor machine KFC1 to obtain a wall clock time under 10s. For KLAT2 forty processors are needed to obtain the desired computational speed. Note that for this example KFC1 does achieve twice the speed of KLAT2.

The $196^3$ case severely stresses our cluster computers. Here we cannot reach the 10s mark on either of the clusters. For KFC1, we can achieve less than 20s / time step on 36 processors, or twice that achievable is 22.7s.
KFC1 has a three-way channel-bonded network. The code scales well for all three test cases as shown in figure 10. Note that we could run the single processor version for the $128^3$ but the performance was so low because of swapping that we decided to scale the larger cases to the minimum number of processors on which the case could reasonable run. Note that the sizeable drop in performance for 34 processors in all three cases. This is related to poor load balance. The only way we can divide our grid is in two slices in j-direction and seventeen slices in the k-direction. Given the cubical symmetry of this problem, the best subdivision is a equal number of cuts in the i, j, k direction. An example for an optimal partitioning is the case for 16 processors. Each sub-block consists of $16^3$ grid points. This example shows the importance of the partitioning on the parallel performance of LESTool.

![Figure 10: Speedup on KFC1 for the $64^3$, $128^3$ and $196^3$ cases](image1)

KLAT2 has a FNN network architecture which is optimized for next neighbor communication. LESTool scales even better on KLAT2 than it does on KFC1 for all three test cases as shown in figure 11. Note that a similar effect caused by uneven load splitting can be seen on this machine as well.

![Figure 11: Speedup on KLAT2 for the $64^3$, $128^3$ and $196^3$ cases](image2)
5. DNS of homogeneous turbulence

The Direct Numerical Simulations (DNS) of decaying isotropic turbulence were presented by Blaisdell et al. [2]. The Blaisdell's DNS initial spectrum of case ia64f was generated using Crecomp [4] are used as the input for LESTool. The velocity fields needed for LESTool are given by Crecomp in such a way that irrespective of the grid density, the energy spectrum produced has the same energy for all the initial spectra. The rms velocities are taken as the input decides the total energy under the curve. This energy is divided across the range of wavelengths available.

![Energy Spectrum Comparison](image)

Figure 12: (64^3 grid) Comparison of energy spectra for various values of C_{DES} with Blaisdell's spectra at t=7

In the above figure, the Blaisdell's DNS result is indicated by the solid black curve running all the way down. All the other curves with different symbols correspond to simulations run using LESTool for various values of grid sensitivity parameter C_{DES} considered. It can be seen that when a 64^3 grid is used, the No Turbulence Model matches best with the Blaisdell's result. When we look at the DES results, the curve with C_{DES}=0.01 almost merges with the No Model case and Blaisdell's curve. This somewhat confirms that the LESTool has generated a DNS solution as did in Blaisdell's simulations.

6. DNS of turbulent channel flow

The results presented in this paper were based on a direct numerical simulation of a fully developed channel flow reported in [7]. The flow has two periodic (x and z) directions and the solid wall condition was imposed in the y direction. The Reynolds number based on the wall shear velocity, \( u_r = \sqrt{\tau_w / \rho} \), is \( Re_r = \rho u_r H / \mu_w = 180 \), where \( H \) is half the channel width. The streamwise and spanwise dimensions of the channel are 4\( \pi H \) and 2\( \pi H \), respectively. The computation is carried out on a grid using 200x121x200 grid points in x, y, and z, respectively. The flow field is initialized with a laminar solution and random fluctuations are superimposed on the pressure field. The governing equations are then integrated in time until a statistical equilibrium is reached (\( tu_r / H > 30 \)).
Figure 13 shows the dimensionless velocity, $u^* = u/u_r$, plotted against dimensionless wall distance, $y^+ = yu_r/\nu$. Also shown in this figure is the comparison of the predicted mean velocity profile against the data cited in [8] for the same Reynolds number. The dotted line represents the desirable profile in the viscous sublayer, $u^* = y^+$, and the dashed line denotes the log-law line, $u^* = \ln y^+ / 0.41 + 5.2$. The figure shows that the current mean velocity profile matches the data of Kim et al. [8] very well. The comparison of the rms values as shown in figure 14 of the velocity shows good agreement with the data of Kim et al.

![Mean velocity profiles](image1)

Figure 13: Mean velocity profiles, solid line, LESTool; circled symbols, DNS of Kim et al.; thin dashed line, viscous sublayer; tick dashed line, log layer.

![Comparison of RMS values](image2)

Figure 14: Comparison of RMS values
7. Simulation of a flow over a cylinder

A more complex basic flow is flow over a cylinder. This is an informative test case for investigating DES, as many of the flow characteristics that have been measured in experiments are not captured by traditional RANS techniques. A critical consideration when working with DES is proving to be grid construction - fundamentally, the local grid dimensions control the location of the transitional region where DES switches from RANS to LES mode. Strelets discusses the challenge of grid design in the conclusion of his 2001 paper [16]. However, it is not yet clear if there is in general a truly grid-independent solution with DES. Rather than asymptotically approaching a single, grid-independent result, sufficiently fine grid outputs might only be confined to a definable range of solutions. This presents an additional challenge to the validation process for Detached-Eddy Simulation and expands considerably the number of simulations necessary to concretely define the advantages and limitations of DES. In figure 15 lines of the vorticity are shown. This shows that the shear layer starts to become turbulent. Further calculations are needed to determine the influence of grid resolution on the performance of the DES mode.

![Figure 15: Vorticity contours of flow around cylinder](image)

8. PaK-B blade cascade simulations

In the current study, the cascade flow is simulated for the PakB blade. The grid resolution is 200x100x100, shown in Figure 16. The blade Reynolds number in the simulations is 25,000. This is typical Reynolds number for high altitude aircraft engines. This case is perfect for the application of Large-Eddy simulation (LES) or Detached Eddy simulation (DES) since the Reynolds number is relatively low, so the resolution requirements are not as demanding as for the design point. A large separation region is visible starting at about 60% of the axial cord and extending to the end of the blade. The flattened region or plateau,
between 60% and 90% of the axial cord, shows the region of boundary layer separation. The numerical results of this study are compared with the experimental case of Bons [3].

![3D view of the grid](image)

Figure 16: 3D view of the grid.

In the inflow plan (grey plane in figure 16) a random pressure field is prescribed to simulate the unsteady turbulent inflow. The blade, shown in red has wall boundary conditions and the green plane shows the outflow condition. The simulation is initialized with a homogeneous turbulence field as discussed in Chapter 6.

In figure 17, the averaged solution of the velocity field in the mid plane is shown and the comparison of the time-averaged pressure coefficient is shown in Figure 18. The comparison with the experimental data seems very good.

![Time-averaged solution of the velocity vector at the center plane](image)

Figure 17 the time-averaged solution of the velocity vector at the center plane.
Figure 18: $C_p$ comparison of calculations with design curve and experimental values.

References


Appendix II - UNCLE
A Center Pressure Based Method for Two/Three-Dimensional Unstructured Incompressible Navier-Stokes Solver

Abstract

A center pressure based method is presented in this paper, and which has been implemented into a new two/three-dimensional parallel unstructured CFD code, UNCLE, which is developed at the University of Kentucky to meet the challenges of physical problems with complex geometries and complicated boundary conditions while maintaining high computational efficiency. Good load balancing across computational nodes is achieved by using METIS. In order to demonstrate the accuracy and performance of center pressure based method, several test cases are presented for validation such as two-dimensional incompressible flow past a flat plate, two/three- dimensional driven cavity flow, and two/three- dimensional flow over a circular cylinder. Notably, an extensive qualitative and quantitative study of two-dimensional flow over a circular cylinder for low Reynolds number is also presented in this paper.

1 Introduction

Continual improvements in computer technologies and computational fluid dynamics (CFD) algorithms have established CFD codes as a reliable tool for fundamental research or industrial applications. To deal with increasing grid sizes and demands for faster output, parallel computation of CFD has become a standard approach. To deal with the different challenge presented by some physical problems with complex geometries and complicated boundary conditions is now approached through unstructured CFD grids due to their ability to smoothly conform to complicated boundaries. However, combining unstructured grids with a parallel code presents still other challenges, such as achieving well-balanced grid decomposition on a distributed system and efficient parallel performance. In order to meet these challenges, a center pressure based method has been implemented into a new parallel unstructured CFD code called UNCLE, which has been developed at the University of Kentucky. UNCLE is designed to meet the challenges of using unstructured grid codes on high-performance parallel computers. It is a two/three-dimensional finite volume unsteady incompressible Navier-Stokes solver with center pressure based SIMPLE algorithm with second order accuracy in both time and space. To increase flexibility in complex geometries, center pressure based method is extended to use a variety of grid types, such as triangular, quadrilateral, tetrahedral, and hexahedral meshes. To obtain good load balancing across computational nodes, METIS [1] is applied for domain decomposition. METIS is a set of programs for partitioning graphs and finite element meshes, and for producing fill-reducing orderings for sparse matrices. The algorithms implemented in METIS are based on multilevel graph partitioning schemes. The key features of METIS include extremely fast partition, high quality partitions, and low fill orderings. The parallel construction of UNCLE is based on message passing interface (MPI) protocols and has worked successfully on systems ranging from commodity PC clusters up to traditional supercomputers. In order to demonstrate the accuracy and performance of center pressure based method, several test cases are presented for validation such as two-dimensional incompressible flow past a flat plate,
two/three-dimensional driven cavity flow, and two/three-dimensional flow over a circular cylinder.

2 Numerical Methods

A center pressure based method for two/three-dimensional finite volume unstructured incompressible Navier-Stokes solver for steady/unsteady flow fields is presented in this paper. It is center pressure based SIMPLE algorithm with second order accuracy in both time and space. In order to compute numerical flux on interfaces, a second order upwind scheme is adopted to compute advection terms and second order central difference scheme is used for diffusion terms. Non-staggered grids with the Rhie and Chow momentum interpolation method [2] is employed to correct the checkerboard solution in the SIMPLE scheme.

2.1 Governing equations

The governing equations for unsteady incompressible viscous flow under the assumption of no body force and heat transfer are as below.

Conservation of Mass

\[ \frac{\partial}{\partial t} \int \rho dV = -\oint \rho u_n dS \]  

Conservation of Momentum

\[ \frac{\partial}{\partial t} \int \rho u_j dV = -\oint \rho u_n u_j dS - \oint p n_j dS + \oint \tau_{ij} n_j dS \]  

Conservation of Energy

\[ \frac{\partial}{\partial t} \int \rho E dV = -\oint \rho u_n E dS - \oint p u_j dS + \oint u_j \tau_{ij} n_j dS \]  

where \( \rho \) is density, \( p \) is pressure, \( u_j \) is component of velocity vector, \( n_i \) is unit normal vector of the interface, \( \tau_{ij} \) is tensor of shear force, and specific internal energy \( E = e + \frac{1}{2}(u^2 + v^2 + w^2) \). Notably, density is constant for incompressible flow.

2.2 Convective and diffusive fluxes

Figure 1(a) shows the schematic diagram for integration area for convective fluxes. By using Taylor series expansion, flow properties on the interface can be obtained by Eq. (3).

\[ \phi^{UNS} = \phi_n + \left. \frac{\partial \phi}{\partial x} \right|_n (x_f - x_n) \]

\[ + \left. \frac{\partial \phi}{\partial y} \right|_n (y_f - y_n) + \left. \frac{\partial \phi}{\partial z} \right|_n (z_f - z_n) + \text{HOT} \]
\[
\phi_{\text{RHS}} = \phi_n + \frac{\partial \phi}{\partial x} |_{x_j} (x_j - x_n) + \frac{\partial \phi}{\partial y} |_{y_j} (y_j - y_n) + \frac{\partial \phi}{\partial z} |_{z_j} (z_j - z_n) + \text{HOT}
\]

where \(\phi\) stands for the velocity components and pressure, the superscript \(\text{RHS}\) and \(\text{LHS}\) denote the approximation from the right-hand side and left-hand side of the interface respectively, and \(\text{HOT}\) represents higher order terms. By substituting Eq. (3) into Eq. (4), interfacial flow properties \(\phi_f\) can be obtained.

\[
\phi_f = \frac{1}{2}(\phi_{\text{RHS}} + \phi_{\text{LHS}}) - \frac{1}{2}\text{sign}(1, \hat{m})(\phi_{\text{RHS}} - \phi_{\text{LHS}})
\]

The gradients at the nodal points (cell centers) are evaluated by the Gauss's divergence theorem as below.

\[
\int \frac{\partial \phi}{\partial x} dV = \int_{\hat{A}} \phi n_x dA
\]

\[
\frac{\partial \phi}{\partial x} = \frac{1}{V} \sum_{i=1}^{N_{\text{face}}} \phi n_{ix} A_i
\]

where \(N_{\text{face}}\) is the total number of interfaces of the cell and \(V\) denotes the volume of the control volume cell.

The schematic diagram for diffusive fluxes is shown in Fig 1(b). The gradients at the interface can be evaluated by using Chain rule as Eq. (6).

\[
\frac{\partial \phi}{\partial x} = \frac{\partial \phi}{\partial \xi} + \frac{\partial \phi}{\partial \eta} + \frac{\partial \phi}{\partial \zeta}
\]

\[
\frac{\partial \phi}{\partial y} = \frac{\partial \phi}{\partial \xi} + \frac{\partial \phi}{\partial \eta} + \frac{\partial \phi}{\partial \zeta}
\]

\[
\frac{\partial \phi}{\partial z} = \frac{\partial \phi}{\partial \xi} + \frac{\partial \phi}{\partial \eta} + \frac{\partial \phi}{\partial \zeta}
\]

where the local coordinate system \((\xi, \eta, \zeta)\) is defined by the type of mesh separately.

For triangular mesh in Fig. 1(b), \(\xi\) is the vector form nodal point \(P_1\) to \(P_2\), \(\eta\) is the vector from vertex \(V_1\) to \(V_2\), and \(\Omega\) is the integration area for diffusive fluxes. The diffusive fluxes can be approximated by Eq. (7).

\[
\left( \frac{\partial \phi}{\partial x} \right)_f = \frac{1}{2\Omega} \left[ (\phi_{\xi} - \phi_{\eta}) (y_{\xi} - y_{\eta}) - (\phi_{\zeta} - \phi_{\eta}) (x_{\zeta} - x_{\eta}) \right]
\]

\[
\left( \frac{\partial \phi}{\partial y} \right)_f = \frac{1}{2\Omega} \left[ (\phi_{\xi} - \phi_{\eta}) (y_{\xi} - y_{\eta}) - (\phi_{\zeta} - \phi_{\eta}) (x_{\zeta} - x_{\eta}) \right]
\]

where \(\phi_{\xi}\) denotes the properties at nodal points and \(\phi_{\eta}\) denotes the properties at vertices. The values of vortices are obtained by averaging surrounding nodal value, in which inverse distances from all surrounding nodal points are considered as weighted function.

2.3 Center pressure based SIMPLE algorithm

By using an initial pressure field, \(P^0\), we can obtain \(u^0\), \(v^0\), and \(w^0\) by solving the momentum equations in an uncoupled form. The momentum equations can be written in the form as Eq. (8).
where the coefficients $a_{nb}$ and $a_c$ are

$$a_{nb} = \max(-m_j, 0) + \mu_j (\xi_j n_j + \xi_z n_z + \xi_n n_n) A,$$

$$a_c = \sum a_{nb},$$

and the RHS term can be written as

$$\text{RHS}_u = -\sum_{i=1}^{N_c} [m_j u^* - (r_{i1}^c - p^c) n_{i1} A - (r_{i2}^c) n_{i2} A - (r_{i3}^c) n_{i3} A] - \frac{\partial p^c}{\partial x} V_c,$$

$$\text{RHS}_v = -\sum_{i=1}^{N_c} [m_j v^* - (r_{i1}^c) n_{i1} A - (r_{i2}^c) n_{i2} A - (r_{i3}^c) n_{i3} A] - \frac{\partial p^c}{\partial y} V_c,$$

$$\text{RHS}_w = -\sum_{i=1}^{N_c} [m_j w^* - (r_{i1}^c) n_{i1} A - (r_{i2}^c) n_{i2} A - (r_{i3}^c) n_{i3} A] - \frac{\partial p^c}{\partial z} V_c,$$

where subscript $c$ denotes the cell we are solving, subscript $nb$ denotes the neighbor cells, and $A$ denotes the interfacial area. In this paper, we solve Eq. (8) by using Gauss-Seidel method. Then, we can obtain $u^*$, $v^*$, and $w^*$ by Eq. (9).

$$u^* = u^* + \Delta u$$

$$v^* = v^* + \Delta v$$

$$w^* = w^* + \Delta w$$

Although at this stage $u^*$, $v^*$, and $w^*$ satisfy the momentum equations, they do not necessarily satisfy the continuity equation. In order to satisfy the mass conservation, one has to interpolate the velocity to the interface. However, this interpolation will lead to the checkerboard solutions. In order to avoid the checkerboard solutions, one has to allow the interfacial velocity to be driven solely by the pressure difference. To achieve this aim without sacrificing the accuracy, one can divide the interpolated interfacial velocity into two components: one is the velocity component without the pressure contribution and the other is solely the pressure contribution. The former, which is at the cell center, can be written as:

$$\bar{u}^* = u^* + \frac{\partial p^c}{\partial x} \frac{V_c}{a_c},$$

$$\bar{v}^* = v^* + \frac{\partial p^c}{\partial y} \frac{V_c}{a_c},$$

$$\bar{w}^* = w^* + \frac{\partial p^c}{\partial z} \frac{V_c}{a_c},$$

The latter is obtained directly from the pressure difference of the two adjacent nodal points, $P_1$ and $P_2$ such that the interfacial velocity can be expressed as:
\[ u^*_f = \tilde{u}_f^* - \left( \frac{\partial p^*}{\partial x} \right)_{a_f} \frac{V_f}{a_f} \]
\[ v^*_f = \tilde{v}_f^* - \left( \frac{\partial p^*}{\partial y} \right)_{a_f} \frac{V_f}{a_f} \]
\[ w^*_f = \tilde{w}_f^* - \left( \frac{\partial p^*}{\partial z} \right)_{a_f} \frac{V_f}{a_f} \]  
\[(11)\]

Where \( V_f \) and \( a_f \) are obtained by interpolation to the interface.

We further assume that there are corrections to \( u_f^* \), \( v_f^* \), and \( w_f^* \), such that the continuity equation can be satisfied by using Eq. (12).

\[ \sum_{i=1}^{N_{\text{tot}}} \rho[ (u_f^* + \Delta u_f^*) n_1 + (v_f^* + \Delta v_f^*) n_2 + (w_f^* + \Delta w_f^*) n_3 ] A = 0 \]  
\[(12)\]

We can rewrite Eq. (12) as

\[ \sum_{i=1}^{N_{\text{tot}}} \rho[ (\Delta u_f^* n_1 + \Delta v_f^* n_2 + \Delta w_f^* n_3 ) A = - \sum_{i=1}^{N_{\text{tot}}} \rho[ u_f^* n_1 + v_f^* n_2 + w_f^* n_3 ] A \]  
\[(13)\]

where the right-hand side in Eq. (13) represents the mass imbalance in the control volume cell.

One assumes there is a corresponding pressure correction field, \( p' \), which drives the velocity corrections according to:

\[ \Delta u_f^* = - \left( \frac{\partial p'}{\partial x} \right) a_f \frac{V_f}{a_f} \]
\[ \Delta v_f^* = - \left( \frac{\partial p'}{\partial y} \right) a_f \frac{V_f}{a_f} \]
\[ \Delta w_f^* = - \left( \frac{\partial p'}{\partial z} \right) a_f \frac{V_f}{a_f} \]  
\[(14)\]

By substituting the velocity correction equations into the equation for the mass imbalance, we can obtain the equations of the pressure correction:

\[ a_c p^* = \sum_{ab} a_{ab} p_{ab}^* + b \]  
\[(15)\]

were \( a_{ab} \) and \( a_c \) in the continuity equation are

\[ a_{ab} = \rho \left( \frac{V_f}{a_f} \xi_1 n_1 + \frac{V_f}{a_f} \xi_2 n_2 + \frac{V_f}{a_f} \xi_3 n_3 \right) A, \]
\[ a_c = \sum_{ab} a_{ab}, \]

and

\[ b = - \sum_{ab} \rho[ u_f^* n_1 + v_f^* n_2 + w_f^* n_3 ] A. \]

Once the pressure correction is obtained, one can update the pressure field by:

\[ p^{*n+1} = p^* + \alpha_p p' \]  
\[(16)\]

where \( \alpha_p \) is the under-relaxation factor for pressure and is generally with a value of 0.5-0.8. Then the velocity correction on the interfaces as well as nodal points will be updated.
2.4 Partitioning approach

Figure 2 shows the schematic diagram of partitioning approach for center pressure based method, which is also implemented into UNCLE. In Fig. 2, blue points indicate vertices, red points indicate nodal points, and white points indicate the boundary points. By using this approach, the control volumes on the boundary are not split. Only communication of nodal values is needed for parallel computation which makes the implement of MPI in an unstructured grid more straightforward.

Excellent load balancing between the subgrids on each node is achieved through using METIS for domain decomposition. METIS can partition an unstructured grid into any integer number of zones without losing load balance. It is compatible with many platforms, convenient for running CFD codes on a variety of supercomputer to cluster architectures. Present partitioning approach has been tested by a number of two/three-dimensional geometries. All results show good load balances. In order to demonstrate the capability of this partitioning approach, this paper will present two cases—one is the grid for two-dimensional flow over circular cylinder in triangular mesh, and the other is the grid for three-dimensional flow over a circular cylinder in tetrahedron mesh. The definition of load-imbalance rate $L_{imb}$ and load-balance rate $L_B$ in this paper is defined as Eq. (17) and (18).

$$L_{imb} = \frac{|N_{node} - N_{avg}|}{N_{avg}} \times 100\%$$  \hspace{1cm} (17)

$$L_B = 1 - L_{imb}$$  \hspace{1cm} (18)

where $N_{node}$ is grid size of the node and $N_{avg}$ is the average grid size. By using load balance rate, we can compare the load balance quantitatively.

Figure 3(a) shows the partitioned grid for 2D flow over a circular cylinder in triangular mesh. The number of total grid points is 51,363 and the number of total cells is approximately 0.1M. The grid is partitioned to 16 zones for parallel computation. The cell distribution is not uniform, denser near the cylinder and coarser away. The load-balance distribution on each node is shown in Fig. 3(b). The x-axis indicates the node number and the y-axis indicates load-balance rate. The resulting load-balance rates are very close to 100% on every node with an average load balance rate of 98.37%. Figure 3(c) shows a partitioned grid for three-dimensional flow over circular cylinder with the internal grid distribution visible in a cut-away. The number of total grid points is approximately 0.3M and the number of total cells is approximately 1.3M. In this case, the grid is partitioned to 32 zones. The average load-balance rate is 97.9%. Our test results show that present partitioning approach has excellent load balance in two/three-dimensional grids with various types of meshes by using METIS for domain decomposition.

2.5 Time discretization

In this paper, a second-order fully implicit scheme is employed for the temporal discretization. In here, we take a one-dimensional equation example:

$$\frac{3\phi^{*n} - 4\phi^n + \phi^{n-1}}{2\Delta t} + \frac{\partial f(\phi^{*n})}{\partial x} = 0$$  \hspace{1cm} (19)

where $\phi$ is primitive variable, $f$ is interfacial flux, and the superscript $n$ indicates the index in time. A deferred iterative algorithm is employed to obtain $\phi^{n+1}$ by substituting (20) into (19),

$$(\phi^{n+1})^{m+1} = (\phi^{n+1})^m + (\Delta \phi)^n$$ \hspace{1cm} (20)

where the subscript $m$ stands for the sub iteration level. The final equation is
\frac{3(\Delta \phi)'' + \partial f(\Delta \phi)''}{2\Delta t} = \frac{(\phi'' - \phi^{*+1})}{2\Delta t} \frac{3((\phi^{*+1})'' - \phi''')}{2\Delta t} - \frac{\partial f((\phi^{*+1})'')}{\partial x} \tag{21}

The right-hand-side of Eq. (21) is explicit and can be implemented in a straightforward manner to discretize the spatial derivative term. The deferred iterative algorithm is strongly stable, and the solution \( \phi^{*+1} \) is obtained by using inner iterations to reach the convergent solution of the right-hand-side of Eq. (21), which means \( \Delta \phi \) is approximate to zero. A sub-iteration is performed at every time step so that this method is fully implicit.

3 Results

3.1 Two-dimensional laminar incompressible flow past a flat plate

In this case, two-dimensional laminar incompressible flow past a flat plate is simulated. The schematic diagram including geometry and boundary conditions is shown in Fig. 4. A uniform free stream boundary condition is imposed on the inlet. Because of the viscous effect, the laminar boundary layer begins to grow at the position \( x=0 \) on the plate. The initial condition for the entire computational domain is uniform free stream. A quadrilateral mesh of 600x200 is used for both Reynolds number (Re) at 5,000 and 50,000 based on a non-dimensional length scale of unity. The minimum distance from the wall is \( 5 \times 10^{-5} \) (corresponding to \( y^+ = 0.1 \)) which is fine enough to capture the phenomena within the boundary layer. In this case, the computational domain is divided into 16 zones. The parallel computation is performed on KFC3a, a 16 nodes PC cluster with Pentium IV 2.4 GHz CPU and gigabit network developed at the University of Kentucky. The results of this computation are compared to the standard Blasius solution for a laminar flat-plate boundary layer. Figure 5(a) shows the \( \eta = \sqrt{\text{Re}} \) versus \( u/U \) plot at \( x=4.5 \) from present results for \( \text{Re}=5,000 \) and 50,000 in comparison with Blasius solution. Both present results are good agreement with Blasius solution. Figure 5(b) shows the plot of momentum thickness (Re0) versus coefficient of skin friction (c_f) with present results and Blasius solution. Good agreement is obtained by comparing our present results with Blasius solution.

3.2 Two-dimensional driven cavity flow

In this section, two-dimensional incompressible flow in a square cavity at a Reynolds number of 400 is simulated. The fluid in the cavity is driven by a moving top with constant speed. Because driven cavity flow lacks an exact solution, an existing accurate numerical solution for this problem is used as a benchmark for comparing our results. Ghia et al. [3] presented numerical studies using the vorticity-stream function formulation for solutions up to \( \text{Re}=10,000 \) with 257x257 grid points, and these simulation results have been widely used as a benchmark for the driven cavity problem. The schematic diagram of this case with geometry and boundary conditions is shown in Fig. 6. The initial condition for the entire computational domain is stationary everywhere. In order to compare Ghia’s results, the number of grid points used is 257x257 or 66,049 and 65,536 cells are used in a quadrilateral mesh. For a triangular mesh, 66,546 cells, which is approximately the same as quadrilateral mesh, and 33,618 grid points are used in our computation. Figure 7(a) shows the u-velocity profile along the horizontal center line for both present results with quadrilateral and triangular mesh and Ghia’s result. Both present
results are in good agreement with Ghia's result. It also shows the present solution is identical and is independent of mesh types. Figure 7(b) shows the u-velocity profile along the horizontal center line; again, the results match. Figure 8 presents the u- and v-velocity contours and streamline plot from the present results on the quadrilateral mesh. In Fig. 8(a), the u-velocity contours ranges from -0.33 to 1.0 with 100 intervals. Solid lines indicate the positive values and dashed lines negative ones. In Fig. 8(b), the v-velocity contours ranges from 0.64 to 0.31 with 100 intervals. The flow structures including the location of the major vortex center, the bubble in the right bottom corner, and a small bubble in the left bottom corner are shown clearly in Figure 8(c), and are in good agreement with the results of Ghia.

3.3 Three-dimensional driven cavity flow
The three-dimensional version of the preceding problem is also a standard case for a new flow solver. In 1987, Ku et al. [4] simulated three-dimensional flow in a cubic cavity by using pseudospectral methods to solve the Navier-Stokes equations for Re = 100, 400, and 1000. In 2003, Shu et al. [5] repeated this problem by using the SIMPLE algorithm with the differential quadrature (DQ) method. They simulated the three-dimensional driven cavity flow at Re = 100, 200, 400 and 1000 and compared their results with Ku's results. In this section, we simulated this problem at Re = 400 and compared our results with those of Ku and Shu. In order to validate the results with different meshes, two grids are used to study this problem. One is a hexahedral mesh with 67x67x67 grid points and 287,496 cells, and the other is a tetrahedral mesh with 79,951 grid points and 446,953 cells. The geometry of this problem is a unit cube. The boundary condition at the y = 1 plane is uniform flow with u=1, v=0, and w=0, and all other boundary conditions are no-slip walls. The initial condition for the entire computational domain is stationary. Figure 9(a) shows the u-velocity profile at the horizontal centerline of the z=0.5 plane for the present hexahedral and tetrahedral mesh results as well as those of Ku and Shu. Figure 9(b) shows the v-velocity profile at the vertical centerline of z = 0.5 for the same set of simulations. Both present simulations show essentially identical solutions, and both are in good agreement Ku and Shu. Figure 10 shows the velocity vectors and pressure contours taken from the present simulation using the hexahedral mesh on the z=0.5, x=0.5, and y=0.5 planes respectively. As before, for the pressure contours dashed lines mean negative values. The established flow structures including the locations of the vortex center and the positive and negative regions in each plane are clearly visible and in good agreement with Ku’s and Shu’s results.

3.4 Two/Three-dimensional incompressible flow over a circular cylinder
Flow over a circular cylinder is a standard unsteady test problem. Many two-dimensional numerical studies have been done on this flow. In 1990, Rogers and Kwak [6] studied flow over a circular cylinder for Re = 200 using an upwind differencing scheme to solve the incompressible Navier-Stokes equations. They compared their results, which were obtained using 3rd order and 5th order upwind differencing schemes, with computational and experimental results. In 1995, Ku [7] studied this problem for Reynolds numbers 100, 250, 500, and 1,000 using a pseudospectral element method. Alonso et al. [8] studied the vortex shedding over a circular cylinder at Re = 500 with a multigrid unsteady Navier-Stokes solver with a flux-limited dissipation scheme in 1995. In 1998, Liu et al. [9] used preconditioned multigrid methods to study unsteady incompressible flows. They investigated flow over a circular cylinder at Re = 200 and their results was in good agreement with other computational and experimental results.

Examples of three-dimensional numerical studies of flow over a circular cylinder are the studies of Braza and Persillon [12] and Henderson [13]. There are also many experimental studies of flow over cylinder problems such as Roshko [14], Wille [15], and Williamson [16], [17]. According to Williamson [16], the phenomena of flow over a circular cylinder can be classified by Reynolds number. Williamson found the laminar vortex shedding region to occur for Reynolds numbers between 49 and 140-194. The three-dimensional wake transition region occurs for Re ~ 190 to 260. For the range between Re= 260 ~ 1000, the three-dimensional disorder of the wake begin to increase in at the fine scales. This evolves into the shear-layer transition regime for Reynolds numbers 1,000 up to 20,000. He also noted that three-dimensional effects occur for Re > 190. Further detailed explanations of the remaining regimes are reported in Williamson [16]. To date, most numerical studies about this problem only focused on a single Reynolds number and either two-dimensional or three-dimensional simulations exclusively. The current results encompass Reynolds numbers 100, 200, 300, 500, 1,000, and 1500 for two-dimensional simulations and 100 and 200 for three-dimensional simulations. These results are compared with the appropriate previous studies as well as current results cross-comparisons to examine the dimensional and flow regime effects.

Figure 11 shows a schematic diagram for flow over a circular cylinder with dimensions and boundary conditions. For three-dimensional simulation, the span of the cylinder is 10D, with D representing the reference length equal to the diameter of the cylinder. The boundary conditions at z = 0 and z = -10D are periodic, eliminating end effects. The initial condition for the entire domain is uniform flow as inflow for all simulations and the time step is 0.005 for all cases. The grid for two-dimensional simulations is a quadrilateral mesh with 22705 cells and 22925 grid points; for three-dimensional simulation, a hexahedral mesh with 1.13M cells and 1.17M grid points is used. Both grids are densely distributed near the cylinder and wake region and coarser near the outer region. All of the two-dimensional simulations are performed on the KFC3a cluster with 16 nodes, and all three-dimensional simulations are performed on KFC2, a 48 node commodity cluster with AMD Athlon 2000+ CPU and a channel-bonded network. It is noted that each three-dimensional simulation took approximately one week with 32 nodes on KFC2. Figure 12 presents the coefficient of Lift (CL) and the coefficient of drag (CD) unsteady histories of the present two-dimensional simulations for Re = 100, 200, 300, and 1,000 respectively. The Strouhal number (St) for these data sets is derived from the frequency of CL. In our simulations, higher Reynolds number cases achieve steady Strouhal numbers faster than lower Re cases, consistent with other computational results. Table 1 presents the summary of our present two- and three-dimensional results along with other computational and experimental results. As shown in Table 1, our current results show good agreement with previous data by comparing St, CL, and CD. Fig. 13 shows the two-dimensional vorticity contours for Re = 100, 200, 300 and 1000. In Fig. 13, the contours are range from -0.3 to 0.3. The three-dimensional \( \omega_z \) contours from the current simulations are similar to those generated by the two dimensional test cases. As seen in Fig. 13, the vortex shedding frequency increases as Re increases. For Re = 100, the vortices decay in the downstream. Because of the limited computational domain, we do not see the vortex
structures merge to large scale vortices in our simulation. For Re = 200, 500 and 1000, not only does the vortex strength decay, but also the vortex structures collapse in the far downstream and start to merge to large scale vortices. This same phenomenon is reported by Inoue and Yamazaki [18].

Figure 14(a)-(c) show the present three-dimensional results of $\omega_x$, $\omega_y$ and $\omega_z$ for Re = 100 and (d)-(f) for Re = 200 in the $y = 0$ plane. The contours of $\omega_x$ and $\omega_y$ range from -0.0005 to 0.0005 for Re = 100 and from -0.02 to 0.02 for Re = 200, and the contours of $\omega_z$ range from -0.3 to 0.3 for both cases where bright regions correspond to positive values and dark regions correspond to the negative values. Obviously, the magnitude of $\omega_z$ is much larger than the magnitudes of $\omega_x$ and $\omega_y$. For Re = 100, we can see that the magnitude of $\omega_x$ decreases in the upstream and the local minimum appears at the position of 5th “roll” in Fig. 14(a). After that, the magnitude of $\omega_x$ begins to grow. In Fig. 14(b), the magnitude of $\omega_y$ grows after the flow past the cylinder. The thickness of the vortex “roll” increases downstream. From Fig. 14(c), the magnitude of $\omega_z$ decays after the flow past the cylinder due to the energy dissipation. Because the three-dimensional effects are very weak at Re = 200, the three-dimensional outcome is very similar to two-dimensional case. For Re = 200, as seen from Fig. 14(d), the magnitude of $\omega_x$ decreases in the beginning and the local minimum also appears near the 5th “roll”. In the far downstream, a transition zone is observed, after which the vortex scales transit from small to large scales. In Fig. 14(e), unlike Fig. 14(b), wavy vortex structures are observed. Figure 14(f) shows that $\omega_z$ decays after the flow past the cylinder. Figure 15 shows the iso-surface of vorticity magnitude for Re = 200 from the present three-dimensional simulation result. The flow structures, especially the vortex streets in the wake regions, observed in our present results are in good agreement with other simulation results.

4 Conclusions

A center pressure based method is presented in this paper, and which has been implemented successfully to a new two/three-dimensional parallel unstructured incompressible Navier-Stokes solver, UNCLE, which has been developed at University of Kentucky. Implementation of using different types of meshes with center pressure based method is feasible and straightforward. In order to increase flexibility in complex geometries, center pressure based method has been extended to use a variety of grid types, such as triangular, quadrilateral, tetrahedral, and hexahedral meshes. Mesh independent tests are also made to prove that current method can generate identical solutions for different mesh types. By using METIS for domain decomposition, excellent parallel load balance is achieved. In this paper, several test cases are presented—laminar incompressible flow past a flat plate, steady two/three-dimensional driven cavity flow, and unsteady two/three-dimensional flow over a circular cylinder for low Reynolds numbers. All these test cases yielded good agreements in comparison with previous computational or experimental results. A complete qualitative and quantitative study of two-dimensional flow over a circular cylinder for low Reynolds number is presented in this paper, which can be further used as a benchmark solution set for the development of new unsteady Navier-stokes solvers.
References

Fig. 1 Schematic diagrams for integration areas. (a) convective fluxes, and (b) diffusive fluxes.

Fig. 2 A schematic diagram of cell-centered partitioning approach.
Fig. 3(a) Partitioned triangular mesh for 2D flow over a circular cylinder

Fig. 3(b) Load-balance distribution on each node in parallel computation

Fig. 3(c) Partitioned tetrahedral mesh for 3D flow over a circular cylinder

Fig. 4 Schematic diagram of laminar incompressible flow past flat plate
Fig. 5(a) Plot of $\eta$ versus $u/U$ at $x=4.5$ with present results and Blasius solution

Fig. 5(b) The plot of $Re_\theta$ versus $c_f$ with present results and Blasius solution

Fig. 6 The schematic diagram of two-dimensional driven cavity flow with boundary conditions
Fig. 7(a) The u-velocity profile along the horizontal center line for present results and Ghia’s result.

Fig. 7(b) The v-velocity profile along the vertical center line for present results and Ghia’s result.

Fig. 8(a) The u-velocity contour plot for two-dimensional driven cavity flow at Re=400.
Fig. 8(b) The v-velocity contour plot for two-dimensional driven cavity flow at Re=400.

Fig. 8(c) Streamline plot for two-dimensional driven cavity flow at Re=400.

Figure 9(a) The u-velocity profile at the horizontal centerline of z=0.5 plane with present results in hexahedral and tetrahedral meshes, and also Ku’s and Shu’s results.
Figure 9(b) The $v$-velocity profile at the vertical centerline of $z=0.5$ plane with present results, Ku's and Shu's results.
Figure 10(a)-(c) The velocity vector and (d)-(f) pressure contours from present results with hexahedral mesh at z=0.5, x=0.5, and y=0.5 planes respectively.
Fig. 11 Schematic diagram of flow over a circular cylinder with dimensions and boundary conditions.
Fig. 12 $C_L$ and $C_D$ history plots of two-dimensional simulation for (a) Re=100, (b) Re=200, (c) Re=300, and (d) Re=1000.

Fig. 13 Vorticity contours for two-dimensional simulation. (a) Re=100, (b) Re=200, (c) Re=300, and (d) Re=1000.
Fig. 14 Vorticity component contours for three-dimensional simulation at y=0 plane, where (a)-(c) $\omega_x$, $\omega_y$, and $\omega_z$ for Re=100 and (d)-(f) $\omega_x$, $\omega_y$, and $\omega_z$ for Re=200.

Fig. 15 The iso-surface of vorticity magnitude for Re = 200 from present three-dimensional simulation result.
<table>
<thead>
<tr>
<th>Re</th>
<th>100</th>
<th>200</th>
<th>300</th>
<th>500</th>
<th>1000</th>
<th>1500</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Present 2D results</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>$C_L = \pm 0.314$</td>
<td>$C_L = \pm 0.642$</td>
<td>$C_L = \pm 0.869$</td>
<td>$C_L = \pm 1.115$</td>
<td>$C_L = \pm 1.378$</td>
<td>$C_L = \pm 1.553$</td>
</tr>
<tr>
<td></td>
<td>$C_D = 1.325 \pm 0.008$</td>
<td>$C_D = 1.318 \pm 0.04$</td>
<td>$C_D = 1.354 \pm 0.072$</td>
<td>$C_D = 1.406 \pm 0.119$</td>
<td>$C_D = 1.489 \pm 0.198$</td>
<td>$C_D = 1.575 \pm 0.247$</td>
</tr>
<tr>
<td></td>
<td>$S_t = 0.165$</td>
<td>$S_t = 0.196$</td>
<td>$S_t = 0.21$</td>
<td>$S_t = 0.224$</td>
<td>$S_t = 0.239$</td>
<td>$S_t = 0.246$</td>
</tr>
<tr>
<td></td>
<td>Present 3D results</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>$C_L = \pm 0.322$</td>
<td>$C_L = \pm 0.664$</td>
<td>$C_L = \pm 0.869$</td>
<td>$C_L = \pm 1.115$</td>
<td>$C_L = \pm 1.378$</td>
<td>$C_L = \pm 1.553$</td>
</tr>
<tr>
<td></td>
<td>$C_D = 1.327 \pm 0.009$</td>
<td>$C_D = 1.324 \pm 0.042$</td>
<td>$C_D = 1.354 \pm 0.072$</td>
<td>$C_D = 1.406 \pm 0.119$</td>
<td>$C_D = 1.489 \pm 0.198$</td>
<td>$C_D = 1.575 \pm 0.247$</td>
</tr>
<tr>
<td></td>
<td>$S_t = 0.164$</td>
<td>$S_t = 0.195$</td>
<td>$S_t = 0.21$</td>
<td>$S_t = 0.224$</td>
<td>$S_t = 0.239$</td>
<td>$S_t = 0.246$</td>
</tr>
<tr>
<td></td>
<td>Computational 2D results</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Rogers and Kwak [6] (5th order)</td>
<td>$C_L = \pm 0.65$</td>
<td>$C_D = 1.23 \pm 0.05$</td>
<td>$S_t = 0.185$</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Alonso et al. [7]</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Ku [8]</td>
<td>$C_L = \pm 0.228$</td>
<td>$C_D = 1.33 - 1.358$</td>
<td>$S_t = 0.1675$</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Liu et al. [9]</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Ronald et al. [10]</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Qian and Vezza [11]</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Computational 3D results</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Braza and Persillon [12]</td>
<td></td>
<td></td>
<td></td>
<td>$S_t = 0.202$</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Henderson [13]</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>$S_t = 0.178$</td>
</tr>
<tr>
<td></td>
<td>Experimental results</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Roshko [14]</td>
<td></td>
<td></td>
<td>$S_t = 0.19$</td>
<td></td>
<td>$C_D = 1.2$</td>
</tr>
<tr>
<td></td>
<td>Wille [15]</td>
<td></td>
<td></td>
<td>$C_D = 1.3$</td>
<td></td>
<td>$S_t = 0.21$</td>
</tr>
<tr>
<td></td>
<td>Williamson [16]</td>
<td></td>
<td>$S_t = 0.166$</td>
<td>$S_t = 0.203$</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Williamson [17]</td>
<td></td>
<td></td>
<td>$S_t = 0.197$</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>