The paper discusses results obtained using a finite-volume method on 3-D unstructured meshes to simulate turbulent viscous flows over typical aerospace configurations. The numerical tool was developed by the CFD group at Instituto de Aeronáutica e Espaço (IAE) to aid the design of aerospace vehicles. One such aerospace configuration of interest to IAE is the first Brazilian Satellite Launch Vehicle (VLS). The VLS launcher is composed of a three-stage, Runge–Kutta time stepping scheme is used to perform the time marching of the flow equations. Spatial discretization of the Reynolds-averaged Navier–Stokes equations can be performed with second-order centered or upwind schemes. Automatic grid refinement routines are considered to adapt the original mesh. A sensor based on density gradients selects the volumes to be refined. The code is able to handle tetrahedra, hexahedra, wedges, and pyramids. A full multigrid scheme is available to accelerate convergence to steady state. Coarse grid levels are constructed through an agglomeration procedure. One- and two-equation turbulence models are implemented to include the turbulent effects into the numerical formulation. The mentioned features integrated in one single code allow the Brazilian aerospace program to simulate complex flow conditions for sounding rockets and satellite launch vehicles with accuracy and reasonable computational resources. Good agreement with theoretical or experimental results is obtained with the present numerical tool.

**I. Introduction**

The paper discusses results obtained with a finite-volume code that simulates viscous, turbulent flows for 3-D, adaptive, unstructured meshes. The implementation uses a cell-centered, face-based data structure. A fully explicit, second-order accurate, five-stage, Runge–Kutta time stepping scheme is used to perform the time marching of the flow equations. Spatial discretization of the Reynolds-averaged Navier–Stokes equations can be performed with second-order centered or upwind schemes. Automatic grid refinement routines are considered to adapt the original mesh. A sensor based on density gradients selects the volumes to be refined. The code is able to handle tetrahedra, hexahedra, wedges, and pyramids. A full multigrid scheme is available to accelerate convergence to steady state. Coarse grid levels are constructed through an agglomeration procedure. One- and two-equation turbulence models are implemented to include the turbulent effects into the numerical formulation. The mentioned features integrated in one single code allow the Brazilian aerospace program to simulate complex flow conditions for sounding rockets and satellite launch vehicles with accuracy and reasonable computational resources. Good agreement with theoretical or experimental results is obtained with the present numerical tool.

**Nomenclature**

\[ a = \text{speed of sound} \]
\[ C = \text{convective operator} \]
\[ C_p = \text{pressure coefficient} \]
\[ D = \text{artificial dissipation operator} \]
\[ d = \text{minimum distance to the wall} \]
\[ e = \text{total energy per unit volume} \]
\[ e_i = \text{internal energy} \]
\[ f = \text{source term of the multigrid method} \]
\[ k = \text{specific turbulent kinetic energy} \]
\[ p = \text{static pressure} \]
\[ P_x = \text{inviscid flux vector} \]
\[ P_y = \text{viscous flux vector} \]
\[ Q = \text{vector of conserved properties} \]
\[ q = \text{heat flux vector} \]
\[ \text{RHS} = \text{right-hand side operator} \]
\[ S = \text{absolute value of the mean strain-rate tensor} \]
\[ S = \text{area vector} \]
\[ S_{ij} = \text{mean strain-rate tensor component} \]
\[ T = \text{static temperature} \]
\[ u, v, w = \text{Cartesian velocity components} \]
\[ V = \text{viscous operator} \]
\[ v = \text{Cartesian velocity vector} \]
\[ x, y, z = \text{Cartesian coordinates} \]
\[ \alpha = \text{angle of attack} \]
\[ \alpha_1 \ldots \alpha_5 = \text{Runge–Kutta control parameters} \]
\[ \gamma = \text{ratio of specific heats} \]
\[ \Delta t = \text{time step} \]
\[ \kappa = \text{von Karman constant} \]
\[ \mu = \text{dynamic viscosity coefficient} \]
\[ \nu = \text{kinematic viscosity coefficient} \]
\[ \nu^* = \text{modified Spalart–Allmaras eddy-viscosity coefficient} \]
\[ \rho = \text{density} \]
\[ \tau = \text{viscous stress tensor} \]
\[ \Phi = \text{gradient ratio for limiter computation} \]
\[ \psi = \text{control volume limiter} \]
\[ \omega = \text{turbulent dissipation} \]
\[ \Omega = \text{absolute value of the mean rotation tensor} \]
\[ \Omega_{ij} = \text{mean rotation tensor component} \]

**Subscripts**

\[ f, k = \text{face index} \]
\[ i, m = \text{grid control volume indices} \]
\[ \ell = \text{laminar property} \]
\[ L, R = \text{interface left and right properties} \]
\[ t = \text{turbulent property} \]
\[ \infty = \text{freestream property} \]

**Superscripts**

\[ (m) = \text{current grid of the multigrid method} \]
\[ n = \text{time instant} \]
\[ * = \text{dimensional property} \]
central body and four strap-on boosters [1]. An illustrative sketch of
the vehicle is presented in Fig. 1. This is the first time that such
thorough analyses of the VLS aerodynamics at representative
transonic and supersonic flight conditions are performed using a
viscous, turbulent CFD tool.

The finite-volume computational code solves the compressible
Reynolds-averaged Navier–Stokes (RANS) equations. A fully
explicit, second-order accurate, five-stage, Runge–Kutta time
stepping scheme is used to perform the time march of the flow
equations. For flux calculations on the volume faces, a Jameson
centered scheme [2] plus explicitly added artificial dissipation terms
[3], or a Roe flux-difference splitting scheme [4] can be used. In the
latter case, second-order accuracy is achieved through a
scheme. An extension of the original multidimensional limiter
formulation of [5] is proposed to keep high accuracy at generic cell
types and to guarantee convergence. Furthermore, computationally
faster implementation of the Roe scheme and its associated limiters is
also proposed. The current methodology allows for large
computational resource savings while maintaining the expected level of
accuracy. The implementation uses a cell-centered, face
based data structure, and the code can use meshes with any
combination of tetrahedra, hexahedra, wedges, and pyramids.
Boundary conditions are set through the use of ghost cells attached to
the boundary faces.

A 3-D automatic mesh adaptation technology is also available.
The CFD group at IAE already has experience with 2-D mesh
refinement techniques [7–9]. Some of these ideas are currently extended for the 3-D case. Basically, the refinement technique
divides the original element in new ones by splitting its constitutive
faces. This splitting operation, however, allows for the presence of
hanging nodes, which clearly creates some additional complexity for
the coding. This additional complexity, nevertheless, is considered a
worthy tradeoff for increasing the overall mesh quality. The
implemented routines handle meshes composed of the previously
described cell element types.

Advanced eddy-viscosity turbulence models are available to
include the turbulence effects into the RANS equations. Viscous
simulations at high Reynolds numbers are typical for aerospace
applications, such as the ones of interest to IAE. Numerical
simulations of such flight conditions which do not consider
turbulence effects may have limited practical application. To obtain
useful viscous simulations results, the Spalart–Allmaras one
equation [10] and the SST two-equation [11] turbulence models are
chosen. These models are suitable for external aerodynamics
applications and they can predict flow separation with acceptable
levels of accuracy. The CFD group at IAE has some previous
experience with such closures for turbulent aerospace flow
simulations [12–14].

A full multigrid (FMG) scheme is included to achieve better
convergence rates for the simulations. To build the mesh sequence for
the multigrid procedure, an agglomeration scheme based on cell
or node seeds is used. The CFD group of IAE has experience with
such technique [12] in other 2-D numerical codes. A robust and
consistent method for 3-D turbulent flow simulations is proposed and
included into the present numerical formulation. This methodology
allows for successful simulations of high-Reynolds number
turbulent flows for highly stretched grids at very acceptable costs.

Extensive validation of this code has already been initiated, and
one is referred to [9,14,15] for a careful analysis of some validation
results. The mentioned features integrated in one single code allow
the Brazilian aerospace program to simulate complex flow
conditions for sounding rockets and satellite launch vehicles with
accuracy at reasonable computational resource expenses [9,14,15]
without resorting to the high costs associated with investments in
commercial flow solvers. In other words, this is a unique simulation
tool developed within the Brazilian aerospace research program,
which is comparable to few other commercial tool options.
Furthermore, an in-house development of such a tool also adds
flexibility to the assessment of the aerospace configurations of
interest to IAE. To demonstrate the mentioned capabilities in an
actual application context, simulation results obtained for the VLS
configuration, as well as other typical aerospace test cases using the
present code, are discussed in this paper. Turbulent viscous transonic
and supersonic flows are considered. The numerical results obtained
show good agreement with the experimental data and they represent
all the relevant aerodynamic features observed in experimental tests.

II. Theoretical Formulation

The objective of the CFD group at IAE is to develop the capability of
simulating 3-D, viscous turbulent flows over general launch
vehicle configurations. An account of the theoretical formulation of
the current 3-D unstructured grid flow solver is presented in the
forthcoming discussions.

A. RANS Equations

The flows of interest in the present context are modeled by the 3-D
compressible Reynolds-averaged Navier–Stokes (RANS) equations.
These equations can be written in dimensionless form, assuming a
perfect gas, as

$$\frac{\partial Q}{\partial t} + \nabla \cdot (P_e - P_s) = 0$$

with the following definitions

$$Q = \begin{bmatrix}
\rho \\
\rho u \\
\rho v \\
\rho w \\
e
\end{bmatrix}, \\
P_e = \begin{bmatrix}
\rho v \\
\rho u v + p_i \\
\rho u v + p_i \\
(\rho e + p) v \\
(e + p)v
\end{bmatrix}, \\
P_s = \begin{bmatrix}
0 \\
\tau_{ii} \\
\tau_{ii} \\
\tau_{ii} \\
\beta_{ii}
\end{bmatrix}$$

where \(i = \{x, y, z\}\) and \(\hat{i} = \{\hat{x}, \hat{y}, \hat{z}\}\) is the Cartesian coordinate unit vector.

The other relations can be given as

$$\tau_{ij} = (\mu_t + \mu_r) \left[ \frac{\partial u_i}{\partial x_j} + \frac{\partial u_j}{\partial x_i} - \frac{2}{3} \frac{\partial u_m}{\partial x_m} \delta_{ij} \right]$$

$$q_i = -\gamma \left( \frac{\mu_t}{P_e} + \frac{\mu_r}{P_r} \right) \frac{\partial (e)}{\partial x_i}$$

and \(\beta_{ij} = \tau_{ij} u_j - q_i\), where \(u_i\) is the Cartesian velocity component, \(x_i\)
the Cartesian coordinate, and \(\delta_{ij}\) is the Kronecker delta. In
the previous definitions, \(\mu_t\) is the molecular dynamic viscosity
coefficient, computed by the Sutherland’s law [14], and \(\mu_r\) is
the eddy-viscosity coefficient, computed by the chosen turbulence
model. The dimensionless pressure, \(p\), can be calculated from the
perfect gas equation of state as

$$p = (\gamma - 1) \left[ 1 - \frac{1}{2} \rho u^2 + v^2 + w^2 \right]$$

In this work, all properties are made dimensionless according to a
set of dimensional reference variables provided by the user. The
necessary dimensional reference variables are composed of a
reference length \(L_\text{ref}\), a reference speed \(V_\text{ref}\), a reference dynamic
viscosity coefficient \(\mu_\text{ref}\), a reference temperature \(T_\text{ref}\), and a
reference density, \(\rho_\text{ref}\). The dimensionless properties are defined
according to [16]. The user must also provide the gas properties,
namely, the constant of the gas \(R\), the specific heat at constant
volume \(C_v\), the specific heat at constant pressure \(C_p\); the
Prandtl number \(Pr\), and the turbulent Prandtl number \(Pr_T\). The gas \(R\), \(C_p\), and \(C_v\) properties are also made dimensionless with the provided

Fig. 1 VLS geometry overview.
reference variables. Furthermore, the Reynolds number is defined as

\[ Re = \frac{\rho U^2 L}{\mu} \]

One should observe that the Euler equations are obtained from the RANS equations as the Reynolds number approaches infinity.

B. Turbulence Modeling

The present work is mainly interested in high Reynolds number simulations of flows over complex aerodynamic configurations. Two turbulence closures are chosen in the present context, namely, the Spalart–Allmaras [10] (SA) one-equation model and the Menter SST [11] two-equation model. Both closures are particularly suited for aerodynamic flow simulations and separation prediction [11]. Furthermore, they are also less restrictive in relation to the grid refinement near the wall than other two-equation models such as the \( k-\varepsilon \) family of models [11].

Both models are solved according to the finite-volume approach. The convective term is discretized using a simplified first-order upwind scheme, and the diffusion term is discretized using a second-order centered scheme. The time march is performed using the implicit Euler scheme. One should observe that the use of an implicit scheme for the time march, in an unstructured mesh, leads to a sparse linear system. The solution for this system of equations is obtained using the biconjugate gradient method [17]. More implementation details on the time integration of the turbulence modeling equations can be found in [9].

Finally, because a measure of the turbulent kinetic energy is used in the SST model, that turbulence quantity should be included in the viscous terms of the RANS equations. Thus, some terms are redefined as

\[ \nu_{ij}^{new} = \nu_{ij} - \frac{2}{3} \sigma_{ij} \nu_{kl} \delta_{kl}, \quad \rho \nu_{i,ij}^{new} = \rho \nu_{i,j} + \left( \mu_{ij} + \sigma_i \mu_i \right) \frac{\partial k}{\partial x_j} \]

and the static pressure is redefined as \( \rho \nu_{ij}^{new} = p - (\gamma - 1) \rho \nu_{i} \). The inclusion of the turbulent kinetic energy as previously shown is unusual in the CFD community. However, these operations augment the physical accuracy of turbulent viscous flow simulations because some turbulent terms extracted from the Reynolds averaging procedure are brought back to the formulation not only via the eddy-viscosity coefficient. This is specially important when higher Mach number flow conditions are considered, in which the turbulent kinetic energy can be comparable to the local mean-flow kinetic energy [18].

III. Numerical Formulation

The forthcoming subsections describe the currently adopted finite-volume method and the temporal discretization of the governing equations.

A. Finite-Volume Discretization

The finite-volume method is used to obtain the solution of the RANS equations. The formulation of the method is obtained by an integration of the flow equations in a finite volume. The application of Gauss’ theorem for each finite volume yields

\[ \int_{V_j} \frac{\partial Q}{\partial t} \, dV + \int_{S_j} (P_e - P_i) \cdot dS = 0 \]

(6)

where the outward-oriented area vector is defined as \( S = \{ S_x, S_y, S_z \} \). The discrete value of the vector of conserved variables for the \( i \)th control volume is defined as the mean value of the conserved variables in the volume as

\[ Q_i = \frac{1}{V_i} \int_{V_i} Q \, dV_i \]

(7)

Hence, the final form of the finite-volume formulation for the RANS equations can be written for an elementary volume and assuming a stationary mesh as

\[ \frac{\partial Q}{\partial t} = -\frac{1}{V_i} \sum_{l=1}^{n_f} (P_e - P_i) \cdot S_k \]

(8)

where \( n_f \) is the number of faces which form the control volume. The code is able to handle grids composed of tetrahedra, hexahedra, wedges, pyramids, or a mix of these types of elements. The previous equation also indicates that the integral is discretized assuming the fluxes to be constant on the faces.

B. Time Integration

The integration in time of Eq. (8) is performed using a five-stage Runge–Kutta-type scheme [2,19] such as

\[ Q_i^{(0)} = Q_i^*, \quad Q_i^{(1)} = Q_i^{(0)} - \alpha_i \frac{\Delta t_i}{V_i} \text{RHS}^{(i-1)}, \quad Q_i^{(n)} = Q_i^{(n-1)} \]

(9)

where \( \ell = 1 \cdots 5 \), and the residue, RHS, is defined as

\[ \text{RHS}_i = C_i - V_i - D_i \]

(10)

Here, \( C_i \), \( V_i \), and \( D_i \) are, respectively, the convective operator, the viscous operator, and the artificial dissipation operator calculated for the \( i \)th control volume. These operators are calculated according to the spatial discretization scheme and they are detailed in the forthcoming sections. The \( \sigma_i \) coefficients are \( 1/4, 1/6, 3/8, 1/2, \) and \( 1 \) for \( \ell = 1, \cdots, 5 \), respectively. The viscous operator is calculated only on the first stage of the Runge–Kutta scheme to save computational resources.

The time step for each volume, \( \Delta t_i \), is calculated assuming a constant Courant–Friedrichs–Lewy (CFL) number throughout the computational domain. Hence,

\[ \Delta t_i = \text{CFL} \frac{\ell_i}{|V_i| + a_i} \]

(11)

where \( a_i \) is the speed of sound, \( |V_i| \) is the magnitude of the local flow velocity, and \( \ell_i \) is the characteristic length, in the \( i \)th cell. The characteristic length is set as the smallest distance between the control volume centroid and the centroids of each face that forms the cell.

C. Spatial Discretization

Both centered and upwind schemes are available in the present numerical method for the computation of the convective fluxes.

1. Centered Scheme

Centered schemes require the explicit addition of artificial dissipation terms to control nonlinear instabilities that may arise in the flow simulation. The centered spatial discretization of the convective fluxes, \( C_i \), in this scheme is proposed in [2]. The convective operator is calculated as the sum of the inviscid fluxes on the faces of the \( i \)th volume as

\[ C_i = \sum_{j=1}^{n_f} P_e(Q_j) \cdot S_j, \quad Q_k = \frac{1}{2}(Q_i + Q_m) \]

(12)

where \( Q_i \) and \( Q_m \) are the conserved properties in the \( i \)th and \( m \)th cells, respectively, that share the \( j \)th face.

The artificial dissipation operator is built by a blend of undivided Laplacian and biharmonic operators. In regions of high-pressure gradients, the biharmonic operator is turned off to avoid oscillations. In smooth regions, the undivided Laplacian operator is turned off to maintain second-order accuracy. A numerical pressure sensor is responsible for this switching between the operators. The expression for the artificial dissipation operator is given by
where $m$ represents the neighbor of the $i$th element, attached to the $k$th face, and $nb$ is the total number of neighbors of the $i$th control volume. Furthermore, the quantities in Eq. (13) are defined as

$$
\epsilon_2 = K_2 \max(v_j, v_m), \quad \epsilon_4 = \max(0, K_4 - \epsilon_2), \quad v_i = \frac{\sum_{m=1}^{nb} |p_m - p_i|}{\sum_{m=1}^{nb} |p_m + p_i|}
$$

with the Laplacian term approximated by an undivided Laplacian computation, such as

$$
\nabla^2 Q_i = \sum_{k=1}^{4ab} |Q_m - Q_4|
$$

In this work, $K_2$ and $K_4$ are assumed equal to $1/4$ and $3/256$, respectively. The $A_i$ matrix coefficient in Eq. (13) is replaced by a scalar coefficient $[3,20]$ such as

$$
A_i = \sum_{k=1}^{n} \left[ v_k \cdot S_k + a_k |S_k| \right]
$$

This formulation is constructed in an attempt to obtain steady-state solutions which are independent of the time step [21].

In the multistage Runge–Kutta time integration previously described, the artificial dissipation operator is calculated only on the first, third, and fifth stages. For the inviscid calculations, the artificial dissipation operator is calculated in the first and second stages only. This approach guarantees the accuracy for the numerical solution while reducing computational costs per iteration [2]. Furthermore, the artificial dissipation model has also been integrated into the multigrid framework. To achieve lower computational costs for the multigrid cycles, only the first-order artificial dissipation model is used in the coarser-mesh levels. This operation is achieved by not computing the biharmonic term in Eq. (13), and by setting $\epsilon_2 = \epsilon_4 + \epsilon_2$ in these coarser levels.

2. Upwind Scheme

The upwind discretization in the present context is performed according to the Roe flux-difference splitting scheme [4]. In this scheme, the inviscid numerical flux in the $k$th face can be written as

$$
P_{ek} = \frac{1}{2}(P_{e_k} + P_{e_k}) - \frac{1}{2} |\tilde{A}_k|(Q_R - Q_L)
$$

where $|\tilde{A}_k|$ is the Roe matrix associated with the $k$th face normal direction, defined as

$$
|\tilde{A}_k|(Q_R - Q_L) = \sum_{j=1}^{4} |\lambda_j| \delta_j r_j
$$

In this formulation, $|\lambda_j|$ represents the magnitude of the eigenvalues associated with the Euler equations, given as

$$
|\lambda| = \text{diag}(|v_n|, |v_n|, |v_n + a|, |v_n - a|)
$$

Similarly, $r_j$ represents the associated eigenvectors, given by

$$
r_1 = \left[ n_z, n_z u + n_z a, n_z v + n_z a, n_z w + n_z a, n_z (\Theta_1 + a(n_z v - n_z w)) \right]^T,
$$

$$
r_2 = \left[ n_z, n_z u - n_z a, n_z v, n_z w + n_z a, n_z (\Theta_1 + a(n_z w - n_z u)) \right]^T,
$$

$$
r_3 = \left[ n_z, n_z u + n_z a, n_z v - n_z a, n_z w, n_z (\Theta_1 + a(n_z u - n_z v)) \right]^T,
$$

$$
r_4 = \left[ 1 \quad u + n_z a \quad v + n_z a \quad w + n_z a \quad H + q_{e_a} a \right]^T,
$$

$$
r_5 = \left[ 1 \quad u - n_z a \quad v - n_z a \quad w - n_z a \quad H - q_{e_a} a \right]^T
$$

where $\Theta_1 = 0.5v \cdot v$. The $\delta_j$ terms represent the projections of the property jumps at the interface over the system eigenvectors, defined as the elements of

$$
\Delta = L[\Delta \rho \quad \Delta (\rho u) \quad \Delta (\rho u) \quad \Delta (\rho u) \quad \Delta \epsilon]^T
$$

where the left eigenvectors are the rows of the $L$ matrix, which are defined as

$$
L_1 = \left[ n_z + \frac{n_z u - n_z a}{u}, -\Theta_1 n_z, \Theta_2 n_z, \Theta_2 n_z, n_z, -\frac{n_z}{a}, \Theta_2 n_z, \Theta_2 n_z, -\Theta_2 n_z \right],
$$

$$
L_2 = \left[ n_z + \frac{n_z u - n_z a}{u}, -\Theta_1 n_z, \Theta_2 n_z, \Theta_2 n_z, \Theta_2 n_z, \Theta_2 n_z, n_z, -\Theta_1 n_z, \Theta_2 n_z, \Theta_2 n_z, -\Theta_2 n_z \right],
$$

$$
L_3 = \left[ n_z + \frac{n_z u - n_z a}{u}, -\Theta_1 n_z, \Theta_2 n_z, \Theta_2 n_z, \Theta_2 n_z, \Theta_2 n_z, \Theta_2 n_z, n_z, -\Theta_1 n_z, \Theta_2 n_z, \Theta_2 n_z, -\Theta_2 n_z \right],
$$

$$
L_4 = \left[ \Theta_1 \Theta_3 - \frac{n_z}{a}, -\Theta_3 u - \frac{n_z}{a}, -\Theta_3 v - \frac{n_z}{a}, -\Theta_3 w - \frac{n_z}{a}, -\Theta_3 \right],
$$

$$
L_5 = \left[ \Theta_1 \Theta_3 + \frac{n_z}{a}, -\Theta_3 u + \frac{n_z}{a}, \Theta_3 v + \frac{n_z}{a}, \Theta_3 w + \frac{n_z}{a}, \Theta_3 \right]
$$

(22)
with \( \Theta_2 = (\gamma - 1)/\sqrt{\gamma}, \Theta_3 = \Theta_2/2, \) and \( \Theta_4 = \Theta_1 \Theta_2. \) In the constant definitions, the \( k \)th subscript, which indicates a variable computed in the face, is eliminated to avoid overflowing the equations with symbols. Properties in the volume faces are computed using the Roe average procedure, as detailed in [4].

In the classical form in which the Roe scheme is presented, such as in Eq. (17), the underlining argument is the numerical flux concept, as also found in other upwind scheme examples [22,23]. Therefore, each time the numerical flux is built, the inherent numerical dissipation is also evaluated. In an explicit Runge–Kutta-type multistage scheme, this fact means that the Roe matrix defined in Eq. (18) is computed in all stages. The present authors rather interpret the Roe scheme as the sum of a centered convective flux, defined by

\[
C_i = \sum_{k=1}^{\infty} P_{ek} \cdot S_k,
\]

and an upwind-biased numerical dissipation contribution, that is given by

\[
D_i = \sum_{k=1}^{\infty} \frac{1}{2} |A_k (Q_{ Ri} - Q_{ Rj})| S_k.
\]

Therefore, the attractive, cheaper, alternate computation of the numerical dissipation in the multistage scheme, as already used for the switched artificial dissipation schemes, is also extended to the upwind flux computation. To the authors’ knowledge, this is a novel proposal and a unique feature of the present numerical tool.

To achieve second-order accuracy in space for the Roe scheme, lineal distributions of properties are assumed at each cell to compute the left and right states in the face. Such states are represented by the \( L \) and \( R \) subscripts, respectively, in the previous Roe definitions. The linear reconstruction of properties is achieved through a MUSCL [6] scheme, in which the property at the interface is obtained through a limited extrapolation using the cell properties and their gradients. To perform such reconstruction at any point inside the control volume, the following expression is used for a generic element, \( q \), of the conserved variable vector, \( \vec{Q} \), in Eq. (1):

\[
q(x, y, z) = q_i + \nu q \cdot \vec{r} \quad \text{(25)}
\]

where \((x, y, z)\) is a generic point in the \( i \)th cell; \( q_i \) is the discrete value of the generic property \( q \) in the \( i \)th cell, which is attributed to the cell centroid; \( \nu q \) is the gradient of property \( q \); and \( \vec{r} \) is the distance of the cell centroid to that generic point. Gradients are computed with the aid of the gradient theorem [24], in which derivatives are converted from the cell centroid to that generic point. Densities are computed through the gradient theorem [24], in which derivatives are converted into line integrals over the cell faces. In the present work, the control volume, \( \nu q \), in which to perform the gradient computation is chosen to be the \( i \)th cell itself [22]. The expressions for the reconstructed properties in the \( i \)th face can be written as

\[
(q_i)_{tk} = q_i + \psi_1 \nabla q_i \cdot \vec{r}_{ki}, \quad (q_i)_{hk} = q_m + \psi_m \nabla q_m \cdot \vec{r}_{km} \quad \text{(26)}
\]

where \( \nabla q_i \) and \( \nabla q_m \) are the gradients computed for the \( i \)th cell and its neighboring \( m \)th cell, respectively; \( \psi_1 \) and \( \psi_m \) represent the limiters in these cells; and \( \vec{r}_{ki} \) and \( \vec{r}_{km} \) are the distance vectors from the \( i \)th and \( m \)th cell centroids, respectively, to the \( k \)th face centroid.

The first-order Roe scheme can be readily obtained by setting the limiter value to zero in Eq. (26). This operation is equivalent to writing \( Q_{ki} = Q_i \) and \( Q_{km} = Q_m \) in the previous formulation. The integration of MUSCL-reconstructed schemes with the multigrid framework is simply accomplished by computing the second-order scheme in the finest grid level and the first-order one in the other coarser levels. This approach guarantees lower computational costs for the multigrid cycles while maintaining the adequate accuracy for the solution at the finest mesh level.

The limiter options that are available in the present context are the minmod, superbee, and van Albada limiters [25]. The respective 1-D definitions for these limiters are

\[
\psi(\Phi) = \begin{cases} 
\min(\Phi, 1), & \text{if } \Phi > 0, \\
\max(\Phi, 0), & \text{if } \Phi \leq 0
\end{cases}
\]

The total variation diminishing (TVD) [5,6] region is limited between the minmod and the superbee curves. In the previous equations, \( \Phi \) is defined as the ratio between the gradients of adjacent control volumes at the interface. One should observe that the minmod and superbee limiters require the evaluation of maximum and minimum functions, which characterizes these limiters as nondifferentiable. The van Albada limiter, on the other hand, is continuous. This aspect is discussed further in the current section.

In a similar sense as discussed for the Roe upwind scheme, the usual way of computing limiters is to perform such calculation every time the new numerical flux should be updated. The limiter computation work, though, is a very expensive task, amounting to more than half of an iteration computational effort, in the present context. Therefore, the idea of freezing the limiter along with the dissipation operator at some stages of the multistage time-stepping scheme is attractive in terms of computational resource savings. Hence, this approach is adopted in the current code framework.

The current extension of the 1-D limiters to the multidimensional case is based on the work of Barth and Jespersen [5]. Azevedo et al. [22] also present some insights into this effort to the 2-D case. The present work, however, presents a further extension of the methodology discussed in [22]. This extension is aimed at allowing for the user the choice of any desired limiter formulation, as well as to solve some Barth and Jespersen [5] limiter drawbacks. One of such disadvantages is that it is not a continuous limiter; other aspects are discussed later.

The difficulty in implementing a TVD method in a multidimensional unstructured scheme is related to how to define the gradient ratio, \( \Phi \). A generalization of \( \Phi \) for an unstructured grid is proposed in [15]. The proposed multidimensional gradient ratio for the \( f \)th face of the \( i \)th control volume is obtained as

\[
\Phi_{ki} = \begin{cases} 
\text{num}^+/\text{den}, & \text{if } \text{den} > 0, \\
\text{num}^-/\text{den}, & \text{if } \text{den} < 0, \\
1, & \text{if } \text{den} = 0
\end{cases}
\]

where

\[
\text{den} = (q_i)_{ki} - q_i, \quad \text{num}^+ = q_i^+ - q_i, \quad \text{num}^- = q_i^i - q_i
\]

Here, the extrapolated property in the face, \( (q_i)_{ki} \), is given by

\[
(q_i)_{ki} = q_i + \nu q_i \cdot \vec{r}_{ki}
\]

and the \( q_i^+ \) variables can be mathematically defined as

\[
q_i^+ = \max(q_i, q_j), \quad q_i^i = \min(q_i, q_j)
\]

The property in the \( f \)th face, \( q_f \), is the arithmetic average of the properties in the neighboring cells, as in Eq. (12), resulting in \( q_f = (q_i + q_{m+})/2 \), where the \( m \)th cell shares the \( f \)th face with the \( i \)th cell. The advantage of the gradient ratio definition in Eq. (28) is that it can be directly used in any other limiter definition, such as the ones presented in Eq. (27). It can also be used to recast the original Barth and Jespersen [5] limiter formulation with a slight modification though. More details of this formulation can be found in [15].

The complete definition for the current multidimensional limiter is finally presented. The following algorithm is implemented for the computation of the limiter in the \( i \)th control volume:

1) The computation of the limiter in the \( i \)th cell is initiated by collecting the minimum, \( q_i^- \), and the maximum, \( q_i^+ \), values for the generic \( q \) variable, in the sense of Eq. (31).

2) For each centroid of the \( k \)th face of the \( i \)th cell, the following steps are performed:

a) The property \( q_f = q(x_i, y_i, z_i) \) in the \( k \)th face centroid is extrapolated, as in Eq. (30).
b) The gradient ratio in the face, $\Phi_i$, necessary to compute the limiter, is obtained through Eq. (28).

c) A limiter value is computed at each face of the $i$th control volume.

3) The limiter value for the $i$th control volume is finally obtained as the minimum value of the limiters computed for the faces.

The nondifferentiable aspect of the minmod, superbee, and Barth and Jespersen [5] limiters poses some numerical difficulties in their use for practical numerical simulations. Their discontinuous formulation allows for limit cycles that hamper the convergence of upwind inviscid and viscous flow simulations to steady state [26]. One option to work around this problem is to freeze the limiter after some code iterations or residue drop, but this technique seems to not always work and to be highly problem dependent [26]. Such characteristics may also inhibit its application in actual production environment due to the need for user input in setting the limiter freezing operation for the simulation of interest. Another option is to use differentiable (or continuous) limiters instead of the ones which require maximum and minimum functions. Some examples can be found in [26], for instance. The limiter formulation in that work, however, seems to somehow pose a tradeoff between convergence and obtaining monotone (oscillation-free) steady-state solutions. The van Albada limiter, modified to include a smoothness augmentation threshold constant, is given by

$$\psi(\text{num}^t, \text{den}) = \frac{\text{num}^t(\text{num}^t + \text{den}) + \epsilon_{\text{LIM}}}{\text{num}^{t+2} + \text{den}^t + \epsilon_{\text{LIM}}} \quad (32)$$

where $\text{num}^t$ or $\text{num}^t$ are employed in the same sense specified in Eq. (28). Furthermore, $\epsilon_{\text{LIM}}$ is the constant limiter control, chosen as $\epsilon_{\text{LIM}} = 10^{-4}$ in the present work. This option seems to be appropriate for all aerospace cases considered by the present and other [27] development groups, always allowing machine-zero steady-state convergence for monotone numerical solutions.

3. Viscous Flux Computation

The viscous operator in the $i$th control volume is calculated as the sum of the viscous fluxes on the faces which constitute the volume

$$V_i = \sum_{k=1}^{nf} P_i(Q_k) \cdot S_k \quad (33)$$

In this case, both the conserved variable vector and its derivatives, on the face, are calculated as arithmetic averages between their corresponding values in the two volumes which contain the face. Derivatives of flow variables, for each control volume, are calculated in the standard finite-volume approach in which these derivatives are transformed, by the gradient theorem, into surface integrals around the control volume [24, 28].

D. Boundary Conditions

“Ghost” volumes are used to enforce the boundary conditions. The boundary conditions for external flows implemented in the 3-D finite-volume code are solid wall conditions for viscous and inviscid flows, nonreflecting farfield, inlet and outlet, symmetry, and extrapolation conditions. Detailed discussions of the boundary conditions for the RANS equations can be found in [9]. Furthermore, detailed discussions on the boundary conditions for the turbulence model equations are found in [14].

IV. Multigrid Technique

A technique that may allow an excellent convergence acceleration for numerical methods is the multigrid procedure [29]. The mathematical concept in which the multigrid technique is based consists basically in eliminating the low-frequency errors of the finest grid by solving the problem in coarser grids. This is based on the knowledge that the time-integration methods available today can only rapidly eliminate high-frequency errors of a computational grid [30], and that the high frequencies associated with the coarse grids are approximately of the same order of magnitude of the low frequencies associated with the finest mesh.

A. Multigrid Methodology

The multigrid algorithm chosen for the present work is of the full approximation storage (FAS) type, which is the recommended method for nonlinear problems [30]. This method is based on exchanging both solution and residue values between different grid levels. It also relies on a good time marching procedure to be effective, such as the current Runge–Kutta time-stepping scheme.

The current multigrid method has been successfully validated with the present 3-D unstructured computational code for inviscid to turbulent viscous simulations, as shown in [31]. To improve the multigrid algorithm as well as the computational method, the simulations start at the coarsest grid level. Some iterations with the Runge–Kutta scheme are performed at this grid, and a high-order interpolation is performed to the next finer grid. Some multigrid cycles are then performed to improve the solution at this grid. This procedure is successively repeated until the finest grid is reached, with a good initial guess to the solution. Multigrid cycles are then performed on the finest mesh until convergence is reached. This technique is usually referred to as a full multigrid (FMG) scheme.

For the problem being solved, written in an operatorlike form, $L^{(0)}q^{(0)} = f^{(0)}$, the algorithm for a “W” cycle of the multigrid solver works as follows:

1) Presmoothing: execute $n_t$ iterations of the time marching procedure in the finest mesh level, $L^{(0)}q^{(0)} = f^{(0)}$.

2) For grid levels $g = 0$ to $M - 1$:
   a) Residue computation: $r^{(0)} = f^{(0)} - L^{(g)}q^{(0)}$
   b) Residue restriction: $r^{(g+1)} = R(r^{(g)})$
   c) Solution restriction: $q^{(g+1)} = R(q^{(g)})$
   d) RHS computation: $f^{(g+1)} = r^{(g+1)} + L^{(g+1)}q^{(g+1)}$
   e) Coarse grid correction: execute $n_c$ iterations of the time marching procedure in level $g + 1$, $L^{(g+1)}q^{(g+1)} = f^{(g+1)}$.

3) Solve the problem for the coarsest grid level, $L^{(M)}q^{(M)} = f^{(M)}$, executing $n_M$ iterations of the time marching procedure.

4) For grid levels $g = M$ to $1$:
   a) Solution correction: $q^{(g-1)} = q^{(g-1)} + P(q^{(g)} - q^{(g-1)})$
   b) Postsmoothing: execute $n_p$ iterations of the time marching procedure in level $g - 1$, $L^{(g-1)}q^{(g-1)} = f^{(g-1)}$.

In this algorithm, $r$ represents the RHS operator at the nth time instant, defined as the negative of the RHS operator in Eq. (10). The multigrid RHS operator, $r$, is also augmented by a multigrid source term, $f$, which represents the residue information exchange between adjacent grid levels. Moreover, $q$ and $q'$ represent the conserved variables at the same time instant, $n$, obtained from two different ways, as described in the previous algorithm. The $g$ superscript denotes the operation for the gth grid level. The number of grid levels of this process is $M + 1$, where $M$ is associated with the coarsest grid level and 0 with the finest one.

In the previous algorithm, exchange operators are used for the connection between two consecutive grid levels. The restriction operator exchanges information from a grid level to the next coarser mesh. The prolongation operator interpolates from a grid level to the next finer one. The operator used in the present work for the conserved property restriction is the volume weighted average. The restricted conserved properties of a coarse mesh volume are equal to the sum of the conserved properties of all the fine mesh cells that form this coarse mesh volume, weighted by their volumes. Mathematically, this operation can be written as

$$q_i^{(g+1)} = \frac{1}{V} \sum_{n=1}^{ncv(i)} V_n q_n^{(g)} \quad (34)$$

where the property in the $i$th volume of the $(g + 1)$th coarser-mesh level is a summation of the property times the volume of the $ncv(i)$ volumes of the next finer $g$th mesh level that belong to the $i$th coarser-mesh cell. This nomenclature is detailed in Fig. 2. On the other hand, the restriction of the residuals is accomplished by simple addition of the finer-mesh residuals. Thus, the residue of a coarse mesh volume
is equal to the sum of the residuals of all the finer mesh volumes that are contained in this coarse mesh volume, such as

\[ r_{ij}^{(g+1)} = \sum_{h=1}^{n_{v(i)}} r_{ih}^{(g)} \]  

(35)

and the residue, \( r_i \), is defined in the previous multigrid algorithm. The restriction operator for the residuals is different from the restriction operator for the conserved properties because the residuals can be interpreted as surface integrals in finite-volume schemes.

The prolongation operator is only applied to conserved property transfers. For each \( f \)th face of the \( m \)th finer-grid cell, the volume-averaged corrections corresponding to the coarser-mesh neighbors, \( i \) and \( nb \), are multiplied by the \( f \)th face area and summed. The result is divided by the total area of the boundary surface of the \( m \)th control volume. This operation, in the \( (g-1) \)th mesh level, is mathematically described as

\[ P(\Delta q_m^{(g)}) = \frac{1}{V_m} \sum_{k=1}^{n_{f(i)}} \Delta q_i^{(g)} \frac{V_i + \Delta q_m^{(g)} V_{nb}}{V_i + V_{ab}} |S_i|, \]  

(36)

The \( m \) subscript represents the \( m \)th volume of the \( (g-1) \)th finer grid level, \( n_f(m) \) represents the total number of faces of this \( m \)th cell, and \( i \) and \( nb \) are the neighboring volumes of the coarser \( g \)th mesh level to the \( k \)th face. Property \( q \) is defined in the previously described multigrid algorithm. Furthermore, \( V \) is the cell volume and \( T \) is the total area of a given cell, given by

\[ T_i = \sum_{k=1}^{n_f(i)} |S_i| \]  

(37)

The nomenclature for this operation is detailed in Fig. 2. This procedure is implemented in a certain way that, if the \( k \)th face is internal to the coarser grid-level cell, as the \( k \)th face in Fig. 2, then automatically \( nb(k1) = i(k1) \).

B. Nonflux-Term Associated Issues

During the development of the multigrid scheme in the present context, problems in the calculation of nonflux terms, such as the artificial dissipation operator, have been observed. These problems are associated with the fact that two neighboring cells may share more than one face in a given agglomerated mesh level. To simplify the explanation of such issue, the following nomenclature is used hereafter: a face represents a real triangular or quadrilateral planar element, whereas an interface represents the complete surface that separates two neighboring volumes. These two definitions are presented in Fig. 3. For the finest grid level, a face and an interface are identical to each other, because two neighboring cells are exclusively separated by a unique face. For the agglomerated grid levels, this is not true anymore and an interface between two cells may contain more than one face, as depicted in Fig. 3. In this figure, the interface between the two given cells is formed by three faces.

For flux-associated terms, this fact does not represent any inconsistency because fluxes are computed through faces and they are multiplied by their surface area. This fact results in a summation of the total area of the given interface, yielding the correct flux computation for that interface. For nonflux terms, such as the artificial dissipation of the centered scheme, the computations do not take into account the face area and they must only be performed for each interface. In the present face-based code structure, one has information about faces and not about interfaces. Furthermore, storing interface information would excessively increase the memory usage of the multigrid solver. The approach in the present context to deal with this issue is to store a smaller array with the number of faces that compose a given interface between two cells. Each face receives this value, then, to be used whenever necessary. For instance, in Fig. 3, faces 1, 2, and 3 would receive the value 3 in the proper array position.

As an example, it is interesting to observe how the computation of the undivided Laplacian term, necessary for the artificial dissipation method, is performed. As already discussed, the Laplacian term is computed as

\[ \nabla^2 Q = \sum_{k=1}^{n_f} \{Q_k - Q_i\} \]  

(38)

where \( Q_i \) is the property in the \( i \)th volume and \( n_f \) is the number of faces of the same element. What happens in the agglomerated cell computation is that, at a given interface, say the interface in Fig. 3, the \( |Q_k - Q_i| \) difference is computed three times, one for each face of the interface. Nevertheless, one can clearly observe that this term is not associated with flux and, thus, this difference is added three times instead of once, which would be the correct procedure. Hence, the correct way to calculate this term would be to replace \( n_f \) by \( nn \), where \( nn \) would represent the number of neighboring cells, or interfaces, of the \( i \)th cell. Unfortunately, this information, along with other associated information, is not available in the code and it would be extremely costly to generate and store such data. The approach in the present context is to compute this term as indicated in Eq. (38) and divide the difference in the \( k \)th face by the number of faces that comprise the interface with the corresponding control volume. This would read, for the \( nm \)-th mesh level,

\[ \nabla^2 Q = \sum_{k=1}^{n_f} \{Q_k - Q_i\} \frac{\text{inface}(k,nm)}{|S_i|} \]  

(39)

where \( \text{inface}(k,nm) \) stores the number of similar faces for the \( k \)th face in the \( nm \)-th grid level and it is generated once in the code startup.

In the initial versions of the code which did not consider the preceding correction on the nonflux terms, strong unphysical property buildup has been observed in all simulations performed. Successful numerical results with the present multigrid method could...
only be obtained with the previously described nonflux correction. This methodology is sufficient to control the unphysical property buildup generated by the imbalance of the artificial dissipation terms due to the inconsistency of the implementation of nonflux related terms for an agglomerated mesh cell.

C. Agglomeration Technique
The coarse mesh levels used by the multigrid scheme are generated by an agglomeration technique. Similar technique has been successfully implemented into a 2-D version of the present 3-D numerical formulation [12]. In the current agglomeration algorithm, a seed volume is chosen in the fine mesh and all the volumes that have at least one node in common with this seed volume are grouped to form the coarse mesh cell. Another seed volume is then selected and the agglomeration procedure continues grouping all the fine mesh volumes. It should be noted that during the agglomeration procedure, only the volumes that are not already agglomerated may be grouped to form a coarse mesh cell. This is a necessary condition to guarantee that there is no volume overlapping in the coarse mesh.

To avoid large mesh growth between successive mesh levels, there is also an option to use seed nodes instead of seed volumes for agglomeration. For meshes composed of tetrahedra, for instance, the mesh growth based on the seed-volume option may result in coarsening ratios of about 20; that is, 20 volumes of the finer mesh level are agglomerated to form the coarser control volume. This behavior adds excessive interpolation errors for the present applications of interest. The seed-node option proved to be effective in avoiding such numerical problems, because less stringent coarsening ratios of about 8 can be obtained with its use.

Better coarse mesh quality can also be obtained if the selection of the seed volumes is not random. Therefore, a list containing all the fine mesh volumes is generated before the agglomeration procedure. For this work, the list is formed such that the first volumes are the volumes next to a solid surface, then other boundaries, and, afterward, the interior volumes. This approach is very simple to implement and adds very low additional computational cost. Although it does not necessarily provide the best agglomeration of the interior volumes, it results in good quality coarse mesh volumes close to the boundaries.

V. Adaptive Mesh Refinement
As is widely known, the quality of discrete numerical simulations is extremely dependent on the discretization mesh [32]. For high-quality numerical solutions, points concentrated in regions where the flow presents sudden variations are required. Such regions may be determined by both geometrical or aerodynamic factors. The concept of the adaptive mesh refinement is to leave to the numerical code, through flow indicatives, the responsibility of concentrating computational points by using automatic routines to modify the mesh. The adaptation can concentrate points in the identified regions by many forms. One way is to move points from flow zones that do not need them to the zones where they are needed. Another way is to create more points in these flow regions. In this work, the option of adding points is chosen because of its ease of implementation in an unstructured grid context.

The coarse mesh indicator sensor uses an undivided density gradient [7] normalized by the largest difference in density verified in the flow. This sensor can be written as

\[
(sensor)_i = \left[ \frac{|Vp|}{\rho_{\text{max}} - \rho_{\text{min},i}} \right] \tag{40}
\]

The sensor automatically refers to control volumes. If it is greater than a threshold value, the volume is indicated for refinement. The types of elements handled by this code are composed of either triangular or quadrilateral faces. To allow their refinement, each face type is split into similar elements, that is, triangular faces yields child triangular faces, and the same is true with quadrilateral ones, as shown in Fig. 4a. It is interesting to point out here that the neighboring unrefined cell of a cell that is marked to be refined a second time, is marked to be refined too. This approach is currently employed to allow a smooth decrease in element size throughout the adapted mesh, as shown in Fig. 4b. The control volume is internally divided compatibly with the face division. The interested reader can find details of the division of all types of elements in [9].

It is interesting to observe the presence of hanging nodes in the split element, as depicted in Fig. 4a. The treatment of hanging nodes is an important aspect, but with a simple implementation. The data structure of the code is face-based; therefore, it is irrelevant for the code if the node is hanging or not. The important aspect is that control volumes in this approach are no longer treated just as tetrahedra, hexahedra, wedges, or pyramids. In fact, each control volume may have the number of faces ranging from 4 (tetrahedron) to 24 (a hexahedra with all its faces divided). Moreover, one is only limited to 24 faces in the present case due to the size decrease criterion. Furthermore, because the data structure is face-based and each face stores the elements that contain it, this means that no memory is wasted. All loops are face-based and this approach makes the flux and dissipation term calculations independent of the element type.

VI. Verification Results
Aerodynamic flows over various aerospace configurations are simulated with the present computational tool. Results are compared to available theoretical or experimental data to assess the quality of the results that can be obtained with the numerical tool. A 1-D shock-tube problem is used to evaluate flux computation scheme results. A flat-plate flow is considered to address the turbulent flow simulation capability. The multigrid scheme is used in the simulations to accelerate convergence to steady state.

A. 1-D Shock Tube
Computations of 1-D shock-tube inviscid flow cases are considered. Numerical results are compared to the analytical solution for this problem. For the numerical simulations, an equivalent 3-D grid composed of a line of 500 hexahedra is used. The initial dimensionless density condition for the left half part of the shock tube is \(\rho_{\text{in},1} = 1\), whereas on the right half, \(\rho_{\text{in},2} = 20\). The reference conditions are taken in the initial state of the low-pressure side of the shock tube. Equal temperatures are assumed at both sides of the shock tube. Several other simulations with different density ratios have also been performed and the results are essentially similar to the ones presented in the forthcoming analyses. A constant dimensionless time step of \(\Delta t = 10^{-4}\) is used for this transient solution, and the forthcoming plots are taken at dimensionless time \(t = 0.1\).

The proposed Roe flux scheme implementation, which uses the concept of centered convective flux plus upwind artificial dissipation terms computed in alternate stages of the Runge–Kutta time marching procedure, is used. The van Albada limiter is chosen for the reconstruction process within the currently proposed multidimensional limiter implementation. The limiter computation is also performed at alternate stages of the Runge–Kutta time step. Numerical results for this formulation are compared to the ones obtained with the centered scheme.

Pressure and density distributions for the upwind and centered schemes are presented in Fig. 5. It can be observed that numerical solutions compare very well with the analytical one. No oscillations
near discontinuities can be found in the upwind numerical results, whereas oscillatory behavior is found for the centered scheme solution. Further detailed analyses of the numerical scheme and limiter formulations can be found in [15]. In that work, other numerical schemes and test cases are carefully studied, including thorough mesh refinement and topology dependency study of the current numerical scheme. Successful results are obtained for shocked and boundary layer flows with various combinations of numerical schemes and mesh refinement levels. It is also interesting to remark here that the proposed alternate computation of the Roe fluxes and limiters within the Runge–Kutta scheme guarantees exactly the same solution as if computing them at all stages. However, such procedure yields a numerical scheme which is 60% faster [15].

B. Flat-Plate Turbulent Flow

A zero-pressure-gradient flat-plate flow at \( Re = 7.62 \times 10^6 \) and low freestream Mach number \( M_\infty = 0.2 \) is considered. This is an important test case because a theoretical solution is provided for the turbulent boundary layer that builds up over the flat-plate surface [33], known as the log-law solution. Simulations with both SA and SST turbulence models are included to provide a comparison of the numerical results for both closures. The hexahedra mesh about the flat plate is clustered near the flat-plate surface to guarantee the condition of \( y^+ \approx 1 \) close to the solid boundary, which is required for integration of turbulence models to the wall. The mesh is also clustered near the flat-plate leading edge to account for the larger velocity gradients that are expected in that region. The resulting grid is composed of 60 cells within the boundary layer, and 80 cells streamwise the flat-plate length. Simulations are carried out with a standard multigrid setting of three grid levels, “V” cycles, and one Runge–Kutta smoother pass at each grid level.

Figure 6 shows the numerical boundary layers obtained for both turbulence models compared to the theoretical solution in [33] and to the experimental data in [34]. The definition of the dimensionless \( u^+ \) and \( y^+ \) variables in this figure can be found in [33]. One can clearly observe in Fig. 6 a striking coherence with the theoretical curve for both turbulence models. This is an indication of the correctness of the implementation of the turbulence models in the present code.

VII. VLS Results

Inviscid flows over the VLS in its first-stage (takeoff) and second-stage flight configurations are simulated for various Mach numbers using the adaptive mesh refinement routines. Turbulent viscous transonic and supersonic flows about the VLS second-stage flight configuration at various angles of attack are also considered. Because of reasonable numerical results and lower computational costs, the SA model is chosen for the forthcoming turbulent flow simulations. These flight conditions are chosen considering the dominant supersonic characteristic of the vehicle flight, and the numerical difficulties that arise at transonic simulations. Experimental data obtained through extensive high-Reynolds-number transonic and supersonic wind tunnel tests are available for this configuration [1]. Detailed description on the test setup and the experimental results for the VLS configuration can be found in [1]. Numerical results are compared to them such that the code effectiveness in the solution of realistic aerospace configuration flows can be assessed.

A. Inviscid Supersonic Flow over the First-Stage Flight Configuration

An inviscid flow at \( M_\infty = 2.0 \) and \( \alpha = 0 \) deg over the VLS is addressed. A view of the rocket nose and booster nose cap regions can be observed in Fig. 8. The mesh has approximately 106,000 nodes, and 581,000 tetrahedral volumes. In the geometry, the engine nozzles are not considered as this component is not adequately simulated with an inviscid formulation [35]. Furthermore, the currently available computational resources do not allow turbulent viscous flow simulations for the entire configuration. Despite the large number of control volumes, the mesh is not refined enough near the body, as can be observed in Fig. 8.

Figure 7 shows density contour results for the current flow case. Detached shock waves in front of the rocket nose and booster regions can be observed in this figure. The flow in the vehicle forebody presents a detached shock wave and a stagnation region in the rocket.

![Fig. 5 Property distributions along the shock tube obtained for different flux schemes.](image1)

![Fig. 6 Flat-plate turbulent boundary layer numerical, theoretical, and experimental results.](image2)
nose region. An expansion occurring along the conical part and over the corners of the payload fairing, and a shock wave at the end of the boattail, can also be observed. Another interesting feature in the flowfield is the high-pressure zone between the boosters, created by shock wave reflection mechanisms. The low-pressure region downstream this high-pressure zone is originated by the outward flow deflection caused by the presence of the high-pressure zone. It should be pointed out that the discussions here are presented in terms of pressure, as pressure and density are directly related by the state equation for ideal gases.

The computational mesh used in the previous results is less than adequately refined for such flight condition. The results show the need for mesh refinement in the booster nose cap and vehicle forebody regions, to adequately solve the complex shock wave structures in those regions. The adaptive mesh refinement technique is applied to the previous flow case to enhance the solution quality. Because of RAM memory restrictions, the authors limit the adaptation to only two refinement passes, although the routines are able to perform as many refinement passes as required. For the current case, the refinement threshold is set to 0.003. The first mesh refinement occurs at 5000 iterations in the original mesh, and the second one at 10,000 iterations in the refined mesh. The final mesh, after two refinement passes, has 362,740 nodes and 1,618,558 volumes, that is, about three times larger than the original coarser one. Detailed views of the adapted mesh over the vehicle forebody and booster nose cap regions are presented in Fig. 8. Clearly, the adaptation routines detect the presence of shock waves and expansions, and the mesh is consecutively refined mainly over these regions. The authors observe that this is a fully tetrahedral mesh. Comparisons of pressure coefficient distributions on the VLS pitching plane for the nonadapted and adapted meshes, are presented in Fig. 9. It can be observed in this figure that the adaptive refinement technique increases the solution quality. This can be seen as thinner shock waves and expansions on the vehicle forebody corners. However, two mesh refinement passes are still not enough to significantly enhance the solution near the booster nose cap zone. This part of the flow is characterized by the intense interactions between the detached shock waves from the boosters. The value of the pressure coefficient in this region is overpredicted by all numerical solutions and the authors believe that only a turbulent viscous simulation could obtain better results for that region.

B. Inviscid Transonic Flow over the First-Stage Flight Configuration

Density contours over the VLS at $M_\infty = 0.9$ and $\alpha = 0 \, \text{deg}$ are presented in Fig. 10. Interesting features of the flow are the shock waves over the payload fairing and boosters. Besides the payload-fairing shock wave, expansions in the corners of the payload fairing and a weak shock wave over the boattail can also be observed. One can also see that the flow rapidly accelerates between the boosters up to the formation of a shock wave. It should be noted that, for this transonic case, there is no high-pressure region downstream the shock wave as clearly marked as in the supersonic case. In that condition, the high-pressure region is formed by intense shock wave reflections in the region between the boosters. A previous work in the group with this test case using the same mesh, presented several problems related to solution oscillations in the domain [36]. A more refined mesh was necessary to yield results with similar quality to the present ones, which indicates the robustness of the current finite-volume code.

The adaptive mesh refinement routines are applied to the current flow case. Only one mesh refinement pass is performed, and the mesh refinement sensor threshold is set to 0.0007. The same grid used in the previous VLS first-stage supersonic simulations, as presented in Fig. 8, is also currently employed. As already discussed, this mesh is composed of 106,000 nodes and 581,000 tetrahedral cells. The adapted mesh after one refinement pass has 242,801 nodes and 768,177 elements. The mesh refinement is applied at 30,000 iterations in the original mesh. Detailed views of the adapted mesh over the vehicle forebody and booster nose cap regions are presented in Fig. 11. The mesh refinement routines detect the relevant regions of the flow, namely, stagnation in the central body and booster nose regions, the transonic shock wave over the payload-fairing.

![Fig. 7 Density contours over the VLS for inviscid flow simulations at $M_\infty = 2.0$ and zero angle of attack.](image)

![a) Forebody region – original mesh](image)

![b) Booster nose cap region – original mesh](image)

![c) Forebody region – refined mesh](image)

![d) Booster nose cap region – refined mesh](image)

![Fig. 8 Original and adapted meshes for inviscid flow simulations at $M_\infty = 2.0$ and zero angle of attack.](image)
component, and the expansion zone between the boosters. Mesh density is subsequently increased in the selected regions.

Comparisons of pressure coefficient distributions in the vehicle pitching plane for the nonadapted and adapted meshes is presented in Fig. 12. The improvement in the solution quality in this case is not as good as in the supersonic case. There are only minor improvements, such as a slightly better position of the shock wave in the payload-fairing cylinder. The inviscid solution presents higher pressure peaks than the experimental data. This behavior is already expected because large boundary layer interaction, mainly for the payload-fairing shock wave, is expected. This difference is diminished using turbulence models, as depicted in the next sections.

C. Turbulent Transonic Flow over the Second-Stage Flight Configuration

Turbulent viscous flows over the VLS at $M_\infty = 0.9$, $Re = 25 \times 10^6$ and zero angle of attack are considered. The mesh used in this case has 100,815 nodes and 89,280 hexahedra. The hexahedra are clustered near solid walls to guarantee $y + \approx 1$ for the interior volumes attached to the solid surface. Furthermore, about 40 hexahedra are placed within the boundary layer, to guarantee sufficient grid resolution in that region. Figure 13 presents Mach number contours over the vehicle forebody. This figure evidences the presence of the boundary layer over the vehicle. A stagnation point occurs in front of the vehicle. There is a supersonic expansion over the end of the conical forebody. This causes the formation of a supersonic region on the payload cylinder which is ended by a shock wave. Because of the boundary layer, this shock wave does not reach the body. The region over the end of the boattail presents very small velocities, but the boundary layer does not separate because of the turbulent characteristics of the flow. In fact, laminar simulations of this flow condition indicate boundary layer separation.

A comparison between pressure coefficient distributions over the VLS surface for inviscid and turbulent numerical solutions, as well as experimental data, is also shown in Fig. 13. The simulation with turbulent effects present more adequate results if compared to the Euler simulation, as already expected. A more consistent solution is obtained, which better captures the strong shock wave over the payload fairing and the weak one slightly downstream of the boattail expansion corner.

D. Turbulent Supersonic Flow over the Second-Stage Flight Configuration

A turbulent flow at $M_\infty = 2.0$, $Re = 30 \times 10^6$ and zero angle of attack over the VLS second-stage flight configuration is simulated. The mesh used in this case has 201,565 nodes and 188,480 hexahedra. The hexahedra are clustered near the solid surface to guarantee $y + \approx 1$, and approximately 40 cells are placed within the boundary layer, for consistent resolution of the turbulent effects in that region.

Figure 14 presents Mach number contours over the vehicle forebody. This flow case is characterized by a detached shock wave...
in the vehicle nose, an expansion on the first and second corners of the payload fairing, and a shock wave at the end of the boattail. A comparison between pressure coefficient distributions over the VLS surface for the turbulent numerical solution and the experimental data is also shown in Fig. 14. There are no relevant differences between the solutions for this case, as expected.

One of the objectives of the CFD group at IAE is to determine the stability derivatives of launch vehicles for many flow conditions. This is usually performed by the determination of the pressure coefficient on the vehicle wall for different angles of attack at the flow conditions of interest. Therefore, a simulation of the VLS flying at angle of attack different from zero is a relevant condition to be tested. As launch vehicles fly at very small angles of attack, the flow over the VLS second-stage flight configuration at $M_\infty = 2.0$, $\alpha = 2.0$ deg, and $Re = 30 \times 10^6$, is considered. The same computational mesh for the zero angle-of-attack case is used in this computation. Figure 15 presents Mach number contours over the VLS forebody in the pitching plane. The detached shock wave is no longer symmetrical. The shock wave on the windside is stronger than on the leeside because of the larger deflection of the flow in that region. Furthermore, the boundary layer on the leeside is thicker than on the windside. A comparison between numerical and experimental pressure coefficient distributions over the vehicle forebody in the pitch plane is also presented in Fig. 15. As one can observe in this figure, the numerical solution once again presents good agreement with the experimental data.

VIII. Conclusions

The paper presents results obtained with a unique, 3-D, finite-volume code developed within the Brazilian aerospace program to solve the RANS equations over typical aerospace configurations. The code uses a Runge–Kutta-type scheme to perform the time march of the flow equations. The code is designed to use unstructured meshes composed of any combination of tetrahedra, hexahedra, wedges, and pyramids. The agglomeration multigrid scheme
provides large convergence acceleration for the numerical simulations. In a general manner, numerical solutions of complicated flows such as transonic turbulent flows about a typical aerospace configurations can be obtained in half the time used by the single-grid simulation.

The fluxes on the volume faces are computed by either a centered scheme plus explicitly added artificial dissipation to control nonlinear instabilities, or a second-order flux-difference-splitting upwind scheme. Application of the upwind scheme in simulations of typical aerospace configurations shows that consistent results can be obtained with this formulation. The upwind scheme presents better results in solving laminar boundary layers due to its more physically coherent artificial dissipation formulation [15]. Therefore, it guarantees further accuracy for the numerical tool with which successful aerospace application results can already be obtained.

Turbulence effects are added to the formulation by eddy-viscosity-based one- and two-equation turbulence models. The Spalart–Allmaras one-equation and the SST two-equation turbulence closures are chosen to include the turbulence effects into the RANS equations. Comparison of numerical boundary layers over a zero-pressure gradient flat-plate flow with the corresponding theoretical log-law solution shows the level of accuracy that can be obtained with the present formulation. Furthermore, the code is also able to correctly solve for more complex flows, such as transonic or supersonic turbulent flows about typical aerospace configurations. Again, good approximation between experimental and numerical results are obtained for such cases.

The integration of all discussed features within a single computational code allow robust and accurate numerical solutions for complex turbulent transonic to supersonic flows with reasonable computational resource usage. This capability is demonstrated with simulations of various typical aerospace configurations. Therefore, the main contributions of the present work are a detailed description of an accurate and robust numerical tool for high-Reynolds-number aerospace applications, a novel formulation and implementation of a MUSCL-type upwind scheme, and a thorough analysis of the VLS launch vehicle aerodynamics. The quality of the numerical solutions that can be obtained with the present computational code are demonstrated. Successful results are presented with good comparison to experimental or theoretical data. VLS flow simulations provide consistent numerical results of acceptable quality for typical transonic and supersonic flight conditions. Simulation results show good approximation to wind tunnel data. The results presented here are a good indication of the capability of simulating turbulent flows about realistic aerospace configurations that has been developed by the CFD group at IAE. These results also show the maturity of the code to be inserted in the design phase of aerospace vehicles as a reliable and accurate numerical tool.

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