Multi-scale flow simulation of automotive catalytic converters
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HIGHLIGHTS
• We model the flow occurring at multiple scales inside catalytic converters.
• A subgrid model is proposed for the flow in the monolith channels.
• Adaptive Mesh Refinement techniques are optimized to capture the main flow features.
• The new model is validated against the experimental results reported by Benjamin.
• The new model allows for significant computational time savings compared to the full simulation.

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ABSTRACT
The flow distribution within the automotive catalytic converter is an important controlling factor on the overall conversion efficiency. Capturing the flow features minimizing the computational cost is the first important step towards the solution of the complex full engineering problem. In this work we present a novel approach that combines physical and numerical multi-resolution techniques in order to correctly capture the flow features inside an automotive catalytic converter. While Adaptive Mesh Refinement techniques are optimized in order to minimize the computational effort in the divergent region, a novel subgrid model is developed to describe the flow inside the catalytic substrate placed between the convergent and divergent regions. The proposed Adaptive Mesh Refinement methods are tested for two test cases representative of the flow features found in the divergent region of a catalytic converter. The performance of the new subgrid model is validated against the non-uniformity index and the radial velocity profile data obtained by Benjamin et al. (2002). The effective coupling of AMR techniques and the subgrid model significantly reduces the error of the numerical predictions to 5–15% in conditions where the full simulation of the problem is out of current computational capabilities.

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1. Introduction

Transportation is responsible for a large part of global emissions (Pachauri and Reisinger, 2007). This problematic has led to governments to establish very stringent conditions for the maximum emissions levels. Post-treatment systems need to be further developed in order to meet with these emissions requirements. A large part of the current studies is devoted to find efficient catalysts to improve the reaction efficiency, but one can also optimize the performance of these equipments by acting on the flow distribution inside the catalytic converter. A few studies (Agrawal et al., 2012; Bella et al., 1991; Guojian and Song, 2005; Karvounis and Assanis, 1993) have focused on the flow distribution effect on the conversion efficiency. However, the interaction between flow and conversion efficiency has not yet been understood completely.

In an ideal situation, the flow at the converter inlet is uniform. However, in practical cases, high Reynolds numbers, pulsating flow, abrupt expansion, and the impact of porous media lead to non-homogeneous and non-uniformity velocity profiles at the inlet converter. Because the velocity profile is non-uniform, we find different inlet velocities (hence different mass flow rates) in the substrate monolith channels. This results in premature degradation of the catalyst in areas of high flow rates and poor volume utilization of catalyst in areas of low flow rates, which turns into a decrease of the system's efficiency (Benjamin et al., 2002; Karvounis and Assanis, 1993). The flow inside the system under these physical and geometrical effects generates large range of scales on the flow in addition to the molecular scales inherent of the chemical reactions that occur inside the catalytic medium. Capturing all the physics and chemistry inside the system is out of reach for the available computational capabilities (Nien et al., 2013; Siemund et al., 1996).

The development of numerical tools and models is crucial in order to optimize the performance of catalytic converters...
Numerical Simulation (DNS) of turbulent AMR to reduce the computational limitations related to the Direct (Fuster et al., 2009a). In this work we investigate the capability of iteration (Fuster et al., 2009b, 2013) and other multiphase computational time savings in problems involving liquid atomization. These techniques have been already shown significant computational cost. A free CFD package (Gerris) is used as a platform to obtain reproducible statistics of the simulations usually have related long simulation runs in order to reach between accuracy and computational effort. Turbulent problems. This fact strengthens the compromise that one has to computationally expensive which impedes their application to real systems, different flow patterns are present due to changes in the cross section from the diffuser inlet to the nozzle outlet.

Fig. 1 depicts a classical example of the flow features found inside the system. In the inlet diffuser the flow expands and becomes turbulent (Neve, 1993; Shuja and Habib, 1996; Ubertini and Desideri, 2000). At the entrance of expansion a turbulent free shear layer develops. A main flow jet region appears close to the axis of symmetry whereas a recirculation flow region appears right after the beginning of the expansion. The intense recirculation induces high energy dissipation rates within the separated flow region (Forrester and Evans, 1997). The flow within the catalytic channel is significantly simpler. The flow is laminarized by viscous forces inside the narrow channels that induce a significant pressure drop across the channel compared to inlet diffuser and outlet nozzle. The characteristic Reynolds numbers in this region do not exceed 500 (Karvounis and Assanis, 1993). Finally, at the outlet nozzle, the section contracts and we find swirl flow (Forrester and Evans, 1997).

In the next subsections, the governing equations and models to solve these equations are discussed in detail.

2. Problem formulation

A typical and ideal automotive catalytic converter systems consist of an inlet pipe, a diffuser, a monolithic substrate, an outlet nozzle and an outlet pipe. The monolithic substrate is either ceramic or metallic and coated with aluminium washcoat which supports the noble metals (catalysts). The monolith comprises numerous parallel narrow channels (of the order of 1 mm) to increase the surface area where reactions occur. In after-treatment systems, different flow patterns are present due to changes in the cross section from the diffuser inlet to the nozzle outlet.

In this study, the general problem is presented first, then the derivation of a simplified model for the flow in catalytic substrate is developed. The accuracy of the model is demonstrated by comparing the results with the full simulation of the flow in this region. Then, we focus our efforts in deriving efficient Adaptive Mesh Refinement methods for the flow characteristics typically found in the divergent region (e.g. recirculation regions and shear layer). The accuracy and efficiency of the method is verified against related test cases with analytical solution. The new model is validated against the experimental data reported by Benjamin et al. (2002). Finally, we present a full numerical example of a typical automotive catalytic system.

3. Governing equations

3.1. Full model

The incompressible fluid assumption is widely used in the literature for the simulation of the flow in catalytic converters (Chakravarthy et al., 2003; Guojiang and Song, 2005; Holmgren et al., 1997; Lai et al., 1992). Treating the exhaust gas as incompressible fluid is a reasonable approximation since the Mach number is smaller than 0.05, acoustic waves have a negligible impact (Chakravarthy et al., 2003) and the variations in pressure are less than 10% of the absolute pressure (Holmgren et al., 1997).
For the sake of simplicity, we assume that the temperature change in the system is not significant and hence the fluid properties are constant. Under these assumptions and considering the gas as a Newtonian fluid, the governing equations for the flow are
\[ \nabla \cdot \mathbf{u} = 0, \]
\[ \rho \left( \frac{\partial \mathbf{u}}{\partial t} + \mathbf{u} \cdot \nabla \mathbf{u} \right) = -\nabla p + \mu \nabla^2 \mathbf{u} + S, \]  
where \( t \) is the time, \( \mathbf{u} \) is the velocity, \( \rho \) is the fluid density, \( p \) is the pressure, \( \mu \) is the viscosity and \( S \) is a momentum source term.

In addition, when chemical reactions inside the system need to be modeled, we have to add \( N \) transport equations, being \( N \) the number of components present in the system. For the i-th component we write
\[ \frac{\partial c_i}{\partial t} = \nabla \cdot (D \nabla c_i) - \nabla \cdot (u c_i) + R_i \]  
where \( c_i \) is the concentration of the i-th component, \( D \) is the diffusion coefficient and \( R_i \) is the reaction rate.

These equations can be solved by imposing proper boundary conditions. Typically we assume that the velocity at the inlet is known and we apply a classical outflow boundary condition at the outlet section (Dirichlet boundary condition for pressure and Neumann boundary condition for the normal velocity). The velocity at the solid walls is imposed equal to zero.

As stated above, the full solution of these equations is exceedingly expensive and we need to propose simplified solution strategies that we apply in regions where the flow features are already well captured by simple models. In particular, the flow inside the catalytic converter is a good candidate for such models. In the next subsection, we present the approach considered to model the flow in this region and how the model is coupled to the full numerical solution of the Navier–Stokes in the diffuser and convergent regions.

3.2. Subgrid models

3.2.1. Pressure drop model for monolithic channels

The flow inside monolithic channels is usually a fully developed laminar flow where the averaged velocity is kept constant by mass conservation. In these conditions, the pressure drop inside the channel is mainly induced by viscous forces and the flow is known to be well represented by the Hagen–Poiseuille pressure drop model (Heck et al., 2001)
\[ \Delta p = \frac{32 L}{\text{Re}_c d} \rho u^2, \]  
where \( \text{Re}_c \) is the Reynolds number inside the channel defined with the channel diameter \( d \) and \( L \) is the channel length.

As expected, the full simulation of pipe flow for the range of Reynolds numbers typically found inside the catalytic channels fits well the theoretical result (Fig. 2). The numerical data of a pipe flow is consistent with the Hagen–Poiseuille model and we can therefore replace the flow inside the channels by a pressure jump model as follows. The catalytic region is replaced by an infinitesimal thin interface along which only transversal velocity is allowed and where the pressure jump is imposed. In the Navier–Stokes equations, this pressure jump can be imposed by adding a source term of the type,
\[ \rho \left( \frac{\partial \mathbf{u}}{\partial t} + \mathbf{u} \cdot \nabla \mathbf{u} \right) = -\nabla p + \mu \nabla^2 \mathbf{u} + C(u)\delta_i \mathbf{n}, \]  
where \( \delta_i \) is a Dirac delta function used to denote that the pressure jump is applied at the interface that represents now the catalytic converter (see Fig. 3), \( \mathbf{n} \) is the normal to this interface and \( C(u) \) is the function that imposes the desired pressure jump as a function of the velocity,
\[ C(u) = \frac{32 L}{\text{Re}_c d} \rho \left( \frac{u}{A_0} \right)^2, \]  
where \( A_0 \) represents the open frontal area of the monolith substrate.

We remark that the subgrid model is naturally coupled with the full solution of the Navier–Stokes equations in the divergent and convergent region, where the source term is set to zero. The validation of the model is discussed in Section 5.

3.2.2. Heterogeneous reaction model inside the catalytic converter

In the simulations included in this work we focus our analysis on the dynamics of the flow, neglecting the reaction mechanisms taking place inside the catalyst monolith. However, we note that it is possible to develop the model further to obtain an approximate prediction of the reaction rates. Replacing the catalytic converter by an infinitely thin interface implies that one needs to model also the global reaction rate that occurs inside the catalytic converter by a simplified source term that we need to plug into Eq. (3).

The calculation of a proper expression for the source term is specific to the problem considered and can be very involved for a general case. However, in a first approximation, if we assume that the influence of homogeneous reactions is negligible compared to heterogeneous reactions, it is possible to find solutions for limiting regimes. The controlling factor of the reaction regime is the temperature. When the catalyst is not sufficiently warm, the reaction taking place on the solid walls is slow and the conversion
4. Numerical method

To solve for the system of equations given by Eqs. (1) and (2) we use the Gerris Flow solver (Popinet, 2009). This solver can be easily adapted to source terms of the form proposed in the previous section due to the similarities found with the pressure jump across gas/liquid interfaces when surface tension effects are present.

Another interesting characteristic of the solver is the capability to perform Adaptive Mesh Refinement (AMR) using quadtree (octree in 3D) meshes. By using AMR in the convergent and divergent regions we expect a significant gain on the computational time with respect to solvers with uniform grids. In the next subsection, we give further details about the use of AMR for the specific problem of catalytic converters as well as the two test cases used for the optimization of the mesh refinement strategies.

4.1. Multi-resolution AMR techniques

In this work we decided to use a Hessian error estimator based on the h-refinement algorithm. These methods basically consist in trying to obtain an estimation of the error contained in the numerical solution by subtracting the numerical solution obtained at two different resolution levels. Octree meshes are suitable for h-refinement methods because by construction it is simple and computationally efficient to travel up and down through the octree structure. Given a leaf cell with a level of refinement \( L \), one can express the solution of second order accuracy around the cell centered coordinate \( \mathbf{x}_c \) using the Taylor expansion as

\[
\hat{f}(\mathbf{x}) = f(\mathbf{x}_c) + (\mathbf{x} - \mathbf{x}_c) \cdot \nabla f(\mathbf{x}_c) + \frac{1}{2} (\mathbf{x} - \mathbf{x}_c)^2 \nabla^2 f(\mathbf{x}_c) + O(\| \mathbf{x} - \mathbf{x}_c \|^3),
\]

where we use an overbar to denote the discretized quantities. Typically, a-posteriori error estimation methods try to get an estimate of \( O(\| \mathbf{x} - \mathbf{x}_c \|^3) \) for every cell in the computational domain. For regular octree meshes of size \( h \), the maximum of the error scales as \( O(h^3) \). This error can be estimated using the Taylor expansion for the parent cell at the \( L-1 \) level and subtracting it from the Taylor expansions at \( L \) level at the leaf cell center. Thus one gets

\[
0 = \hat{f}(\mathbf{x}_c) + (\mathbf{x}_c - \mathbf{x}_c^{-1}) \cdot (\nabla \hat{f}(\mathbf{x}_c^{-1}) - \nabla f(\mathbf{x}_c^{-1})) + O(\| \mathbf{x}_c - \mathbf{x}_c^{-1} \|^3).
\]

Using \( \hat{f}^{-1}(\mathbf{x}_c) \) to denote the linear interpolation of the solution at the \( L-1 \) level at a given location and using that, for octree structures, \( \mathbf{x}_c - \mathbf{x}_c^{-1} = h/2 \) is always satisfied, we obtain the following estimation of the discretization error:

\[
O\left(\frac{h^2}{4}\right) \approx \| \hat{f}^{-1}(\mathbf{x}_c) - f(\mathbf{x}_c) \|.
\]

The error above can be interpreted as a measure of the error at the \( L-1 \) which can be corrected extrapolating the error at the \( L \) level taking into account the spatial order of the method used, \( \alpha \), so that,

\[
\epsilon = \| \hat{f} - f \|_{L_0} \approx \| \hat{f}^{-1} - f \|_{L_0}^{\alpha 2}.
\]

In this paper, we denote with \( \eta_{\text{hes} - L_0} \) the error estimation measured in the \( L_0 \) norm using this Hessian method approach.

\[
\eta_{\text{hes} - L_0} = \| \epsilon \|_{L_0}, \quad \eta_{\text{hes} - L_0} := \| \hat{f} - f \|_{L_0}
\]

For the simulations shown in this work we choose the \( L_1 \) norm. The question about which quantity \( f \) provides optimal results remains open. Typically, previous works use the error on the primitive variables (e.g. velocity) to define criteria to adapt the grid, but other choices are possible. For instance, in this work we have chosen the error in the vorticity field \( \omega = \partial_x u - \partial_y v \) as a criterion to adapt the grid. For two-dimensional simulations, vorticity is a conserved scalar quantity. To write the Navier–Stokes equations in terms of the equivalent vorticity equation has been shown some desirable numerical properties in terms of accuracy and speed (Davies and Carpenter, 2001). Recent formulations (Oshanski and Rebholz, 2010) have proposed the use of Helicity, which is the corresponding invariant of Euler’s equations in three dimensions, to derive efficient numerical schemes especially designed to capture turbulent structures. These works reveal that the use of vorticity (or helicity) is usually advisable when one wants to capture turbulent structures. Thus, we propose here the use of the residual of the vorticity field as a good candidate to minimize the numerical error introduced when discretizing the vortices appearing in the divergent region.

4.2. Test cases for adaptive mesh refinement

As mentioned previously, the two main flow patterns found in the expansion region of catalytic converters are a shear layer and a recirculation region. In order to validate the accuracy and the efficiency of the proposed AMR method for these flow patterns, we decide to use the following tests. The first test is the measure of the energy dissipation by a Lamb–Oseen vortex. This vortex model is representative of the vortices existing in the recirculation region in the inlet diffuser. The second test is the growth of random noise perturbations in a shear velocity region. The AMR efficiency of this example is also relevant for catalytic converters due to the shear layers generated in the abrupt expansion in the diffuser region.

4.2.1. Lamb–Oseen vortex

This example represents a solution to the two-dimensional viscous Navier–Stokes equations where axial and radial velocities are zero. Setting the initial vorticity field with a known circulation \( \Gamma \) to \( \omega(x, 0) = \Gamma \delta(x) \delta(y) \) it is possible to obtain the analytical solution of the velocity temporal
evolution as (Meunier and Villermaux, 2003)
\[ u_\theta = \frac{\Gamma}{2\pi r} \left[ 1 - \exp\left( -\frac{r^2}{4\nu t} \right) \right]. \]  
(12)

where \( \nu \) is the kinematic viscosity and \( r \) and \( \theta \) are respectively the radial and the azimuthal coordinates. A characteristic length for this problem can be obtained by setting the radial distance at which the velocity norm is maximal,
\[ l_c = \frac{2.2418}{\nu t_0}. \]  
(13)

For the simulations contained in this section we choose \( \nu t_0 = 0.5 \) in order to define a characteristic length different from zero. In addition, we set the circulation equal to \( \Gamma = 1 \) in a square domain with nondimensional length \( L/l_c = 600 \) where we impose Neumann boundary conditions for the velocity at all boundaries.

The initial velocity field is initialized according to Eq. (12). Fig. 4 represents the theoretical azimuthal velocity profiles as a function of the radius for different times. We note that the theoretical solution extends to infinity. This fact introduces a certain error at the domain boundaries when setting the domain size to a finite distance. This error cannot be attributed to the discretization method and therefore, cannot be captured by the error estimator. To solve this problem, we decide to measure the efficiency of the error estimators in a circular region of radius \( R_l/l_c = 100 \).

Fig. 5 depicts a classical velocity field and the corresponding grid distribution for a given example. The grid is significantly refined at the domain center where the velocity variations are important. Near the boundaries, the mesh is coarsened which significantly decreases the number of unknowns to be solved in the discrete form of the equations.

Fig. 6 depicts the convergence curves obtained for a fixed grid and the AMR criteria, where an effective grid size is defined as \( \Delta x = L/\sqrt{N_{cell}}, \ N_{cell} \) being the total number of cells in the domain. The convergence order for a fixed mesh is lower than two for coarse grids given that at very low resolutions, the discrete solution becomes a step function for which the maximum convergence order achievable is one. For sufficiently high resolution, the grid captures well the flow features at the vortex center and we observe the second order convergence expected from the second order discretization methods implemented in Gerris. When applying AMR, we significantly reduce the numerical error for a constant resolution. Although the convergence order is similar to that of uniform grids, AMR starts showing a second order convergence for lower resolutions than a fixed discretization, which turns into smaller discretization errors for a fixed number of cells. It is also worthy noting that in the low resolution region, where the convergence order is lower than two, the global accuracy for AMR simulations is significantly improved compared to a uniform grid for a given number of cells (Fig. 6(b)). The main reason for that is that the computational effort is mainly focused at the vortex center, where the solution variation is abrupt, whereas the solution is significantly coarsened close to the domain boundaries.

For this particular example, where the analytical solution is known, it is also possible to investigate the local performance of the proposed error estimator. We note that local analyses are more challenging than global error estimations given that in addition to the total error we want to know how this error is distributed in the computational domain. In order to measure the local efficiency of the criterion to capture the real error, we measure the coefficient of determination, \( R^2 \), obtained from the correlation between the predicted error \( \epsilon_{\text{estimated}} \) with respect to the true error \( \epsilon_{\text{true}} \), which is defined as a difference between the numerical and analytical solution. Fig. 7(a) shows that the correlation between the error estimated and the true error for all tested resolutions is linear. In addition, the coefficient of determination is close to the ideal value (\( R^2 = 1 \)). Fig. 7(b) shows that the proposed AMR method correctly quantifies the amount of error introduced by the numerical solution irrespective of the grid resolution. We therefore conclude that the proposed AMR method exhibits a good performance in order to minimize the total error in Lamb–Oseen like structures, giving also a good estimation of the local error distribution.

4.2.2. Growth of random noise perturbations in a shear layer

In this numerical test we investigate the capability of AMR to correctly capture the growth of small perturbations by a Kelvin–Helmholtz instability (KHI). KHI is defined as flow instability consequence of velocity difference between two parallel
A measure of the perturbation growth can be obtained by measuring the maximum amplitude of the vertical velocity $V$ in time. In particular, Fig. 8(b) depicts the temporal evolution of the $V$ component of the velocity integrated across the entire domain. After an initial transient state, the instability develops and generates well resolved structures that grow in time exponentially (Fig. 8(c)–(e)). The numerical growth-rate is obtained by fitting the perturbation growth in a time interval $t \Delta U / \delta_c = [30 : 50]$. The error is then computed by comparing the numerical and theoretical growthrate.

Fig. 9 depicts an example of the grid distribution typically generated. The grid is significantly refined at the velocity shear region where the velocity variations are important and it is coarsened gradually as variations become less important. Fig. 10 shows the error in the growth-rate obtained for a uniform mesh and a non-uniform mesh adapted according to the Hessian error estimator in the vorticity field. As in the previous test case, the AMR grid provides more accurate than a fixed mesh for a given resolution (Fig. 10). AMR is able to start displaying second order convergence for lower resolutions given that it is able to concentrate the grid cells in those regions where the error is large. For coarse grids the convergence order is significantly degraded but in any case the AMR provides significantly more accurate solutions than a fixed grid for a fixed number of cells (Fig. 10(b)).

5. Validation of the subgrid pressure jump model

In this section we concentrate on the validation of the pressure jump model using the experiments presented by Benjamin (2003) and Benjamin et al. (2002), where only the expansion stage and
the monolith substrate region are considered. These works were devoted to examine the flow distribution in automotive catalyst systems and to provide experimental data for verification of computational fluid dynamic simulations. The experimental set-up consists of a diffuser and a substrate. The diffuser is axially symmetric with a total angle 60° and length 61.5 mm. The inlet pipe diameter on which Reynolds number are based is 48 mm. The velocity profiles are measured 45 mm upstream of the diffuser throat. A tube is used to hold flow straighteners to achieve uniform flow. Two ceramic substrates of different lengths are used (152 mm and 102 mm respectively). Both substrates have a nominal cell density of 400 cspi made of square channels of 1 mm. The diameter of both substrates is 118 mm. Physical and geometrical properties of the experimental set-up are indicated in Table 2.

During the numerical investigations, we impose a uniform velocity profile at the inlet pipe. The walls of the system are treated as impermeable solid walls with no-slip boundary condition. For the outlet section, an outflow boundary condition is applied.

To evaluate the effect of geometry and mass flow rate on the flow distribution a non-uniformity index \( \sigma_v \) is defined using the variance of the velocity profile across a transversal section with respect to the averaged uniform velocity profile

\[
\sigma_v = \frac{1}{m} \int_A \left[ U - U_e \right]^2 \, dA. \tag{17}
\]

The non-uniformity index over the cross section of the substrate, \( \psi \) is defined as

\[
\psi = \frac{\sigma_v}{U_e} \times 100. \tag{18}
\]
This number increases as the flow becomes less uniformly distributed. For a perfectly distributed flow this number is zero.

Fig. 11 compares the vorticity fields obtained at various times. The flow patterns observed in the divergent region are similar to those observed by Benjamin (2003). The model is able to act as a porous wall that induces strong recirculation in the expansion region without the necessity to explicitly simulate the flow inside the monolith channels.

To validate the developed numerical tool we measure the non-uniformity index $\psi$ as a function of the Reynolds number for the two monolith substrates tested in Benjamin et al. (2002). We start showing the convergence of the results for the monolith of length 152 mm for $Re=10,000$. In this case, the model converges to a value relatively close to the experimental observations for all Reynolds numbers tested here. These results show that the AMR method allows us to obtain relatively accurate results with approximately 100 times less number of grid cells compared to a fixed mesh. The computational gain increases as the Reynolds increase. The reason for this effect is that the characteristic scales of the flow become smaller as the Reynolds increases and therefore, the efficiency of AMR increases for low resolution simulations.

Although the results provided by the model converge to a value relatively close to the experimental observations, we systematically observe that simulation results under-predict the experimental value for experimental conditions. To gain further insight about the source of this disagreement, we perform a systematic comparison of the non-uniformity index $\psi$ for the two different substrates and as a function of the Reynolds number. Even though the simulation results are consistent with the experimental results, Fig. 13 shows that as the substrate length decreases, the accuracy of the model decreases too. This effect is important for short monoliths and high Reynolds numbers. This reveals a limitation of the pressure jump model: while the model assumes a fully developed laminar flow inside the channels, in real conditions there is certainly a transition region at the channels entrance in which the pressure loss is not correctly captured by the Hagen-Poiseuille pressure drop model. As expected, as the length of the channels increases it is less important the transition region on the total pressure drop and the model predictions become more accurate. The accuracy of the solution is also larger as the Reynolds number decreases. At any event, we can conclude that the accuracy of the numerical model is satisfactory given the significant amount of computational time save with respect to the cost that would be implicated in the simulation of the flow inside the channels.

A finer analysis about the source of discrepancy between numerical and experimental results shows that the thickness of the boundary layer at the inlet has an impact on the results obtained. For instance, Fig. 14 compares the experimental and numerical velocity radial distribution for $Re=79,900$ assuming that the velocity profile at the inlet pipe corresponds to

$$U(y) = U_0 \text{erf} \left( \frac{y - R_{inlet}}{\delta} \right),$$

where $\delta$ is the boundary layer thickness at the inlet pipe. This parameter is important given that it controls the growthrate of the instability, which finally has an impact on the flow distribution downstream. The value of $\delta$ that provides the best fitting between simulation and experimental results is $\delta=5$ mm. In this case, numerical results converge to the velocity profile given by Benjamin et al. (2002) as we reduce the tolerance of the AMR criterion (Table 3). Unfortunately, the boundary layer thickness is not explicitly given in Benjamin et al. (2002), which means that the numerical fitting of $\delta$ may hide errors introduced by the models used and also by the discretization method. In any case, the good fitting between numerical and experimental results is remarkable due to the low number of grid points used and the large Reynolds number tested.

### Table 1
Simulation conditions used to test the subgrid model in Fig. 11.

<table>
<thead>
<tr>
<th>Dimensions of the solution domain</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pipe diameter ($D_{mon}$)</td>
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<tr>
<td>Pipe throughout a monolith substrate length</td>
</tr>
<tr>
<td>Catalyst diameter ($D_{catalyst}$)</td>
</tr>
<tr>
<td>Catalyst length</td>
</tr>
<tr>
<td>Reduced catalyst length in simulation</td>
</tr>
<tr>
<td>Catalyst channel diameter</td>
</tr>
<tr>
<td>Reynolds number at inlet pipe diameter ($Re$)</td>
</tr>
<tr>
<td>Inlet velocity profile</td>
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</tbody>
</table>

### Table 2
Simulation of experimental set-up.

<table>
<thead>
<tr>
<th>Dimensions of the geometry</th>
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<tbody>
<tr>
<td>Inlet pipe diameter</td>
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<tr>
<td>Inlet pipe length</td>
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<tr>
<td>Diffuser length</td>
</tr>
<tr>
<td>Total diffuser angle</td>
</tr>
<tr>
<td>Substrate diameter</td>
</tr>
<tr>
<td>Substrate length</td>
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<tr>
<td>Nominal substrate cell density</td>
</tr>
<tr>
<td>Fluid properties</td>
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<tr>
<td>Reynolds number at inlet pipe diameter ($Re$)</td>
</tr>
<tr>
<td>Inlet velocity profile</td>
</tr>
</tbody>
</table>
6. Numerical example

As an example of the capabilities of the developed models and numerical tools we present in this section the simulation results obtained for the simulation conditions included in Table 1 for using the whole geometry. The characteristic length of the system is chosen to be substrate diameter $D_{\text{max}}$.

All simulations were performed on a Dell Precision T5500 Westmere with a processor of Two Intel Xeon E5645 and total memory of 48 Go.

Fig. 15(a) depicts a sequence of snapshots of the resulting grid distribution obtained by vorticity based Hessian estimator. The grid is preferentially concentrated in the recirculation region, where most of the energy dissipation occurs (Fig. 15(b)) and where it is important to capture the flow features if one wants to correctly predict the flow distribution inside the different catalytic converter channels.

The combined use of the pressure drop model and AMR techniques significantly reduce the computational time (Fig. 16). We emphasize that this gain is expected to be even more significant in full three dimensional computations. The save in computational time is not at expenses of accuracy. As shown in Fig. 17 the velocity profiles at the medium plane obtained for the full numerical simulation and the simplified geometry match well. As it can be seen in Figs. 11 and 15(b) the reduced model captures remarkably well the influence of the substrate on the flow patterns upstream. The results are consistent with the flow patterns observed from the full simulation considering the substrate (Fig. 1) and also the flow patterns observed experimentally by Benjamin (2003).

7. Conclusion

In this paper we investigate a novel technique to couple the full solution of the Navier–Stokes equations in the divergent and convergent regions of a catalytic converter with a simplified physical model for the catalytic substrate.

The introduction of a new source term in the momentum equation allows us to capture the pressure drop induced by the catalytic substrate in the flow without the need of simulating the flow inside the catalytic channels. The model has been validated against the experimental results reported by Benjamin et al. (2002). The model still captures the influence of the substrate on the main flow features observed upstream saving a significant computational time.

In addition to the model for the catalytic substrate, specific multi-resolution techniques have been developed and validated against test cases related to the flow features observed in the diffuser region of catalytic converter systems. We conclude that by optimizing the grid distribution we can accelerate the simulation time by factor 10. This factor is expected to be larger for three dimensional computations.

To sum up, the coupling of models for the flow inside the catalytic substrate and the use of adaptive mesh refinement combined with efficient criteria for mesh adaptation produce optimum grid distributions that make possible to simulate complex flow problems involving different lengthscales in reasonable computational times. This work is currently being used as a solid
Fig. 14. Temporal evolution of the total grid points and convergence of the radial velocity profile at Re = 79,900 (based on inlet pipe diameter). The simulation results reproduce relatively well experimental results of Benjamin et al. (2002). (a) Number of total grid points as a function of the AMR tolerance (Table 3) and (b) velocity profiles for the three AMR tolerances contained in Table 3 ($R$ represents the radius of the substrate).

Table 3
Simulations for radial velocity profile convergence.

<table>
<thead>
<tr>
<th>Simulation</th>
<th>AMR tolerance $\eta$</th>
<th>Time averaged number of total grids</th>
</tr>
</thead>
<tbody>
<tr>
<td>Run 1</td>
<td>$\eta &lt; 0.7$</td>
<td>7970</td>
</tr>
<tr>
<td>Run 2</td>
<td>$\eta &lt; 0.5$</td>
<td>14,651</td>
</tr>
<tr>
<td>Run 3</td>
<td>$\eta &lt; 0.2$</td>
<td>28,799</td>
</tr>
</tbody>
</table>

Fig. 15. Grid size distribution (a) and non-dimensional vorticity (b) snapshots. The color scale for grid distribution represents the level of refinement, $l$, so that $l^2 = D_{max}/\Delta x$. The non-dimensionless vorticity is calculated as $\omega D_{max}/u_0$. The grid is preferentially concentrated in those regions where the vortical structures are present. (a) Grid distribution and (b) vorticity fields.
platform in which to implement simplified chemical reaction models capable of providing a complete description of all the physical and chemical processes taking place inside the catalytic system such as diesel oxidation catalyst (DOC), NOx trap catalyst, and selective catalytic reduction (SCR) catalyst.

References


