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1. INTRODUCTION
The Euler equations for compressible inviscid flow in a two dimensional domain $\Omega$,

$$q_x + f_x(q) + g_y(q) = 0,$$  \hspace{1cm} (1.1)

form a quasi-linear hyperbolic system of conservation laws. The state of the fluid at a point $(x,y) \in \Omega$ is given by $q(x,y) = (\rho, \rho u, \rho v, E)^T$. The fluxes in the $x$- (resp. $y$-) direction are

$$f(q) = (\rho u, \rho u^2 + p, \rho u v, u(E + p))^T$$

and

$$g(q) = (\rho v, \rho u v, \rho v^2 + p, v(E + p))^T.$$ 

For a perfect gas, $p$ and $E$ are related by the equation of state

$$p = (\gamma - 1)(E - \frac{1}{2}\rho(u^2 + v^2)).$$

With $(n_1, n_2)$ a pair of direction cosines, the flux in the $(n_1, n_2)$-direction is given by $n_1 f + n_2 g$. It is easily verified that the Euler equations are invariant under rotation of the independent variables. This means that, with the change of variables

$$\begin{pmatrix} x' \\ y' \end{pmatrix} = \begin{pmatrix} n_1 & n_2 \\ -n_2 & n_1 \end{pmatrix} \begin{pmatrix} x \\ y \end{pmatrix},$$

we obtain

$$q_x' + f_x'(q') + g_y'(q') = 0,$$

where

$$q' = \begin{pmatrix} 1 & 0 & 0 & 0 \\ 0 & n_1 & n_2 & 0 \\ 0 & -n_2 & n_1 & 0 \\ 0 & 0 & 0 & 1 \end{pmatrix} q.$$
Solutions of (1.1) are not necessarily smooth functions and it makes sense to generalize (1.1) to its weak form:

$$\frac{d}{dt} \int q(x,y) dx dy + \oint_{\partial \Omega} (qD_{\partial t} + n_2 q) ds = 0,$$

for all $\Omega \subset \Omega$.

where $(n_1,n_2)$ is the direction of the outward normal along $\delta \Omega$. Equation (1.2) allows also nonphysical solutions and it can be shown that a physical solution should also satisfy an entropy condition [6,14].

In symbolic form we write (1.1) or (1.2) as

$$q_t + N(q) = 0.$$  \hspace{1cm} (1.3)

The steady Euler equations are given by

$$N(q) = 0.$$  \hspace{1cm} (1.4)

Here $N: X \rightarrow Y$ is a nonlinear operator, $X \subset [L^2(\Omega)]^d$ is the space of possible fluid states and $Y = [L^2(\Omega)]^d$ is the Banach space of rates of change (of state).

\textbf{NOTATION}

$\Omega \subset \mathbb{R}^2$ the domain of definition for the Euler equations.

$\Omega_{ij}$ a cell in the partitioning of $\Omega$.

$\partial \Omega_{ij}$ the boundary of $\Omega_{ij}$.

$\Omega_{ijk}$ a neighbouring cell of $\Omega_{ij}$, $k=N,S,E,W$.

$\Gamma_{ijk} = \Omega_{ij} \cap \Omega_{ik}$.

$q_{ij}$ an approximation for the mean state in $\Omega_{ij}$

$$q_{ij} \text{ meas (} \Omega_{ij} \text{)} = \int q(x,y) dx dy.$$  \hspace{1cm} q_{ij}

$q_{ijk}$ an approximation for the mean state in $\Omega_{ijk}$.

$q_k = \{ q_{ij} \mid \Omega_{ij} \subset \Omega \}$.  \hspace{1cm} q_k

$p$ pressure.

$\rho$ density.

$u,v$ speed of fluid in $x,y$ direction.

$E$ total energy per unit volume.

$c = \sqrt{\gamma p / \rho}$ speed of sound.

$z = \ln(p \rho^{-\gamma})$ specific entropy.

$\gamma$ ratio of specific heats, $\gamma = 1.4$.

\textbf{2. Discretization}

In order to achieve a discretization of a hyperbolic system of conservation laws that has a significant meaning even if the meshsize is coarse, it is important to use a fully conservative method which discretizes the weak form of the equations. A simple but effective technique is found in the finite volume discretization. This technique allows for an irregular partitioning of the domain $\Omega$ in a number of disjunct cells. For ease of notation and implementation, we divide the bounded domain $\Omega$ in quadrilateral cells $\Omega_{ij}$, such that the result is topological equivalent with a partitioning in regular
squares. In each cell the state of the fluid is represented by \( q_{ij} \), the mean value of \( q(x,y) \) over \( \Omega_{ij} \). The semi-discrete form of the space-discretized equation is

\[
\text{meas} \left( \Omega_{ij} \right) \frac{d}{dt} q_{ij} + \int_{\partial \Omega_{ij}} (n_1 f + n_2 g) ds = 0, \text{ for all } \Omega_{ij} \subset \Omega, \tag{2.1}
\]

where \( n_1 f + n_2 g \) denotes the normal flux outward \( \Omega_{ij} \). For the steady equation on our mesh this reduces to

\[
\sum_{k=E,S,W} \int_{\Gamma_{ij}} (n_1 f + n_2 g) ds = 0, \text{ for all } i,j, \tag{2.2}
\]

Here \( \int (n_1 f + n_2 g) ds \) denotes the rate of transport of \( q \) over the boundary \( \Gamma_{ij} \) from cell \( \Omega_{ij} \) to the neighbouring cell \( \Omega_{ijk} \), i.e. the flux from \( \Omega_{ij} \) to \( \Omega_{ijk} \) multiplied by \( \text{meas} (\Gamma_{ijk}) \). In the discretization this flux \( n_1 f + n_2 g \) is approximated by the numerical flux \( f(q_{ij}, q_{ijk}) \). In a first order discretization scheme this numerical flux over \( \Gamma_{ij} \) depends only on the unknown states \( q_{ij} \) and \( q_{ijk} \). For a given choice of the numerical flux, the discrete system for the steady equations (2.2) can formally be written

\[
N_k(q_k) = 0. \tag{2.3}
\]

This is the discrete system of nonlinear equations of which the solution is required. The operator \( N_k : X_k \rightarrow Y_k \) is the nonlinear discrete operator, \( X_k \) is the (finite dimensional) linear space of mean states in cells \( \Omega_{ij} \), and \( Y_k \) is the (finite dimensional) linear space representing the rate of netto transport into the cells \( \Omega_{ij} \).

To obtain a good discretization, the selection of a good numerical flux is essential. Such a flux should take into account the fact that, depending on the local characteristics, the flux at \( \Gamma_{ij} \) is -in a specific way - depending on \( q_{ij} \) and \( q_{ijk} \). E.g. in case of a supersonic flow from \( \Omega_{ij} \) to \( \Omega_{ijk} \) this flux depends only on \( q_{ij} \).

One way to determine the numerical flux is to consider the flux computation at \( \Gamma_{ijk} \) as a locally one-dimensional problem and to solve the Riemann problem of gasdynamics: compute the flux at \( \Gamma_{ijk} \), \( 0 < t < t_0 \) with at \( t = 0 \) the initial conditions \( q = q_{ij} \) in \( \Omega_{ij} \) and \( q = q_{ijk} \) in \( \Omega_{ijk} \). The use of this computed flux as the numerical flux in (2.3) yields the Godunov discretization. A disadvantage is the expensive solution of the Riemann problem at each cell boundary. Several less expensive approximate Riemann solvers have been proposed. These lead to various well known flux (difference) splitting methods [2,8,9,11,12,15,17,18].

Selecting a numerical flux we should take care that it should (1) yield shocks which satisfy correct jump conditions, (2) find only physically correct shocks, i.e. shocks satisfying an entropy condition, and (3) yield nonlinear stability (a monotone nonlinear system (2.3)). Further it is an advantage to have a differentiable function \( f(q_{ij}, q_{ijk}) \) if a Newton-type technique is used in the solution process for (2.3).

A very good numerical flux, that satisfies these desired properties, is generated by Osher’s approximate Riemann-solver [8,9]. This numerical flux is based on the Riemann-invariants which relate different states that are connected by simple waves [14]. The good qualities of this numerical flux are often considered to be offset by its relative complexity. However, we found this disadvantage improved by the use of a proper set of dependent variables, viz. \( u, v, c \) and \( z \) [3].

An additional advantage of the numerical flux computation based on Osher’s approximate Riemann solver is the fact that the treatment of boundary conditions, i.e. the computation of the boundary fluxes, can be done in a way that is completely consistent with the computation of fluxes over interior cell walls. For this purpose the equations are considered quasi-one-dimensional, normal to \( \partial \Omega \), and a state \( q_{ijB} \) at the boundary \( \delta \Omega_{ij} \cap \delta \Omega \) can be calculated such that \( q_{ijB} \) satisfies the boundary conditions and such that \( q_{ijB} \) and \( q_{ij} \) are connected by Riemann invariants that correspond to negative (right boundary) or positive (left boundary) eigenvalues [3].
It appears that in the MG-iteration such a consistent treatment of the equation and the boundary conditions makes a special relaxation treatment of the boundary superfluous. (A similar effect is seen when an elliptic equation together with its boundary conditions is consistently discretized by the finite element method.)

3. Solution methods and linearization

Globally there are three ways to solve the nonlinear system (2.3). First, one can start from the semi-discretized form (2.1) and solve the large system of ordinary differential equations

\[ D_h \frac{d}{dt} q_h + N_h(q_h) = 0 \]  

(3.1)

by some explicit time integrator; \( D_h \) is a diagonal matrix. In this way the time-dependent behaviour of the flow is followed, starting from some initial condition. Integration over a long enough time interval may make the solution of (3.1) converge to the solution of (2.3). An advantage is that intermediate values of \( q_h \) allow a physical interpretation. If (2.3) has a non-unique solution, the proper choice of an initial condition may select the solution required. Another advantage is that this method needs only evaluations of the operator \( N_h(q_h) \). If the time-dependent solution is not wanted, it is a disadvantage that usually many time-steps are needed to obtain a sufficiently converged final solution \( q_h \). Stability conditions may prevent the use of large time steps. Multigrid may provide a technique to accelerate the convergence of this time stepping [4,13].

Secondly, the equation (2.3) may be solved by some implicit time integrator, or -what is closely related- the system (2.3) or (3.1) can be solved by some global linearization. Examples are the application of a Newton-type method or the use of the SER- (Switched Evolution Relaxation) scheme as used by Mulder and Van Leer [7],

\[ [D_h + \Delta t \cdot N_h(q_h^{(\alpha)})] (q_h^{(\alpha+1)} - q_h^{(\alpha)}) = -\Delta t \cdot N_h(q_h^{(\alpha)}) . \]  

(3.2)

In these methods large linear systems are to be solved and the construction of the Jacobian matrix

\[ N_h(q_h) \]

is required. For the solution of the linear system (e.g. in eq. 3.2), several techniques are available. Multigrid in its linear form (the Correction Scheme) [1] can be used to accelerate iterative methods for the solution of the linear systems.

In particular Newton’s method will give very fast convergence, provided that a sufficiently accurate initial estimate is available (and that a singular Jacobian is avoided). If the accurate initial estimate is not available, continuation or time-stepping techniques may slacken the solution process as was the case with the explicit time integration.

Finally, the equation (2.3) may be solved directly by means of a non-linear relaxation method. If (2.3) sufficiently satisfies stability (monotonicity) conditions, simple relaxation methods may converge. Rather than a good global convergence rate, we may expect that local relaxations -such as point Gauss-Seidel methods- will be able to smooth the error. Multigrid in its nonlinear form FAS (the Full Approximation Scheme) [1] is a proper technique to accelerate the convergence of these relaxations. The nonlinear relaxation iteration with FAS seems the most direct way to solve (2.3) and it is this approach that we follow in this paper.

In a local relaxation sweep, for each \( \Omega_j \) one set of four equations,

\[ (N_h(q_h))_{ij} = 0 , \]  

(3.3)

is solved (approximately). For these four equations no natural ordering exists and -hence- we solve these equations simultaneously (collective relaxation). Now only the local linearization, i.e. linearization of (3.3) with respect to \( (q_h)_{ij} \) is needed.
Besides the reduced sensitivity for accurate initial estimates [16], it is an additional advantage of nonlinear relaxation that the storage requirements are significantly less. Only 4*4-systems are solved and no global Jacobian matrix needs to be kept, whereas a global linearization requires 80*N real numbers to store the Jacobian; N is the number of cells in the mesh. Probably a global linearization only pays off in a final stage of the solution process for (2.3), when Newton’s method guarantees quadratic convergence. For a further discussion of the choice between Newton-Multigrid or Multigrid-Newton see [5].

Independent of the type of linearization that is actually used, it is important to see the structure of the Jacobian matrix of (2.3). For the finite volume discretization, the global Jacobian matrix is assembled in a way that is similar to the assembling process for finite elements. The finite volume Jacobian is the sum of small block-2*2 cell wall matrices (for FEM: element matrices). The regular structure of the mesh used induces a block-5-diagonal structure in which each block entry itself is a 4*4 matrix. The Jacobian and the rhs are assembled simultaneously and for each cell wall, $\Gamma_{ijk}$, the entries $\pm f(q_{ij}, q_{ijk})$

$$\pm A_{ijk}^+ = \pm \frac{\partial}{\partial q_{ij}} f(q_{ij}, q_{ijk}) \text{ and } \pm A_{ijk}^- = \pm \frac{\partial}{\partial q_{ijk}} f(q_{ij}, q_{ijk})$$

are evaluated and added as a contribution to the rhs or the Jacobian matrix respectively (see figure 1). The fact that all column sums are zero, except for boundary contributions, reflects that the discretization is conservative.

A row from the Jacobian, corresponding to the cell $\Omega_{ij}$, is now seen to be of the form

$$-A_{i,j-\frac{1}{2},j}^- + A_{i,j+\frac{1}{2},j}^+$$

$$-A_{i-\frac{1}{2},j}^+ , \ -A_{i,j-\frac{1}{2}}^- + A_{i,j+\frac{1}{2}}^+ , \ +A_{i,j+\frac{1}{2}}^- , \ +A_{i,j+\frac{1}{2},j}^+$$ (3.4)

This structure of the Jacobian matrix is to be exploited when relaxation methods for the system are analyzed.

Figure 1. Assembling the rhs and the Jacobian of the nonlinear system.

In figure 1 the notation $\begin{bmatrix} P & P \\ P & P \end{bmatrix} \begin{bmatrix} P \\ P \end{bmatrix}$ is short for

$$\begin{pmatrix} +A_{ijE} & +A_{ijE} \\ -A_{ijE} & -A_{ijE} \end{pmatrix} \begin{pmatrix} f(q_{ij}, q_{ijk}) \\ -f(q_{ij}, q_{ijk}) \end{pmatrix}$$ (3.5)
where \( A_{ij} = A_{i,j+\frac{1}{2}} \).

The boundary condition treatment is straightforward as soon as we take the view that for each \( \Gamma_{ij} = \Omega \cap \Omega_{ij} \) we can determine a \( q_{ij} \) as is introduced in section 2. This boundary state depends on the state in the neighbouring cell: \( q_{ij} = q_{ij}(q_{ij}) \). Hence, we have for the contribution from \( \Gamma_{ij} \) to (2.3)

\[
f_{ij} := f(q_{ij}(q_{ij}), q_{ij})
\]

and

\[
\frac{\partial}{\partial q_{ij}} f_{ij} = \frac{\partial}{\partial q_{ij}} f(q_{ij}, q_{ij}) + \frac{\partial}{\partial q_{ij}} f(q_{ij}(q_{ij}), q_{ij}).
\]

(3.6)

\[
= A_{ij} + A_{ij} \frac{\partial q_{ij}}{\partial q_{ij}}.
\]

This completes the description of the linearization of (2.3).

The local linearization of (3.3) gives a coefficient matrix as appears in the main diagonal of (3.4). For cells near \( \delta \Omega \) these systems are augmented by boundary terms as given in (3.6).

4. THE NESTED SEQUENCE OF DISCRETIZATIONS

For the P-variant of Osher's scheme [3], when the flow field is sufficiently smooth (no shocks present), for any \( q_{h} \), defined on the grid \( \Omega_{h} = (\Omega_{ij}) \), piecewise constant states \( q_{h},k \) can be defined at the cell boundaries \( \Gamma_{ij} \), such that \( f(q_{h,k}) = f(q_{ij}, q_{ij}) \) or, at the boundary of \( \Omega_{h} \), \( q_{h,k} = q_{ij} \). Under these assumptions the finite volume method can be seen as a formal weighted residual method for the discretization of \( N(q) = 0 \).

We can write the discrete operator \( N_{h} \) as a Galerkin approximation to \( N \),

\[
N_{h}(q_{h}) = R_{h}(N_{h} q_{h}),
\]

(4.1)

where \( P_{h} : X_{h} \rightarrow X \) relates to each \( q_{h} \) a function \( P_{h} q_{h} \) on \( \Omega_{h} \) for which

\[
\begin{cases}
P_{h} q_{h}(s) = q_{ij} & \text{for } s \in \Omega_{ij} \\
P_{h} q_{h}(s) = q_{h,k} & \text{for } s \in \Gamma_{ij}
\end{cases}
\]

where \( q_{h,k} \) is such that \( f(q_{h,k}) = f(q_{ij}, q_{ij}) \). I.e. piecewise constant functions with upwind continuity at the boundaries for characteristic information.

The restriction \( R_{h} : Y \rightarrow Y_{h} \) is defined by

\[
(\bar{R}_{h} r)_{ij} = \int_{\Omega_{h}} r(x, y) \, dx \, dy,
\]

so that, formally,

\[
(\bar{R}_{h} N(q))_{ij} = \int_{\Omega_{h}} f_{x}(q) + g_{y}(q) \, dx \, dy
\]

\[
= \int_{\Omega_{h}} n_{x} f(q) + n_{y} g(q) \, dx
\]

\[
= \sum_{k \in \Gamma_{h}} \int_{\Gamma_{h}} (n_{x} f + n_{y} g) q \, ds
\]

and

\[
(\bar{R}_{h} N(P_{h} q_{h}))_{ij} = \sum_{k \in \Gamma_{h}} \int_{\Gamma_{h}} (n_{x} f + n_{y} g)(P_{h} q_{h}) \, ds
\]

(4.2)
\[
\sum_k \int_{\Gamma_{ik}} (n_f^2 + n_2 q) (q_{ijk}) \, ds \\
= \sum_k \text{meas} (\Gamma_{ijk}) f(q_{ijk}).
\]

Now, a regular sequence of nested discretizations is found in the following way. Start with a partitioning of \( \Omega \) in cells \( \{ \Omega_{ij} \}_{0 < i \leq n_1 2^l, 0 < j \leq n_2 2^l} \). This yields the finest level of discretization (level 1). For each \( k = l, l - 1, l - 2, \ldots, 1 \), a coarser level of discretization \( k - 1 \) is defined by deleting each second meshline, so that each time 4 cells in the finer mesh correspond to a single coarser cell. In this way sequences of discrete spaces \( X_h, X_{2h}, X_{4h}, \ldots \), and \( Y_h, Y_{2h}, Y_{4h}, \ldots \), are obtained. With a prolongation between the discrete solution spaces \( P_{h,2h} : X_{2h} \to X_h \), defined by "piecewise constant interpolation" (distribute a coarse cell value \( q_{ij} \) as the same value over 4 finer cells), and a restriction \( R_{2h,h} : Y_h \to Y_{2h}, \) defined by adding the 4 fine cell values to obtain the coarse cell value, we have the relation

\[
P_{2h} = P_h P_{h,2h} \quad \text{and} \quad R_{2h} = R_{2h,h} R_h.
\]

In this way we obtain a commutative diagram for the -now nested- set of discretizations \( N_h, N_{2h}, \ldots \).

![Diagram](image)

Figure 2. The nested set of discretizations.

The prolongation and restriction introduced in this way are the natural operators that correspond with the physical meanings of \( X \) and \( Y \). An \( q_h \in X_h \) is associated with the mean states of the fluid in the cells \( \{ \Omega_{ij} \} \), whereas \( r_h \in Y_h \) is associated with the netto transport rate of the conservative quantities into the cells \( \{ \Omega_{ij} \} \).

![Diagram](image)

Figure 3. The first order finite volume prolongation \( P_{h,2h} \) and \( R_{2h,h} \) restriction.
The operators $P_{h,2h}$ and $\widetilde{R}_{2h,h}$ are also used in section 5 as grid transfer operators in the FAS multigrid algorithm. In the case of CS-multigrid they can be used for the efficient construction of the coarse grid Jacobian matrices. Then the operator $R_{2h,h}N_h(q_h)p_{h,2h}$ can be used instead of $N_{2h}(q_{2h})$. This Galerkin coarse grid Jacobian has the usual (additional) advantage that -after a coarse grid correction- the restriction of the residual vanishes and -thus- the residual contains mainly Fourier components with high frequencies.

5. THE MULTIGRID METHOD
Based on the sequence of discretizations $N_h, N_{2h}, N_{4h}, \ldots$ the Full Approximation Scheme (FAS) multigrid algorithm has been used to accelerate the nonlinear point Gauss Seidel relaxation.

One iteration step of the FAS-algorithm for the solution of $N_h(q_h) = r_h$ is defined as

1. Execution of $p$ pre-relaxation steps.
2. Execution of a coarse grid correction, i.e.
   2a. for some given $q_{2h}$ compute
       $$r_{2h} = N_{2h}(q_{2h}) + \widetilde{R}_{2h,h}(r_h - N_h(q_h));$$
   2b. determine by $\sigma$ FAS-iteration steps $\tilde{q}_{2h}$, the approximate solution of
       $$N_{2h}(q_{2h}) = r_{2h};$$
   2c. replace $q_h$ by $q_h + P_{h,2h}(\tilde{q}_{2h} - q_{2h});$
3. execution of $q$ post-relaxation sweeps.

On the coarsest grid no coarse grid correction is executed.

Experiments show that $\sigma > 1$ generally gives no more efficient results than $\sigma = 1$ (the V-cycle). A strategy that usually yields efficient results is $p = q = \sigma = 1$, where collective symmetric Gauss-Seidel was used as a relaxation method. The ordering of the lexicographical Gauss-Seidel relaxation was from north-west to south-east -vice versa- in the pre-relaxation, and from north-east to south-west -vice versa- during post-relaxation. This strategy was adopted as a standard strategy and it is compared with various other variants in section 6.

Initial estimates are obtained by the Full Multi Grid (FMG) technique. For $k = 0, 1, 2, \ldots, l - 1$ the initial approximation on level $k + 1$ is obtained by

1. Application of a single FAS-cycle to the solution on level $k$, and
2. Interpolation of the approximate solution on level $k$ to level $k + 1$.

For this interpolation not the prolongation $P_{h,2h}$ is used. The piecewise constant prolongation is not sufficiently accurate to interpolate the first order accurate coarse grid solution to the finer mesh. For the interpolation operator, a bilinear blockwise interpolation is used: the bilinear interpolation of the solution found on a coarse $2 \times 2$ block of cells $\Omega_{ij}$ is transferred to a $4 \times 4$ block of cells on the finer grid.

6. NUMERICAL RESULTS
To show some properties of our method, we computed flows through a channel with a circular bump. As a first test, a standard testproblem [10] was chosen. This problem concerns a transsonic flow. Further tests were made by variation of parameters, such that the flow became supersonic or subsonic. Tests were made both on adapted grids as described in [10] and on non-adapted regular square grids. In this paper, concerned with the convergence of the method rather than with the representation of the solution, we restrict ourselves to regular grids. For tests on the adapted grids the reader is referred to [3].

In a sense, the tests on a regular grid are harder because it is less clear that the coarse grid discretizations have a significant meaning for the problems on the finer grids. In particular, on the
coarsest level the meshsize was so large that the discretization cannot recognize the circular bump(!).

The problem description is given in figure 4.

Figure 4 The testproblems

On the northern wall \( v = 0 \) is specified, on the southern wall the ratio \( v/u \) is specified in such a way that it is in agreement with the direction of the wall (dependent on \( x \)). At the inflow boundary \( u, v, \gamma \) are specified. For supersonic flow on this boundary also \( \gamma \) is given. For transonic or subsonic flow the pressure \( p \) is given at the outflow boundary. The thickness of the bump is \( \delta = 0.042 \).

The coarsest grid is a \( 4 \times 2 \) grid, as indicated in figure 4. Finer grids were obtained by regular subdivision of the coarse cells. The boundary condition at the bump was satisfied at the mid-cell-wall. As a consequence, on the coarsest grid a uniform flow is the solution of the discrete problem!

In the figures 5 to 12 we show the convergence histories of the FAS iteration for the following testproblems.

**Problem 1** Supersonic flow: \( u = 3.0, v = 0.0, \gamma = 1.0, z = -\gamma \ln(\gamma) \)

**Problem 2** Transsonic flow: \( u = 0.85, v = 0.0, \gamma = \ln(\gamma^{\gamma-1}), p = 1.05 \)

**Problem 3** Subsonic flow: \( u = 0.3, v = 0.0, z = \ln(\gamma^{\gamma-1}), p = 1.05 \)

In the figures the norm of the residual, \( ||N(q_h)|| \), is plotted against the iteration number. The norm used is the maximum of the four \( L_1 \)-norms of the components in the residual. The numbers 2, 3, 4 or 5 in the figures denote the number of levels used.

The P-variant of Osher's approximate Riemann solver [3] was used for the discretization. The three problems have been run for the (standard) strategy, as described in section 5. Further experiments have run with Red-Black Gauss-Seidel relaxation. Other experiments have run with the W-cycle instead of the V-cycle.

From the experiments we conclude that for the supersonic and transsonic flow the rate of convergence of FAS is independent of the meshwidth. Convergence is slower and dependent of the meshwidth for small Machnumbers. Red-black Gauss-Seidel relaxation is slower than Symmetric Gauss Seidel relaxation, but for vector architectures it may still be competitive.

If boundary conditions are over-specified, i.e. if complete states of flow are specified at the inflow and outflow boundaries (and, hence, the upwind scheme may select the best defined boundary conditions), the problem - being better posed - converges faster, as long as the number of meshpoints is small enough.

Experiments for which no figures are given, show that W-cycles give almost the same convergence behaviour as V-cycles. Bilinear prolongations in FAS give almost the same convergence behaviour as
piecewise constant prolongations. Also, when the O-variant of Osher's scheme was used [3], no significant differences were observed.

7. CONCLUSION
An efficient multigrid method for the solution of the steady Euler equations could be developed. The backbone of the method is a nested sequence of Galerkin discretizations, which is constructed by the finite volume technique. To obtain a stable discretization and a good representation of the solution (sharp captured shocks, no sonic glitches, etc.) and to find a boundary condition treatment that is consistent with the interior discretization, the use of an powerful numerical flux is essential. Such a numerical flux was found in Osher's approximate Riemann-solver.

When the FMG-technique is used for the computation of initial estimates, it appears that a few FAS cycles are often enough to obtain a solution of the discrete system that is accurate upto truncation error.

Numerical experiments show that in several transonic testcases the rate of convergence of the FAS iteration is independent of the gridsize. For a subsonic testcase, \( M = 0.3 \), this could not be observed.

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Residual (abscissa) versus number of FAS-cycles (ordinate)
Residual (abscissa) versus number of FAS-cycles (ordinate)