Comparison of Node-Centered and Cell-Centered Unstructured Finite-Volume Discretizations: Inviscid Fluxes

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Cell-centered and node-centered approaches have been compared for unstructured finite-volume discretization of inviscid fluxes. Regular and irregular grids are considered, including mixed-element grids and grids with random perturbations of nodes. Complexity, accuracy, and convergence rates of defect-correction iterations are studied for eight nominally second-order accurate schemes: two node-centered schemes, NC and NC-WLSQ, with respective unweighted and weighted least-square methods for gradient reconstruction and six cell-centered schemes – two node-averaging schemes with and without clipping and four schemes that employ different stencils for least-square gradient reconstruction. The cell-centered nearest-neighbor (CC-NN) scheme has the lowest complexity; a version of the scheme that involves smart augmentation of the least-square stencil (CC-SA) has only marginal complexity increase. All other schemes have larger complexity; complexity of node-centered schemes are somewhat lower than complexity of cell-centered node-averaging (CC-NA) and full-augmentation (CC-FA) schemes.

On highly anisotropic grids typical of those encountered in grid adaptation, discretization errors of five of the six cell-centered schemes converge with second order on all tested grids; the CC-NA scheme with clipping degrades solution accuracy to first order. The node-centered solutions converge with second order on regular and/or triangular grids and with first order on perturbed quadrilateral and mixed-element grids. All schemes may produce large relative errors in gradient reconstruction on grids with perturbed nodes. Defect-correction iterations for schemes employing weighted least-square gradient reconstruction diverge on perturbed stretched grids. Overall, the CC-NN and CC-SA schemes offer the best options of the lowest complexity and small second-order discretization errors.

On anisotropic grids over a curved body typical of turbulent flow simulations, the discretization errors are comparable for all schemes except the CC-NA scheme; the latter may produce large discretization errors explained by node-averaging degeneration. Accurate gradients can be reconstructed by least-square methods with a local approximate mapping; without mapping, only the NC-WLSQ scheme provides accurate gradients. Defect-correction iterations may diverge for the CC-NA and NC-WLSQ schemes; the iterations converge fast for the CC-SA and CC-FA schemes on all grids, and converge slower for other schemes on at least some irregular grids.

I. Introduction

Both node-centered and cell-centered finite-volume discretization schemes are widely used for complex three-dimensional turbulent simulations in aerospace applications. The relative advantages of the two approaches have been extensively studied in the search for methods that are accurate, efficient, and robust over the broadest possible range of grid and solution parameters. The topic was discussed in a panel session at the 2007 AIAA Computational Fluid Dynamics conference, but a consensus did not emerge. One of the difficulties in assessing the two approaches is that comparative calculations were not completed in a controlled environment, i.e., computations were made with different codes and different degrees of freedom and the exact solutions were not known.

In this paper, we provide a controlled environment for comparing a subset of the discretization elements needed in turbulent simulations, namely that of the inviscid discretization. In particular, we consider a constant-coefficient convection equation as a model for inviscid fluxes. This paper is second in a series of papers on comparison of cell-centered and node-centered finite-volume discretizations. It follows Ref. [1], which considered viscous discretizations.

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The ultimate objective of the effort is to identify methods suitable for a uniformly second-order accurate and efficient unstructured-grid solver for the Reynolds-Averaged Navier-Stokes equations.

In this work, we use the method of manufactured solution so the exact solutions are known and chosen to be smooth on all grids considered, i.e., no accuracy degradation occurs because of lack of solution smoothness. Computational studies are conducted on two-dimensional grids ranging from structured (regular) grids to irregular grids composed of arbitrary mixtures of triangles and quadrilaterals. Highly irregular grids are deliberately constructed through random perturbations of structured grids to bring out the worst possible behavior of the solution. Two classes of tests are considered. The first class of tests involves smooth manufactured solutions on both isotropic and highly anisotropic grids with discontinuous metrics, typical of those encountered in grid adaptation. The second class of tests concerns solutions and grids varying strongly anisotropically over a curved body, typical of those encountered in high-Reynolds number turbulent flow simulations.

There are eight main schemes considered — two representative node-centered schemes with weighted and unweighted least-square methods for gradient reconstruction and six cell-centered schemes. The cell-centered schemes include node-averaging schemes with and without clipping and four least-square gradient reconstruction schemes that are named according to the stencil used for the least-square fit: a nearest-neighbor scheme uses only face-neighboring cells; a smart-augmentation scheme minimally augments the nearest-neighbor stencil; two full augmentation schemes with and without weighting use larger stencils that include all node-sharing cells. Each of the schemes considered is nominally second-order accurate.

For the second class of tests, the approximately mapped least-square approach introduced in Ref. [1] is used to improve gradient reconstruction accuracy on curved high-aspect-ratio grids. The mapping uses the distance function commonly available in practical codes and can be used with any scheme.

The properties to be compared in this study are computational complexity (operation count) and discretization accuracy at equivalent numbers of degrees of freedom as well as convergence rates of defect-correction iterations with a first-order driver.

The material in this paper is presented in the following order. Section II introduces the computational grids used in the current study. A brief explanation of finite-volume discretizations in Section III is followed by the estimates of discretization complexity for two- and three-dimensional grids given in Section IV. Section V outlines the analysis methods used in this study. A brief introduction of the model equation in Section VI precedes results provided in Section VII on accuracy of finite-volume solutions and gradients and on convergence rates of defect-correction iterations observed on isotropic irregular grids. Section VIII compares the finite-volume discretizations on stretched highly anisotropic grids in rectangular geometries. Section IX provides comparisons for irregular high-aspect-ratio grids in curved geometries. Conclusions and recommendations are offered in Section X.

II. Grids

This paper studies finite-volume discretization (FVD) schemes for inviscid fluxes on grids that are defined as irregular. A grid is classified as regular if it can be derived by a smooth mapping from a grid with (1) a periodic node connectivity pattern (i.e., the number of edges per node changes periodically) and (2) a periodic cell distribution (i.e., the grid is composed of periodically repeated combinations of cells). Regular grids include, but are not limited to, grids derived from Cartesian ones — triangular grids obtained by diagonal splitting with a periodic pattern, smoothly stretched grids, skewed grids, smooth curvilinear grids, etc. Grids that are not regular are called irregular grids. We are especially interested in unstructured grids, e.g., grids with the number of edges changing from node to node with no pattern.

The regular and irregular grids considered in this paper are derived from an underlying (possibly mapped) Cartesian grid in two dimensions (2D) with mesh sizes \(h_x\) and \(h_y\) and the aspect ratio \(A = h_x/h_y\); both mesh sizes of the underlying grid are assumed to be small, \(h_y \ll 1, h_x \ll 1\). Irregularities are introduced locally and do not affect grid topology and metrics outside of a few neighboring cells. A local grid perturbation is called random if it is independent of local perturbations introduced beyond some immediate neighborhood. For computational grids generated for the reported studies, local and random grid irregularities are introduced in two ways: (1) the quadrilateral cells of the underlying grid are randomly split (or not split) into triangles; (2) the grid nodes are perturbed from their original positions by random shifts, where the shifts are fractions of a local mesh size.

Four basic grid types are considered: (I) regular quadrilateral (i.e., mapped Cartesian) grids; (II) regular triangular grids derived from the regular quadrilateral grids by the same diagonal splitting of each quadrilateral; (III) random triangular grids, in which regular quadrilateral are split by randomly chosen diagonals, each diagonal orienta-
tion occurring with probability of half; \((IV)\) random mixed-element grids, in which regular quadrilateral are randomly split or not split by diagonals; the splitting probability is half; in case of splitting, each diagonal orientation is chosen with probability of half. Nodes of any basic-type grid can be perturbed from their initial positions by random shifts, thus leading to four additional perturbed grid types which are designated by subscript \(p\) as \((I_p)-(IV_p)\). Grids of types \((III) - (IV)\) and \((III_p) - (IV_p)\) are irregular (and unstructured) because there is no periodic connectivity pattern. All perturbed grids are irregular because there is no periodic cell distribution. The representative grids are shown in Figure 1.

![Figure 1. Typical regular and irregular grids.](image)

Our main interest is the accuracy of FVD schemes on general irregular (mostly unstructured) grids with a minimum set of constraints. In particular, we do not require any grid smoothness, neither on individual grids nor in the limit of grid refinement. The only major requirement for a sequence of refined grids is to satisfy the consistent refinement property. The property requires the maximum distance across the grid cells to decrease consistently with increase of the total number of grid points, \(N\). In particular, the maximum distance should tend to zero as \(N^{-1/2}\) in 2D computations. For unstructured grids in three dimensions (3D), the consistent refinement property has been studied elsewhere. On 2D grids, the effective mesh size, \(h_e\), is computed as the \(L_1\) norm of the square root of the control volumes. Note that on consistently refined grids considered in this paper, control volumes converge in grid refinement with the same rate in any norm.

The locations of discrete solutions are called data points. For consistency with the 3D terminology, the 2D cell boundaries are called faces, and the term “edge” refers to a line, possibly virtual, connecting the neighboring data points. Each face is characterized by the directed-area vector, which is directed outwardly normal to the face with the amplitude equal to the face area.

The random node perturbation in each dimension is defined as \(\frac{1}{\sqrt{2}}\rho h\), where \(\rho \in [-1, 1]\) is a random number and \(h\) is the local mesh size along the given dimension. With these perturbations, triangular cells in the rectangular geometry can approach topological degeneration and zero volume. The random perturbations are introduced independently on all grids in grid refinement implying that grids of types \((I_p) - (IV_p)\) are grids with discontinuous metrics, e.g., ratios of neighboring cell volumes and face areas are random on all grids and do not approach unity in the limit of grid refinement.
III. Finite-volume discretization schemes

The FVD schemes are derived from the integral form of a conservation law

\[ \int_{\partial \Omega} F \cdot \hat{n} \, ds = \iint_{\Omega} f \, d\Omega, \]

where \( \Omega \) is a control volume with boundary \( \partial \Omega \), \( F \) is the flux, \( \hat{n} \) is the outward unit normal vector, \( ds \) is the area differential, and \( f \) is a forcing function. The general FVD approach requires partitioning the domain into a set of non-overlapping control volumes and numerically implementing equation (1) over each control volume.

![Control-volume partitioning for finite-volume discretizations. Numbers 0 – 12 and letters A – L denote grid nodes and primal cell centers, respectively. The control volume for a node-centered discretization around the grid node 0 is shaded. The control volume for a cell-centered discretization around the cell center A is hashed.](image)

Cell-centered (CC) discretizations assume solutions are defined at the centers of the primal grid cells with the primal cells serving as the control volumes. The cell center coordinates are typically defined as the averages of the coordinates of the cell’s vertices. Note that for mixed-element grids, cell centers are not necessarily centroids.

Node-centered (NC) discretizations assume solutions are defined at the primal mesh nodes. For NC schemes, control volumes are constructed around the mesh nodes by the median-dual partition: the centers of primal cells are connected with the midpoints of the surrounding faces. These non-overlapping control volumes cover the entire computational domain and compose a mesh that is dual to the primal mesh. Both cell-centered and node-centered control-volume partitions are illustrated in Figure 2.

The flux, \( F^h \cdot \hat{n} \), at a control-volume face is computed according to the Roe scheme,

\[ (F^h \cdot \hat{n}) = \frac{1}{2} \left[ (F_R \cdot \hat{n}) + (F_L \cdot \hat{n}) \right] - \frac{1}{2} |\hat{A}| \left( Q_R - Q_L \right), \]

where \( Q_L \) and \( Q_R \) are the “left” and “right” solution reconstructions; \( F_L \) and \( F_R \) are the corresponding “left” and “right” numerical fluxes; \( |\hat{A}| \) is the Roe’s approximate Riemann solver matrix. The solutions \( Q_L \) and \( Q_R \) are linearly reconstructed at the face by using solutions defined at the control volume centers and solution gradients reconstructed at each control volume. Various FVD schemes differ in the way they reconstruct gradients at the control volumes.

For cell-centered schemes, the face-based flux integration over a control-volume face is approximated as the inner product of \( F^h \) computed at the face center and the face directed area vector. The integration scheme is second-order accurate on grids of all types. For node-centered schemes, the edge-based flux integration scheme approximates the integrated flux through the two faces linked at an edge midpoint by multiplying \( F^h \) computed at the edge midpoint with the combined-directed-area vector. The integration scheme is computationally efficient and second-order accurate on regular and triangular grids of types (I), (II), (III), (IIp), and (IIIp); the integration accuracy degenerates to first order on mixed-element and perturbed quadrilateral grids of types (IV), (IVp), and (Ip). Note that node-centered schemes that avoid accuracy degradation can be constructed but at a higher cost (see estimates in Section IV).

The forcing term integration over the control volume is approximated as the value at the control-volume center multiplied by the volume \( |\Omega| \). This approximation is second-order accurate when the control-volume center coincides...
with the centroid. On general irregular grids, the control-volume center is not necessarily the centroid, and the approximation becomes locally first-order accurate. However, with grid irregularities introduced locally and randomly (thus, implying a zero-mean distribution of the deviations between control-volume centers and centroids), the integral of the forcing term over any sub-domain of size $O(1)$ is approximated with second order.

### A. Cell-centered schemes

#### 1. Node averaging schemes

In the cell-centered node-averaging (CC-NA) schemes, the solution values are first reconstructed at the nodes from the surrounding cell centers. With respect to Figure 2, the solution at the node $0$ is reconstructed by averaging solutions defined at the cell centers $A$, $B$, and $C$. The solution reconstruction proposed in Refs. [6, 7] and used in Ref. [8] is an averaging procedure that is based on a constrained optimization to satisfy some Laplacian properties. The scheme is second-order accurate and stable when the coefficients of the introduced pseudo-Laplacian operator are close to 1. It has been shown$^9$ that this averaging procedure is equivalent to an unweighted least-square linear fit.

The gradient of solution $U$ at the cell $\Omega$ is reconstructed by the Green-Gauss formula,

$$\nabla U = \frac{1}{|\Omega|} \int_{\partial \Omega} U \hat{n} ds,$$

where

$$|\Omega| = \iint_{\Omega} d\Omega$$

is the cell volume. For second-order accuracy, the solution at a face is computed by averaging the values at the face nodes and the integral over the face is approximated by the product of the solution and the face directed area. For simplicial grids, an explicit formula relating the gradient with the nodal solution values is given elsewhere.$^8$

On highly stretched and deformed grids, some coefficients of the pseudo-Laplacian may become negative or larger than 2, which has a detrimental effect on stability and robustness.$^{10,11}$ Holmes and Connell$^6$ proposed to enforce stability by clipping the coefficients between 0 and 2. The CC-NA scheme with clipping (CC-NA-CLIP) represents a current standard in practical computational fluid dynamics for applications involving cell-centered finite volume formulations.$^{12}$ As shown further in the paper, clipping seriously degrades the accuracy of the solution.

#### 2. Least-square schemes

An alternative approach relies on a least-square method for gradient reconstruction, in which the linear approximation obtained at a control volume coincides with the solution value at the control volume center. In this paper, both weighted and unweighted least-square methods are considered. The weighted method is designated as WLSQ herein and the unweighted method is used as default without designation. In the WLSQ method, the contributions to the minimized functional are weighted with weights inversely proportional to the distance from the control-volume center. In the unweighted method, all contributions are equally weighted.

The stencils used in the least-square minimization are discussed with respect to Figure 2. Three types of stencils are considered — nearest neighbor (NN), full augmentation (FA), and smart augmentation (SA) stencils. The NN stencil involves only centers of face-neighbor cells; the FA stencil includes all the cells that share a vertex with the given cell, i.e., all the cells involved in CC-NA gradient reconstruction; the SA stencil is an adaptive stencil that provides a minimal extension of the NN stencil necessary to improve convergence rates of the defect-correction iterations (DCI) with the first-order cell-centered FVD scheme as the driver. For cell-center $A$, the NN stencil includes neighbors $B$, $C$, $D$, and $E$; the FA stencil includes additionally neighbors $F$, $G$, $H$, $I$, $J$, $K$, and $L$; the SA applies an augmentation test to the NN stencil and expands it only if necessary by choosing only appropriate cells from the augmentation pool provided by the FA method.

Initially, the CC-SA scheme is identical to the CC-NN scheme. In stencil augmentation at each cell, the augmentation test computes the quantity $C_{i_c} = |1 - d_{S,A}/d_1|$, where $d_{S,A}$ and $d_1$ are the respective main-diagonal coefficients of full linearizations of the current CC-SA and the first-order driver schemes for a constant-coefficient convection operator. The test is applied for a number of representative convection directions indexed by $i_c$, e.g., all the edge directions associated with the cell. In the algorithm implemented for this paper, the current CC-SA scheme is considered sufficiently augmented if the augmentation indicator...
\[ AI = \max_{i} C_{i} < \epsilon, \]  

where \( \epsilon = 0.4 \) is a user-defined tolerance. Smaller values of \( \epsilon \) imply larger CC-SA stencils. If augmentation is required, a cell from the augmentation pool is added to the stencil. The cells from the pool are tested one by one until a cell that brings \( AI \) below the \( \epsilon \)-threshold is found. If no such single cell has been found, the cell that makes the best improvement in \( AI \) is added to the stencil, and the augmentation procedure repeats. Note that it is possible that at the end, the user-defined tolerance has not been achieved. Even in these instances, the smart augmentation adds only cells that reduce \( AI \), thus, providing a much smaller stencil than CC-FA stencil even in the worst-case scenario.

B. Node-centered schemes

For the node-centered computations, the current standard employs a least-square gradient reconstruction. The typical stencil at a control volume involves all nodes linked by an edge. For example, with reference to Figure 2, the least-square fit for the shaded control volume centered at node \( 0 \) includes nodes 1, 2, and 4. Both weighted and unweighted least-square methods are evaluated.

IV. Complexity

A. Flux integration complexity

In this section, the complexity associated with flux integration in 3D cell-centered or node-centered FVD schemes is estimated. The complexity is measured as the number of flux-reconstruction instances required for one residual evaluation. Flux reconstructions are the main contributors to the operation counts associated with flux integration; other aspects of the discretization, such as determining the solution values or solution-gradient values require additional considerations. Three types of primal meshes are considered: (1) fully-tetrahedral, (2) fully-prismatic, (3) fully-hexahedral.

An underlying Cartesian grid is considered and split into the various elements. The splitting into tetrahedra assumes each hexahedral defined by the grid is split into 5 tetrahedra with one of the tetrahedra being completely interior to the hexahedral (i.e., its faces are not aligned with any of the hexahedral faces – see Figure 3). Note that there are other partition strategies that lead to different number of tetrahedra per hexahedral; for example, dividing the hexahedral into two triangular prisms with subsequent division of each of the prisms into 3 tetrahedra leads to 6 tetrahedra per hexahedral. In this section we do not consider other possible partitions.

![Figure 3. Splitting hexahedral into 5 tetrahedra.](image)

Table 1 shows complexity estimates for one cell-centered and two node-centered FVD schemes. Only interior discretizations are estimated; boundary effects are neglected. The cell-centered formulation uses a face-based flux integration scheme with one flux reconstruction per control face. The first node-centered scheme is a linear FVD scheme with edge-based flux integration; the second node-centered scheme is a linear FVD scheme with face-based flux integration.

Both node-centered FVD schemes assume a median-dual partition of the domain. In such a partition, the constituent dual control volumes are bounded by generally non-planar dual faces formed by connecting 3 types of points:
(1) edge midpoints, (2) element-face centroids, and (3) element centroids. FVD schemes with edge-based flux integration approximate integration over all of the constituent dual faces adjacent to an edge by evaluating the flux at the edge midpoint using a lumped directed area. FVD schemes with face-based flux integration reconstruct fluxes at each of the constituent dual faces separately and use local directed areas. For the present estimation, we assume that each flux-reconstruction instance requires the same operation count; in particular, the approximate Riemann solver is applied at each reconstruction point. In fact, significant savings for face-based flux integration can be achieved if the dissipation matrix is computed once for all control surfaces surrounding an edge.

Two estimates of complexity are given. The first estimate assumes that any constituent quadrilateral face in the control surface is broken into two triangular faces. The second estimate (in parentheses) assumes any constituent quadrilateral face is approximated as planar. The former is required to ensure a precise (water-tight) definition of the control surface and can serve as a measure of the complexity in integration of the physical flux terms. The latter can serve as an estimate of the complexity associated with numerical dissipation terms, in which details of the control-surface can be neglected.

<table>
<thead>
<tr>
<th>Elements</th>
<th>Cell-centered face-based flux integration</th>
<th>Node-centered edge-based flux integration</th>
<th>Node-centered face-based flux integration</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tetrahedral</td>
<td>4 (4)</td>
<td>12</td>
<td>120 (60)</td>
</tr>
<tr>
<td>Prismatic</td>
<td>8 (5)</td>
<td>8</td>
<td>72 (36)</td>
</tr>
<tr>
<td>Hexahedral</td>
<td>12 (6)</td>
<td>6</td>
<td>48 (24)</td>
</tr>
</tbody>
</table>

Table 1. Number of flux-reconstruction instances per equation for FVD discretizations with precise (water-tight) definition of the control-volume boundary; in parentheses, the number of flux-reconstruction instances with planar approximations to the control-volume boundary.

The complexities of cell-centered and node-centered FVD schemes with edge-based flux integration are reasonably close. Unfortunately, as shown in this paper and also previously,\textsuperscript{2,4,5} the accuracy of the edge-reconstruction FVD scheme degenerates to first order on perturbed quadrilateral and general mixed-element grids. To maintain the second-order accuracy on general grids, one can employ the node-centered scheme with face-based flux integration, but the integration complexity of this formulation substantially exceeds the complexity of the cell-centered FVD scheme. These results are in agreement with the observations made by Delanaye and Liu\textsuperscript{13} leading to the selection of a cell-centered discretization.

B. Size of inviscid stencil

Another important measure of complexity of an FVD scheme is the size of the full-linearization stencil. The size of the 2D and 3D full-linearization stencil is examined for the inviscid cell-centered and node-centered FVD schemes. Cartesian meshes are split into triangular and tetrahedral elements, as in the previous section, again neglecting boundary effects. Estimates are compared to numerical calculations on an actual 3D grid that includes boundary effects; the grid is a viscous fully-tetrahedral grid composed of 16,391 nodes.

In 3D half of the grid nodes have 18 adjacent edges (32 adjacent tetrahedra) and half have 6 adjacent edges (8 adjacent tetrahedra). Each of the tetrahedra interior to an originally-hexahedral cell is defined by four nodes, each with 18 adjacent edges. Each of the four surrounding tetrahedra within an originally-hexahedral cell is defined by three nodes with 18 adjacent edges and 1 node with 6 adjacent edges.

For reference, Table 2 shows the average and maximum number of edges, \( n_{edge} \), connecting to a grid node. The average number of connecting edges sets the least-square stencil size for the node-centered scheme as \( n_{edge} + 1 \). The number of connecting edges is an important factor for the NC, CC-NA, and CC-FA schemes because it directly defines the size of the full-linearization stencil. Generally speaking, the number of edges is not bounded in 3D and, thus, the corresponding NC, CC-NA, and CC-FA stencil sizes are not bounded.

<table>
<thead>
<tr>
<th>Dimension</th>
<th>( n_{edge} ) (Average)</th>
<th>( n_{edge} ) (Maximum)</th>
</tr>
</thead>
<tbody>
<tr>
<td>2D</td>
<td>6</td>
<td>8</td>
</tr>
<tr>
<td>3D</td>
<td>12</td>
<td>18</td>
</tr>
</tbody>
</table>

Table 2. Edges connecting to a grid node in the split Cartesian grids. Figure 4 provides a 2D illustration.
For the inviscid discretization, the DCI with a first-order driver is generally used to converge the residual; thus, it is important to consider first-order and second-order linearizations. For the first-order cell-centered FVD scheme, the size of the linearization stencil is simply the number of faces plus one (to account for the central node). For the first-order node-centered discretization, the size of the linearization stencil is the number of edges connecting to a node plus one. Table 3 shows 2D and 3D linearization stencil sizes. The cell-centered discretization has nearly a factor of 3 smaller stencil in 3D.

<table>
<thead>
<tr>
<th>Elements</th>
<th>Node-centered</th>
<th>Cell-centered</th>
</tr>
</thead>
<tbody>
<tr>
<td>Estimate 2D</td>
<td>7</td>
<td>4</td>
</tr>
<tr>
<td>Estimate 3D</td>
<td>13</td>
<td>5</td>
</tr>
<tr>
<td>Numerical 3D</td>
<td>14</td>
<td>5</td>
</tr>
</tbody>
</table>

Table 3. Average size of the inviscid first-order FVD stencil on triangular/tetrahedral grids in 2D/3D.

For second-order accuracy, all schemes reconstruct gradients at the control volumes. The node-centered discretizations use a least-square approach and require solutions at the neighbor-of-neighbor nodes and a correspondingly large linearization stencil. The cell-centered CC-NA schemes have even larger linearization stencils which include all cells contributing to solution reconstruction at any node of a face-neighboring cell. Stencils of CC-FA schemes are the same as CC-NA stencils. The CC-NN stencil also uses a least-square approach to fitting the gradient in reconstruction, but requires a much smaller stencil which includes only neighbor-of-neighbor cells. Table 4 shows stencil sizes for 2D and 3D. The splittings used in the estimates are shown in Figures 3 and 4 for 3D and 2D, respectively. In 3D, the NC stencil is significantly smaller than the CC-NA and CC-FA stencils. In both 2D and 3D, the CC-NN stencil is the smallest.

<table>
<thead>
<tr>
<th>Elements</th>
<th>NC</th>
<th>CC-NA</th>
<th>CC-NN</th>
</tr>
</thead>
<tbody>
<tr>
<td>Estimate 2D</td>
<td>23</td>
<td>25</td>
<td>9</td>
</tr>
<tr>
<td>Estimate 3D</td>
<td>75</td>
<td>139</td>
<td>15</td>
</tr>
<tr>
<td>Numerical 3D</td>
<td>63</td>
<td>118</td>
<td>15</td>
</tr>
</tbody>
</table>

Table 4. Average size of the inviscid second-order stencil for 2D/3D discretizations with triangular/tetrahedral elements.

The numbers are so striking that it is useful to show the 2D stencils for a single shaded control volume in Figure 4 for each approach. The stencil sizes are 25, 25, and 9 for the NC, CC-NA, and CC-NN schemes, respectively. Note that the stencil size for the NC control-volume with four edges adjacent to the one shown in Figure 4 is 21; thus, the average of 23 is shown in Table 4. Also, for the 3D NC schemes, the nodes with 6 and 18 edges have stencil sizes of 57 and 93, respectively; thus, the average of 75 is shown in the table. For the CC-NA and CC-FA schemes, the cells at the corners of the original Cartesian cell have a stencil size of 149 and those fully interior to the original Cartesian cell have a stencil size of 99. Since there is one interior tetrahedron for each of the four corner tetrahedrons, the average of 139 is shown in the table.

V. Analysis

Accuracy of FVD schemes is analyzed for known exact or manufactured solutions. The forcing function and boundary values are found by substituting this solution into the governing equations, including boundary conditions. The discrete forcing function is defined at the data points.

A. Discretization error

The main accuracy measure is the discretization error, \( E_d \), which is defined as the difference between the exact discrete solution, \( U^h \), of the discretized equations (1) and the exact continuous solution, \( U \), to the corresponding differential equations.

\[
E_d = U^h - \bar{U}^h, \tag{6}
\]

where \( U^h \) is \( U \) sampled at data points.
B. Truncation error

Another accuracy measure commonly used in computations is truncation error. Truncation error, $E_t$, characterizes the local accuracy of approximating the differential equations. For finite differences, it is defined as the residual obtained after substituting the exact solution $U$ into the discretized differential equations.\(^{(14)}\) For FVD schemes, the traditional truncation error is usually defined from the time-dependent standpoint.\(^{(15,16)}\) In the steady-state limit, it is defined (e.g., in Ref. [18]) as the residual computed after substituting $U$ into the normalized discrete equations (1),

$$E_t = \frac{1}{|\Omega|} \left[ - \iiint_{\Omega} f^h d\Omega + \oint_{\partial \Omega} (F^h \cdot \hat{n}) \, ds \right],$$

(7)

where $F^h$ is evaluated at the control-volume boundary, $f^h$ is an approximation of the forcing function $f$ on $\Omega$, and the integrals are computed according to some quadrature formulas. Note that convergence of truncation errors is expected to show the order property only on regular grids;\(^{(17)}\) on irregular grids, it has been long known that the design-order discretization-error convergence can be achieved even when truncation errors exhibit a lower-order convergence or, in some cases, do not converge at all.\(^{(18-23)}\)

C. Accuracy of gradient reconstruction

Yet another important accuracy measure is the accuracy of gradient approximation at a control-volume. For second-order convergence of discretization errors, the gradient accuracy is usually required to be at least first order. For each control-volume, accuracy of the gradient is evaluated by comparing the reconstructed gradient, $\nabla r$, with the exact gradient, $\nabla_{\text{exact}}$, computed at the control-volume center. The accuracy of gradient reconstruction is measured as the relative gradient error:

$$E_{\text{rel}} = \frac{\| \epsilon \|}{\| G \|},$$

(8)

where functions $\epsilon$ and $G$ are amplitudes of the gradient error and the exact gradient, respectively, evaluated at control-volume centers;

$$\epsilon = |\nabla r U^h - \nabla U|, \quad \text{and} \quad G = |\nabla U|;$$

(9)

$\| \cdot \|$ is a norm of interest.

D. Convergence of defect-correction iterations (DCI)

Besides accuracy, an important quality of a practical discretization is availability of an affordable solver. For FVD schemes with low complexity, such as CC-NN and CC-SA, an efficient solution method would use a full linearization in relaxation of the target FVD scheme. For FVD schemes with high complexity, such as CC-NA, CC-FA, and NC
schemes, iterations with the full linearization are not affordable; DCI schemes with linearized first-order drivers are common methods used in practical computations.

Let \( u^h \) be the current solution approximation. The DCI method is defined in the following two steps:

1. The correction \( v^h \) is calculated from
   \[
   L^h_d u^h = R^h (u^h),
   \]
   where \( R^h (u^h) \) is the residual of the target FVD scheme and \( L^h_d \) is a driver scheme. Equation (10) is solved to zero residual.

2. The current approximation is corrected
   \[
   u^h = u^h + v^h.
   \]

The DCI method considered in this paper is applied to second-order FVD schemes and uses the corresponding first-order upwind FVD scheme as a driver. In tests, the forcing term and the boundary conditions are set to zero so the exact solution is zero. The initial solution perturbation is random.

VI. Convection equation

The linear convection equation
   \[
   (a \cdot \nabla) U = f
   \]
   is considered as a model for inviscid fluxes; \( a = \left( \sin \left( \frac{\pi x}{16} \right), \cos \left( \frac{\pi y}{16} \right) \right) \) is a constant vector, \( f \) is a forcing function. Boundary conditions are over-specified conditions, in which solution at any control volume that includes a nodes connected by an edge to the boundary is over-specified from the manufactured solution.

VII. Isotropic irregular grids

A. Grid refinement

All computations in this section are performed for the manufactured solution \( U = -\cos(2\pi x - \pi y) \). Sequences of consistently refined grids of types \((III_p)\) and \((IV_p)\) are generated on the unit square \([0, 1] \times [0, 1]\). Irregularities are introduced at each grid independently, so the grid metrics remain discontinuous on all the grids. The ratio of areas of neighboring faces can be as large as \(3\sqrt{2}\); because a control volume can be arbitrarily small, the ratio of the neighboring volumes can be arbitrarily high. Two node-centered and six cell-centered schemes are considered: NC, NC-WLSQ, CC-SA, CC-NN, CC-FA, CC-FA-WLSQ, CC-NA and CC-NA-CLIP. On grids of type \((III_p)\), CC-SA scheme augments about 50% of the interior least-square stencils and CC-NA-CLIP clips about 10% of the interior nodes. On grids of type \((IV_p)\), CC-SA scheme augments between 25% and 30% of the interior least-square stencils and CC-NA-CLIP clips about 3% of the interior nodes. On grids of both types, about 80% of the augmented stencils increase the stencil size just by one cell, about 20% by 2 cells, and less than 1% by more than 2 cells.

B. Gradient reconstruction accuracy

Figure 5 shows convergence of the \(L_\infty\) norms of relative gradient errors on grids of types \((III_p)\) and \((IV_p)\). Only errors computed with the CC-NA-CLIP scheme do not converge in grid refinement. Similar absence of convergence has been observed and reported previously\(^1\) for gradients reconstructed with the clipped CC-NA scheme within control-volume faces. All other methods provide first-order gradient approximations on grids of both types.

C. Convergence of truncation and discretization error

Numerical tests evaluating convergence of truncation and discretization errors are performed for the constant-coefficient convection equation (12). Figures 6 and 7 show convergence of the \(L_1\) norms of truncation and discretization errors, respectively.
Truncation errors of all the cell-centered schemes (except the CC-NA-CLIP scheme) converge with first order on grids of both types and truncation errors of the node-centered schemes converge with first order on triangular grids of type $(III_p)$; the corresponding discretization errors converge with second order. As predicted in Refs. [2, 5], truncation errors of node-centered schemes do not converge on mixed-element grids; the corresponding discretization errors converge with first order. The reason for this convergence degradation is the edge-based flux integration scheme, which is second-order accurate on simplex (triangular and tetrahedral) grids, but only first-order accurate on perturbed quadrilateral and general mixed-element grids. As shown in Ref. [5], with a more accurate face-based flux integration scheme, second-order accuracy is achieved with NC schemes on arbitrary grids. Although barely discernible, convergence of truncation and discretization errors of the CC-NA-CLIP scheme deteriorates on finer grids. Detailed tests performed on finer grids and reported in Section E show that truncation error convergence stagnates and discretization error convergence deteriorates to first order. Although not shown, convergence of the $L_\infty$ norms of the CC-NA-CLIP scheme show signs of deterioration on coarser grids. For other schemes, convergence slopes are the same for all norms and do not change on finer grids.

All second-order discretization error plots are close to each other indicating similar accuracy on grids with equivalent number of degrees of freedom. For reference, Figures 7(a) and 7(b) include the convergence plots of “ideal” discretization errors computed with the cell-centered exact-gradient (CC-EG) scheme that uses gradients evaluated at each cell from the manufactured solution. These plots represent the best-possible second-order convergence, which can be achieved on given grids. Close proximity of the actual and the ideal second-order discretization errors indicates that the accuracy is nearly optimal.

**D. Convergence of defect-correction iterations**

Convergence of DCI is studied on isotropic grids of types $(III_p)$ and $(IV_p)$ with $65^2$ nodes. Convergence rates are shown in Figure 8 for all schemes. As was mentioned above, the CC-SA and CC-NN schemes have small stencils and can be relaxed with full linearization of the target second-order operators.

The DCI method for all schemes converges fast with an average convergence rate per iteration better than 0.6. The convergence plots can be divided into three parts: initial convergence, transition, and asymptotic convergence. Initial convergence is typically fast for random initial solutions. The number of iterations within the transition region grows slightly on finer grids. Asymptotic convergence rates for all schemes are around 0.5 per iteration. Note, that on grids of type $(I)$, all studied discretization schemes correspond to the Fromm discretization of the convection equation. A detailed study of DCI for the Fromm discretization on Cartesian grids has been reported elsewhere. Note, also, that reported problems with stability of DCI for the WLSQ schemes and for the CC-NA scheme without clipping are not evident on these isotropic grids.
Figure 6. Convergence of $L_2$-norms of truncation errors on isotropic irregular grids. Manufactured solution is $U = -\cos(2\pi x - \pi y)$.

Figure 7. Convergence of $L_1$-norms of discretization errors on isotropic irregular grids. Manufactured solution is $U = -\cos(2\pi x - \pi y)$.

E. Effects of clipping

The tests reported in this section are performed for the CC-NA and CC-NA-CLIP schemes and demonstrate detrimental effects of clipping on convergence of gradient-reconstruction, truncation, and discretization errors in grid refinement. Irregular triangular grids of type ($III_p$) are considered. These grids are characterized by a higher percentage of clipped nodes; about 10% of the interior nodes are clipped. Figure 9(a) shows an example of a grid of type ($III_p$) with $17^2$ nodes; nodes where clipping occurs are circled.

Figure 9(b) shows that the gradients reconstructed by the CC-NA-CLIP scheme do not approximate the exact gradients. The CC-NA scheme provides a first-order accurate gradient reconstruction, which is sufficient for second-order discretization accuracy. Figures 9(c) and 9(d) exhibit convergence of the $L_1$ norms of truncation and discretization errors, respectively. The CC-NA scheme demonstrates first-order convergence of truncation errors and second-order convergence of discretization errors, which is consistent with results reported by Mitchell\textsuperscript{26} for Ringleb flow. Truncation errors are very similar on coarse grids, but start to diverge on finer grids. Truncation errors of the CC-NA
scheme demonstrate clear first-order convergence; truncation errors of the CC-NA-CLIP scheme converge slower on finer grids and eventually stagnate. The discretization error convergence of the CC-NA-CLIP scheme exhibits second order on the coarse grids, but then degrades to first order. Although not shown, the $L_\infty$ norm of discretization errors of the CC-NA-CLIP scheme shows degradation on coarser grids in grid refinement; asymptotically, $L_\infty$ norms of both node-averaging schemes converge with the same orders as the corresponding $L_1$ norms. Note that on grids with a small percentage of clipped nodes, convergence degradation becomes visible only on very fine grids. This may explain why such degradation has not been reported for practical computations.

VIII. Anisotropic irregular grids

A. Grid stretching

In this section, we study FVD schemes on stretched grids generated on rectangular domains. Figure 10 shows an example grid of type $(III_p)$ with the maximal aspect ratio $A = 10^3$. The manufactured solution is $U = \sin(\pi x + 2\pi y)$. A sequence of consistently refined stretched grids is generated on the rectangle $(x, y) \in [0, 1] \times [0, 0.5]$ in the following 3 steps.

1. A background regular rectangular grid with $N = (N_x + 1) \times (N_y + 1)$ nodes and the horizontal mesh spacing $h_x = \frac{1}{N_x}$ is stretched toward the horizontal line $y = 0.25$. The $y$-coordinates of the horizontal grid lines in the top half of the domain are defined as

   \[ y_{\frac{N_y}{2} + j} = 0.25; \quad y_j = y_{j-1} + \delta_y \beta^{j-1/2} \left( \frac{N_y}{2} + 1 \right), \quad j = \frac{N_y}{2} + 2, \ldots, N_y, N_y + 1. \]  

   Here $\delta_y = \frac{h_x}{A}$ is the minimal mesh spacing between the vertical lines; $A = 10^3$ is a fixed maximal aspect ratio; $\beta$ is a stretching factor, which is found from the condition $y_{N_y+1} = 0.5$. The stretching in the bottom half of the domain is defined analogously.

2. Irregularities are introduced by random shifts of interior nodes in the vertical and horizontal directions. The vertical shift is defined as $\Delta y_j = \frac{1}{4} \rho \min(h_y^{-1}, h_y^j)$, where $\rho$ is a random number between $-1$ and $1$, and $h_y^{-1}$ and $h_y^j$ are vertical mesh spacings on the background stretched mesh around the grid node. The horizontal shift is introduced analogously, $\Delta x_i = \frac{1}{4} \rho h_x$. With these random node perturbations, all perturbed quadrilateral cells are convex.

3. Each perturbed quadrilateral is randomly triangulated with one of the two diagonal choices; each choice occurs with a probability of one half.
B. Gradient reconstruction accuracy

A recent study\textsuperscript{27} assessed accuracy of gradient approximation on various irregular grids with high aspect ratio $A = h / h_y \gg 1$. The study indicates that for rectangular geometries and functions predominantly varying in the direction of small mesh spacing ($y$-direction), gradient reconstruction is accurate. For manufactured solutions significantly varying in the direction of larger mesh spacing ($x$-direction), the gradient reconstruction may produce extremely large $O(Ah_x)$ relative errors affecting the accuracy of the $y$-directional gradient component. Figure 11 shows examples of first-order accurate gradient approximations that exhibit large relative errors on high-aspect-ratio grids of type $(III)$. 

Figure 11 demonstrates the evaluation of gradient reconstruction accuracy performed with the methodology of downscaling described in detail elsewhere.\textsuperscript{2,5} The computational tests are performed on a sequence of downscaled narrow domains $L \times (L/A)$ centered at the focal point $(x, y) = (0.3, 0.5)$. The scale $L$ changes as $L = 2^{-n}$, $n = 0, \ldots, 8$ and the considered aspect ratios are $A = 10^6$ and $A = 10^3$; the latter corresponds to the highest aspect ratio observed at the central line of the stretched grid shown in Figure 10. On each domain, an independent high-aspect-ratio random grid of type $(III)$ with $9^2$ nodes is generated; the grid aspect ratio is fixed as $A$ on all scales. The gradient reconstruction accuracy

Figure 9 shows the accuracy of CC-NA schemes on isotropic irregular triangular grids of type $(III_p)$. Manufactured solution is $U = -\cos (2\pi x - \pi y)$. 

B. Gradient reconstruction accuracy

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was measured at the interior control volumes. Only weighted-least-square schemes, NC-WLSQ and CC-FA-WLSQ, provide accurate gradients independent of the aspect ratio $\mathcal{A}$, the relative errors of gradient reconstructions provided by other schemes are several orders of magnitude larger and directly proportional to $\mathcal{A}$. All methods converge with first order in grid refinement.

A summary of the results concerned with gradient accuracy on anisotropic grids is presented in Table 5. All considered gradient reconstruction methods may generate large relative errors on perturbed grids of types (III) – (IV)$. Only the NC-WLSQ scheme provides gradient reconstruction accuracy on all unperturbed grids. On perturbed grids, there are topologies, where all stencil points are essentially equidistant from the stencil center, and the WLSQ method is ineffective. Such situations occur more frequently for cell-centered schemes; all cell-centered schemes may generate large gradient errors even on unperturbed mixed-element grids of type (IV). The CC-NN, CC-NA, and CC-FA-unweighted methods may also have large relative errors on random triangular grids of types (III); the CC-FA-WLSQ method always provides accurate gradients on these grids.
for the CC-NA scheme. Convergence rates of DCI for stable schemes are similar to those observed on isotropic grids of types VII (subsection E), discretization-error convergence order for the CC-NA-CLIP scheme is expected to deteriorate of order accurate solutions have been previously reported of 

\[ 10^{-6}, 10^{-5}, 10^{-4}, 10^{-3} \]

Figure 12. Convergence of discretization errors for solution \( U = \sin (\pi x + 2\pi y) \) on irregular stretched grids.

A poor gradient reconstruction accuracy, however, does not necessarily imply large discretization error. Second-order accurate solutions have been previously reported on grids with large gradient reconstruction errors. Here, we observe similar results for cell-centered and node-centered FVD schemes for constant-coefficient convection. Convergence histories of the \( L_1 \) norms of discretization errors for the manufactured solution \( U = \sin (\pi x + 2\pi y) \) on a sequence of consistently refined stretched grids of types (III) and (IV) are shown in Figure 12. On grids of type (III), all discretization errors converge with second order. Note that, from the convergence results reported in Section VII (subsection E), discretization-error convergence order for the CC-NA-CLIP scheme is expected to deteriorate to first order on finer grids. Discretization errors of the NC-WLSQ scheme are not shown in Figure 12 because the NC-WLSQ scheme does not converge in DCI on grids of types (III) and (IV). The NC scheme converges with first order, as expected. Discretization errors of all cell-centered schemes converge with second order, close to each other and to the ideal discretization errors (CC-EG).

### Table 5. Relative error of gradient reconstruction on anisotropic irregular grids.

<table>
<thead>
<tr>
<th>Grid Types</th>
<th>(I)</th>
<th>(II)</th>
<th>(III)</th>
<th>(IV)</th>
<th>(Ip) – (IVp)</th>
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<td>( O(A h_x) )</td>
<td>( O(A h_x) )</td>
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</table>

C. Convergence of discretization errors

D. Convergence of defect-correction iterations

The DCI method applied to NC-WLSQ and CC-FA-WLSQ schemes diverges on perturbed stretched grids with triangular elements (types (IIp), (IIIp), and (IVp)); the method converges fast for all schemes on unperturbed grids of types (I) – (IV). Somewhat surprisingly, in rectangular geometry, no convergence problems have been detected for the CC-NA scheme. Convergence rates of DCI for stable schemes are similar to those observed on isotropic grids.
Figure 13 shows convergence histories on a 33 × 129 grids of types $(III_p)$ and $(IV_p)$. The asymptotic rates for all converging schemes are around 0.5 per iteration.

**IX. Grids with curvature and high aspect ratio**

In this section, we discuss accuracy of FVD schemes on grids with curvature and high aspect ratio. The grid nodes are generated from a cylindrical mapping, where $(r, \theta)$ denote polar coordinates with spacings of $h_r$ and $h_\theta$, respectively. The grid aspect ratio is defined as the ratio of mesh sizes in the circumferential and the radial directions, $A = R h_\theta / h_r$, where $R$ is the radius of curvature.

The curvature-induced mesh deformation is characterized by the parameter $\Gamma$:

$$\Gamma = \frac{R (1 - \cos(h_\theta))}{h_r} \approx \frac{R h_\theta^2}{2h_r} = A \frac{h_\theta}{2}.$$  (14)

The following assumptions are made about the range of parameters: $R \approx 1$, $A \gg 1$, and $\Gamma h_r \ll 1$, which implies that both $h_r$ and $h_\theta$ are small. For a given value of $A$, the parameter $\Gamma$ may vary: $\Gamma \ll 1$ indicates meshes that are locally (almost) not deformed. As a practical matter, grids with $\Gamma < 0.2$ can be considered as nominally Cartesian. In a mesh refinement that keeps $A$ fixed, $\Gamma = O(A h_\theta)$ asymptotes to zero. This property implies that on fine enough grids with fixed curvature and aspect ratio, the discretization error convergence is expected to be the same as on similar grids generated on rectangular domains with no curvature.

Our main interest is the discretization errors of solutions varying predominantly in the radial direction on grids with $\Gamma \gg 1$ corresponding to meshes with large curvature-induced deformation. Computations and analysis reported earlier\textsuperscript{25,28,29} conclude that the unweighted-least-square gradient approximation is zeroth-order accurate for such solutions on grids with high $\Gamma$. The manufactured solution, $U = \sin(5\pi r)$, is used in all tests in this section.

Four basic types of 2D grids are studied in the cylindrical geometry. In distinction from the computational grids used in the rectangular geometry, random node perturbation is not applied to high-$\Gamma$ grids because even small perturbations in the circumferential direction may lead to non-physical control volumes.

Computational grids used in the grid-refinement study are radially stretched grids with radial extent of $1 \leq r \leq 1.2$ and angular extent of $20^\circ$ with a fixed maximal aspect ratio $A \approx 1, 100$. The grids have four times more cells in the radial direction than in the circumferential direction. The maximal value of $\Gamma$ changes approximately from 24 to 1.5. The grid stretching ratio is changing as $\beta = 1.25, 1.11, 1.06, 1.03$, and 1.01. Representative grids of types $(III)$ and $(IV)$ are shown in Figure 14.
To improve the accuracy of gradient reconstruction, a general approximate mapping (AM) method has been introduced in Ref. [1]. The method constructs a local mapping based on the distance function that supplies the distance from a field point to designated boundaries and is readily available in practical codes. The AM method applies the least-square minimization in a local coordinate system, \((\xi, \eta)\), where \(\eta\) is the coordinate normal to the boundary and \(\xi\) is the coordinate parallel to the boundary (see sketch in Figure 15). The unit vector normal to the boundary, \(\hat{n}_0\), is constructed as

\[
\hat{n}_0 = \frac{(r_0 - r_0')}{|r_0 - r_0'|},
\]  

Figure 15. Sketch of coordinate system used in approximate mapping method.
where the position of the control-volume center is denoted \( \mathbf{r}_0 \) and the position of the closest point on the boundary is denoted \( \mathbf{r}_{0}^* \). The unit vector normal to \( \mathbf{n}_0 \) is denoted as \( \mathbf{t}_0 \). For constructing the least-square minimization at a control-volume with the center \( \mathbf{r}_0 \), the local coordinates of a stencil point \( \mathbf{r}_i \) are defined as

\[
\xi_i = (\mathbf{r}_i - \mathbf{r}_0) \cdot \mathbf{t}_0, \\
\eta_i = (s_i - s_0),
\]

where \( s_i \) denotes the distance function at location \( \mathbf{r}_i \). Thus the \( \eta \)-coordinate corresponds to the distance from the boundary and the \( \xi \)-coordinate is the projection onto the plane parallel to the surface at \( \mathbf{r}_0 \). The least-square minimization yields gradients in the \( (\xi, \eta) \) directions or, equivalently, through a coordinate rotation, in the \( (x, y) \) Cartesian directions.

To reconstruct an interior state for flux computation at a control-volume face location, \( \mathbf{r}_f \), the coordinates associated with the mapping at \( \mathbf{r}_0 \) are introduced, as

\[
\xi_f^0 = (\mathbf{r}_f - \mathbf{r}_0) \cdot \mathbf{t}_0, \\
\eta_f^0 = (s_f - s_0).
\]

The coordinate \( s_f \) should approximate the distance from the actual surface consistently. A possible approximation at the interface location is

\[
s_f = (s_0 + s_1)/2,
\]

where, for node-centered schemes, \( s_0 \) and \( s_1 \) correspond to the distance function of the two nodes defining the edge, and, for cell-centered schemes, \( s_0 \) and \( s_1 \) correspond to the distance function of the two cell centers adjacent to the face. The interior state is linearly reconstructed using the state at \( \mathbf{r}_0 \) and gradients in the \( (\xi, \eta) \) directions. Direct reconstruction using Cartesian coordinate gradients is also possible, yielding identical results for cell-centered schemes on grids constructed with advancing-layer techniques. The AM method is applied only to the cell-centered schemes in this paper and the reconstruction uses Cartesian coordinate gradients directly.

At boundaries for node-centered schemes, the normal computed from (15) must be replaced because \( \mathbf{r}_0 = \mathbf{r}_0^* \). One method is to replace (15) with an area-weighted sum of the normals at boundary cells adjacent to \( \mathbf{r}_0 \).

### B. Accuracy of gradient approximation

The accuracy of gradients reconstructed in the global Cartesian coordinate system on high-\( \Gamma \) grids of types \( (I) \) – \( (IV) \) is summarized in Table 6. Convergence of the maximum gradient errors is tabulated.

The NC-WLSQ scheme reconstructs accurate gradients on grids of all types. All other schemes show large \( O(1) \) errors on mixed-element grids of type \( (IV) \). On grids of type \( (III) \), the CC-FA-WLSQ also provides accuracy for gradient reconstruction. Schemes using unweighted least-square gradient reconstruction produce large gradient errors even on regular grids.

Gradient accuracy is dramatically improved with the AM method. Table 7 shows accuracy orders for gradients reconstructed with cell-centered least-square methods in the local coordinates. All tested schemes provide accurate gradients on grids of all types. We do not show results for the CC-NA scheme with approximate mapping in this paper.

For illustration, Figure 16 shows relative accuracy of gradients reconstructed on grids of type \( (IV) \). Note that the CC-NA scheme produces erratic convergence and large gradient errors. This behavior is explained by node-averaging degeneration on high-\( \Gamma \) mixed-element grids. On these grids, there are topologies where the node solution is averaged from four neighboring cells. The four cell centers involved in such averaging may be located on a straight line, thus leading to degeneration.

### C. Discretization error convergence

Convergence of \( L_1 \)-norms of discretization errors of FVD schemes with and without approximate mapping is shown in Figure 17. Discretization errors of the NC-WLSQ scheme in Figure 17(a) are shown only for grids of type \( (III) \) with relatively low \( \Gamma \); on higher-\( \Gamma \) grids, the NC-WLSQ solutions could not be obtained because DCI do not converge.
Table 6. High-Γ grids: relative errors of gradient reconstruction in global Cartesian coordinates. Manufactured solution is \( U = \sin(5\pi \gamma) \).

<table>
<thead>
<tr>
<th>Grid Types</th>
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<th>( (III) )</th>
<th>( (IV) )</th>
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<td>( O(h^3_a) )</td>
<td>( O(h^3_a) )</td>
<td>( O(h^3_a) )</td>
</tr>
<tr>
<td>CC-SA</td>
<td>( O(1) )</td>
<td>( O(1) )</td>
<td>( O(1) )</td>
<td>( O(1) )</td>
</tr>
<tr>
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<td>( O(1) )</td>
<td>( O(1) )</td>
<td>( O(1) )</td>
</tr>
<tr>
<td>CC-FA</td>
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<td>( O(1) )</td>
<td>( O(1) )</td>
<td>( O(1) )</td>
</tr>
<tr>
<td>CC-FA-WLSQ</td>
<td>( O(h^3_a) )</td>
<td>( O(h^3_a) )</td>
<td>( O(h^3_a) )</td>
<td>( O(1) )</td>
</tr>
<tr>
<td>CC-NA-CLIP</td>
<td>( O(h^3_a) )</td>
<td>( O(h^3_a) )</td>
<td>( O(1) )</td>
<td>( O(1) )</td>
</tr>
<tr>
<td>CC-NA</td>
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<td>( O(h^3_a) )</td>
<td>( O(1) )</td>
<td>( O(1) )</td>
</tr>
</tbody>
</table>

Table 7. High-Γ grids: relative errors of gradient reconstruction in local approximate-mapping coordinates. Manufactured solution is \( U = \sin(5\pi \gamma) \).

<table>
<thead>
<tr>
<th>Grid Types</th>
<th>( (I) )</th>
<th>( (II) )</th>
<th>( (III) )</th>
<th>( (IV) )</th>
</tr>
</thead>
<tbody>
<tr>
<td>CC-SA</td>
<td>( O(h^3_a) )</td>
<td>( O(h^3_a) )</td>
<td>( O(h^3_a) )</td>
<td>( O(h^3_a) )</td>
</tr>
<tr>
<td>CC-NN</td>
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<td>( O(h^3_a) )</td>
<td>( O(h^3_a) )</td>
<td>( O(h^3_a) )</td>
</tr>
<tr>
<td>CC-FA</td>
<td>( O(h^3_a) )</td>
<td>( O(h^3_a) )</td>
<td>( O(h^3_a) )</td>
<td>( O(h^3_a) )</td>
</tr>
<tr>
<td>CC-FA-WLSQ</td>
<td>( O(h^3_a) )</td>
<td>( O(h^3_a) )</td>
<td>( O(h^3_a) )</td>
<td>( O(h^3_a) )</td>
</tr>
</tbody>
</table>

Figure 16. Convergence of relative gradient errors on high-Γ stretched grids of type (IV) with maximum aspect ratio \( A = 1, 100 \). Manufactured solution is \( U = \sin(5\pi \gamma) \). The magnitude of the NC-WLSQ errors is comparable with the magnitudes of the errors of the mapped schemes.

With the exception of the CC-NA scheme on high-Γ grids of type (IV), all schemes show very similar discretization errors.

Large erratic discretization errors of the CC-NA scheme are caused by degeneration of the node-averaging stencil on mixed-element grids mentioned in the previous section. This explanation is supported by the evidence of accurate solutions obtained with the CC-NA scheme on low-Γ grids and on triangular grids of type (III), where such degeneration is impossible.

On grids of the same size, the discretization errors of schemes using the AM method show less variation. The level
Figure 17. Convergence of $L_1$-norms of discretization errors on high-$\Gamma$ stretched grids with maximum aspect ratio $\lambda = 1, 100$. Manufactured solution is $U = \sin(5\pi r)$.

of discretization errors obtained by the schemes with $O(1)$ error in the gradient reconstruction is not much different from the discretization error level obtained by the schemes with either the AM method or the exact gradient.

D. Convergence of defect-correction iterations

Convergence rates of DCI on irregular high-$\Gamma$ grids are shown in Figure 18. The DCI method diverges for the CC-NA scheme on grids of both types and, for the NC-WLSQ scheme, on grids of type (III); on grids of type (IV), the NC-WLSQ scheme slowly converges. Clipping was originally introduced to stabilize computations on irregular grids and DCI with the CC-NA-CLIP scheme converge for both grids considered.

For all schemes, except the CC-SA and CC-FA schemes, convergence rates of DCI are significantly slower than $O(1)$ on grids of type (IV). DCI with the CC-NN scheme is stable in all 2D computations shown here, although much slower than DCI with the CC-SA scheme. We have encountered several 3D computations in which DCI is unstable with the CC-NN scheme but stable with either CC-NA-CLIP or the CC-FA or CC-SA schemes. This observation prompted the development of the CC-SA scheme.
Figure 18. DCI: convergence of $L_1$-norms of residuals on high-$\Gamma$ stretched grids with maximum aspect ratio $A = 1, 100$.

X. Conclusions

Two node-centered and six cell-centered schemes have been compared for finite-volume discretization of a constant-coefficient convection equation as a model of the inviscid flow terms. The cell-centered nearest-neighbor (CC-NN) scheme has the lowest complexity; in particular, its stencil involves the least number of neighbors. A version of the scheme that involves smart augmentation of the least-square stencil (CC-SA) has only marginal complexity increase. All other schemes have larger complexity; the complexity of node-centered (NC) schemes are somewhat lower than complexity of cell-centered node-averaging (CC-NA) and full-augmentation (CC-FA) schemes. Defect-correction iterations (DCI) with a first-order driver are typically used to solve second-order finite-volume discretization (FVD) schemes. Convergence of DCI is an important consideration. Thus, we have assessed convergence of DCI for all of the schemes considered here. Note, however, because of their low complexity, the CC-NN and CC-SA schemes are promising as candidates to be iterated with full second-order linearization.

Comparisons of accuracy and convergence rates of DCI have been made for two classes of tests: the first class is representative of adaptive-grid simulations and involves irregular grids with discontinuous metrics; the second class is representative of high-Reynolds number turbulent flow simulations over a curved body. All tests have been performed on consistently refined grids for smooth manufactured solutions.

For the tests of the first class performed in rectangular geometries, the following observations have been made:

1. Discretization errors of five of the six cell-centered schemes converge with second order on all tested grids. The CC-NA scheme with clipping fails to approximate gradients and degrades solution accuracy to first order. As expected, the NC discretization errors converge with second order on triangular and regular quadrilateral grids and with first order on mixed-element (types $(IV)$ and $(IV_p)$) and perturbed quadrilateral (type $(I_p)$) grids. Second-order discretization errors are quantitatively similar on grids with the same number of degrees of freedom and closely approach “ideal” second-order errors on given grids exhibited by the cell-centered scheme with exact gradients (CC-EG).

2. All schemes may produce $O(A h_x)$ large relative errors in gradient reconstruction on perturbed grids of types $(I_p)$ – $(IV_p)$; here $A$ is the grid aspect ratio and $h_x$ is the larger mesh spacing.

3. The DCI method for the schemes with the weighted least-square gradient reconstruction (CC-FA-WLSQ and NC-WLSQ) diverges on perturbed stretched grids. DCI converge fast for all other schemes, including CC-NN and CC-SA; a typical asymptotic convergence rate is 0.5 per iteration.

4. As a recommendation for computations in geometries with no curvature, the CC-NN and CC-SA schemes offer the best options of the lowest complexity and second-order discretization errors.
The tests of the second class have been performed on consistently refined stretched grids generated around a curved body, typical of those generated by the method of advancing layers. The range of grid parameters has been chosen to enforce significant curvature-induced grid deformations, characterized by the parameter $\Gamma$. All tests have been performed for a manufactured solution smoothly varying in the radial direction.

1. With one notable exception, the magnitudes of discretization errors on high-$\Gamma$ grids are close to the magnitudes of the ideal errors obtained by the CC-EG scheme. The exception is the CC-NA scheme on mixed-element grids, explained by a degeneration of the node-averaging procedure.

2. With least-square minimizations performed in Cartesian coordinates, only the NC-WLSQ scheme provides accurate gradients on all grids; all other schemes generate $O(1)$ errors in gradient reconstruction for mixed-element grids of type ($IV$). Although demonstrated here only for cell-centered schemes, least-square minimizations performed in local approximate-mapping coordinates provide accurate gradients on all grids.

3. Convergence rates of DCI for the CC-SA and CC-FA schemes are fast across the grid types. The DCI convergence rates for other schemes are slower on mixed-element grids of type ($IV$). Consistent with previous observations, the DCI method diverges for the CC-NA scheme on grids of types ($III$) and ($IV$) and, for the NC-WLSQ scheme, the DCI method diverges on grids of type ($III$).

4. Among other schemes, the CC-SA scheme is the most attractive; it offers low complexity, accuracy comparable with that of the ideal CC-EG scheme, and fast convergence of DCI. However, the particular tests performed in the curved geometry do not discriminate between the different schemes sufficiently. The poor gradient reconstruction is not reflected in the magnitude of discretization errors as would be expected, for instance, for tests with a gradient source term. In such tests, least-square schemes with approximate mapping would be expected to be better.

References


