# Multigrid as a useful solver/preconditioner for blocks

Iterative Methods for Large-Scale Saddle-Point Problems

#### Fabio Durastante

Università di Pisa



#### **Overview**

### 1. The Multigrid Idea

- 1.1 Multigrid based on geometry
- 1.2 A general two grid formulation
- 1.3 Smooth and stable components

#### 2. Algebraic multigrid

- 2.1 Ruge-Stuben Splitting Algorithm
- 2.2 Aggregation-based multigrids
- 2.3 Compatible relaxation
- 2.4 Available libraries

#### 3. Extreme scale applications

- 3.1 The Poisson benchmark
- 3.2 A Large Eddy Scale simulation



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These methods start from a simple idea, but the confusion of their explanation grows exponentially with the degree of generality and abstraction that one wants to impose.

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#### Our plan

Therefore we will start explaining them in a completely abstract framework. Therefore we will start from the simplest 1D example, and work on that.

Good introductions are contained in the books (Briggs, Henson, and McCormick 2000; Trottenberg, Oosterlee, and Schüller 2001; Vassilevski 2008), for the more theoretically inclined the best high-level presentation is in (Xu and Zikatanov 2017).

Let us consider the following boundary value problem

$$\begin{cases} -u_{xx}(x) = f(x), & x \in (0,1), \\ u(0) = u(1) = 0. \end{cases}, f \in C^{1}([0,1]).$$

If we apply **standard centered finite difference** discretization on the grid  $\Omega_h = \{x_k\}_{k=0}^{n+2} = \{kh\}$  and h = 1/(n+2), that gives rise to the linear system

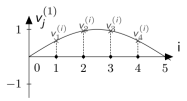
$$\frac{K}{h^2} A_n \mathbf{u} = \mathbf{f}, \qquad A_n = \begin{bmatrix}
2 & -1 & 0 & \cdots & 0 \\
-1 & 2 & -1 & \cdots & 0 \\
0 & \ddots & \ddots & \ddots & \vdots \\
\vdots & \ddots & -1 & 2 & -1 \\
0 & \cdots & 0 & -1 & 2
\end{bmatrix}$$

The matrix  $A_n$  is a **very peculiar type of matrix** for which we now everything, specifically:

$$A_n \mathbf{v}^{(i)} = \lambda_i \mathbf{v}^{(i)},$$

with eigenvalues and eigenvectors

$$\lambda_i = 2 - 2\cos\left(\frac{i\pi}{n+1}\right), \quad \mathbf{v}_j^{(i)} = \sin\left(\frac{ij\pi}{n+1}\right), \ i, j = 1, \dots, n.$$

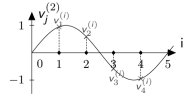


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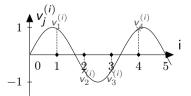
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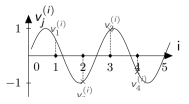
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We divide **arbitrarily** this set of frequencies in two subsets:

Now we have put some notation in place, but let us try solving our system with the simplest method we know: Jacobi!

$$\mathbf{u}^{(k+1)} = \mathbf{u}^{(k)} + D_n^{-1} (\mathbf{f} - A_n \mathbf{u}^{(k)}) = \left( I_n - \frac{1}{2} A_n \right) \mathbf{u}^{(k)} + \frac{1}{2} \mathbf{f},$$

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• The iteration matrix is then  $J_n = I_n - A_n/2$ , with the information we have on the spectrum, we observe that:  $\rho(J_n) \to 1$  for  $n \to +\infty$  so slow convergence! A sorry state of affairs.

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$$\mathbf{u}^{(k+1)} = \left[ (1 - \omega) I_n + \omega \left( I_n - \frac{1}{2} A_n \right) \right] \mathbf{u}^{(k)} + \frac{1}{2} \omega \mathbf{f},$$

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- Maybe we can **weight the iteration** to make things better. The iteration matrix is now  $J_n^\omega = I_n \frac{\omega}{2} A_n$ . The best spectral conditioning for  $\mu_1^\omega = \lambda_{\max}(J_n^\omega)$  is obtained for  $\mu_1^1 < \mu_1^\omega \ \forall \omega \in (0,1)$ . Therefore, we have only made the convergence worse...

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We hold the line, let us write everything in the eigenvector basis:

$$\mathbf{e}^{(0)} = \sum_{i=1}^{n} \alpha_i \mathbf{v}^{(i)}, \ \ J_n^{\omega} = V \Lambda_n^{\omega} V^T \ \ ext{where} \ \Lambda_n^{\omega} = \mathrm{diag}(\mu_i^{\omega}),$$

at the kth step (for  $\overline{\mathbf{u}} = A_n^{-1} \mathbf{f}$  the true solution) we find

$$\overline{\mathbf{u}} - \mathbf{u}^{(k)} = \mathbf{e}^{(k)} = (J_n^{\omega})^k \mathbf{e}^{(0)} = V(\Lambda_n^{\omega})^k V^T \mathbf{e}^{(0)} = V(\Lambda_n^{\omega})^k \alpha$$
$$= \sum_{i=1}^n \left( 1 - 2\omega \sin^2 \left( \frac{i\pi}{2(n+1)} \right) \right)^k \alpha_i \mathbf{v}^{(i)}.$$

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#### 🖢 Idea!

The *i*th entry of  $e^{(k)}$  is defined in terms of the *i*th eigenvalues of  $J_k^{\omega}$ :

$$\beta_i = \left(1 - 2\omega \sin^2\left(\frac{i\pi}{2(n+1)}\right)\right)^k \alpha_i \approx \left(1 - \omega \frac{\pi^2}{2}h^2\right)^k \alpha_i.$$

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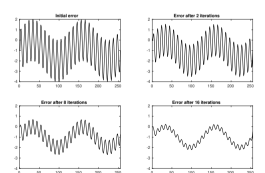
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#### Working only in the high-frequency

We choose an *optimal*  $\omega$  that minimizes the absolute values of the  $\beta_i$  in the high frequencies, i.e.,  $\omega_{\text{opt}} = 2/3$ .

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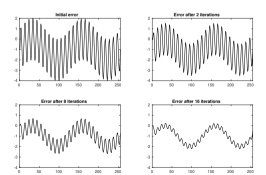


#### Solver as smoother

In the domain of the high frequencies, for whatever value of k, we find that the error has become a smooth function.

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In the domain of the high frequencies, for whatever value of k, we find that the error has become a smooth function.

But the convergence is always bad...

### Multigrid based on geometry: Error equation

• Let us suppose that we have computed an approximation  $\widetilde{\mathbf{u}}$  of the solution  $\overline{\mathbf{u}}$  through some iterations of the optimally weighted Jacobi.

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If we could compute in some way the error e then the solution of the linear system could be obtained as  $\overline{u}=\widetilde{u}+e$ 

If we wanted to compute it, we could solve the linear system

$$A_n \mathbf{e} = A_n \overline{\mathbf{u}} - A_n \widetilde{\mathbf{u}} = \mathbf{f} - A_n \widetilde{\mathbf{u}} = \mathbf{r}.$$

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#### Still no gain?

We are back solving a linear system with the same coefficient matrix. **Nevertheless**, having swapped from the need of computing  $\mathbf{u}$  to  $\mathbf{e}$  gives us the chance to exploit the information that the error has been smoothed.

### Multigrid based on geometry: Grids

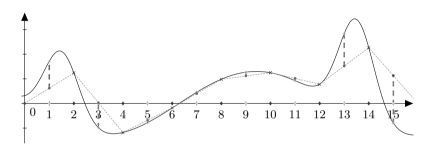
Let us consider the grids (for an odd n):

$$\Omega_{h} = \left\{ \frac{i\pi}{n+1} : i = 1, \dots, n \right\}, 
\Omega_{2h} = \left\{ \frac{2i\pi}{n+1} : i = 1, \dots, \frac{n-1}{2} \right\} = \left\{ \frac{i\pi}{\frac{n-1}{2}+1} : i = 1, \dots, \frac{n-1}{2} \right\},$$

We restrict the matrix and the residual vector on the coarse grid to solve the error equation

$$\Omega_h \rightsquigarrow A_n \mathbf{u} = \mathbf{f}, \quad \mathbf{u}, \mathbf{f} \in \mathbb{R}^n,$$
  
$$\Omega_{2h} \rightsquigarrow A_{\frac{n-1}{2}} \widetilde{\mathbf{e}} = \widetilde{\mathbf{r}}, \quad \widetilde{\mathbf{e}}, \widetilde{\mathbf{r}} \in \mathbb{R}^{\frac{n-1}{2}}.$$

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### Multigrid based on geometry: the whole idea

- 1. We apply the **smoother** to smooth the error in the high frequency,
- 2. We use our **coarsening strategy** for the **error equation** 
  - 2.1 a restriction operator  $I_h^{2h}: \Omega_h \to \Omega_{2h}$ ,
  - 2.2 a prolongation operator  $I_{2h}^h:\Omega_{2h}\to\Omega_h$ ,
  - 2.3 the discretization matrix at the lower level, i.e.,  $A_{n-1/2}$ .
- 3. We can make additional distinctions between high and low frequencies for the error equation with respect to the actual grid  $\Omega_{2h}$  and a coarser grid  $\Omega_{4h}$  to iterate our coupling of smoothing iterations and iterative refinement by coarsening
- 4. We do something peculiar on the **coarsest grid** in which we face a very small linear system, possibly a single linear scalar equation, that can be solved efficiently by a direct method.

## Multigrid based on geometry: the algorithm

```
Data: \{A_k\}_{k=1}^0, I, \{S_k^{(1)}\}_{k=1}^0, \{S_k^{(2)}\}_{k=1}^0, \{I_k^{(k-1)}\}_{k=1}^0 and \{I_{k=1}^k\}_{k=1}^0, \{I_k^{(k-1)}\}_{k=1}^0, initial guess \mathbf{u}^{(j)}.
Output: Approximation \mathbf{u}^{(j+1)} to the solution of \mathbf{x}_l.
Input: \mathbf{u}_{k}^{(j+1)} = \text{MGM}(A_{k}, \mathbf{b}_{k}, \mathbf{x}_{k}^{(j)}, k, \nu_{1}, \nu_{2}, \gamma)
\nu_1 steps of presmoother S_{\iota}^{(1)} applied to A_{\iota}\widetilde{\chi}_{\iota}^{(j)} = \mathbf{b}_{\iota}:
                                                                                                                                                          // Presmoothing
Compute the residual \mathbf{r}_{k}^{(j)} = \mathbf{b}_{k} - A_{k} \widetilde{\mathbf{x}}_{k}^{(j)}:
                                                                                                                                    // Coarse Grid Correction
Restrict the residual \mathbf{r}_{k-1}^{(j)} = \mathbf{I}_{k}^{k-1} \mathbf{r}_{k}^{(j)};
if k = 1 then
       Direct solver for A_{k-1}e_k^{(j)}:
else
      for i = 1, \ldots, \gamma do
  \mathbf{e}_{k-1}^{(j)} = \text{MGM}(\mathbf{A}_{k-1}, \mathbf{r}_{k-1}, 0, k-1, \nu_1, \nu_2, \gamma)
end
Prolong the error \mathbf{e}_{k}^{(j)} = I_{k-1}^{k} \mathbf{e}_{k-1}^{(j)}:
Update the approximation \mathbf{x}_{h}^{(j)} = \widetilde{\mathbf{x}}_{h}^{(j)} + \mathbf{e}_{h}^{(j)}:
\nu_2 steps of postsmoother S_i^{(2)} applied to A_{\nu} \widetilde{x}_i^{(j+1)} = \mathbf{b}_{\nu} with initial guess \mathbf{x}_i^{(j)}:
                                                                                                                                                    // Postsmoothing
```

We express the previous algorithm as the product by an **iteration matrix**  $M_I$ :

$$\begin{cases} M_0 = 0, & k = 0, \\ M_k = (S_k^{(1)})^{\nu_1} \left( I_k - I_{k-1}^k (I_{k-1} - M_{k-1}^{\gamma}) A_{k-1}^{-1} I_k^{k-1} A_k \right) (S_k^{(2)})^{\nu_2} & k \ge 1. \end{cases}$$

**Q Idea:** it is a stationary method, thus it converges iff  $\rho(M_k) < 1$ .

#### Convergence theorem

Let  $A \in \mathbb{R}^{n \times n}$  be SPD. Assume that the prolongation operators  $I_{k-1}^k$  have full rank and that the Galerkin conditions holds

$$I_{k-1}^k = (I_k^{k-1})^T$$
,  $A_{k-1} = I_k^{k-1} A_k I_{k-1}^k$ ,  $\forall k = l-1, \dots, 0$ ,

Furthermore, if the orthogonal projector  $\Pi_k = I - I_{k+1}^k A_{k+1}^{-1} I_k^{k+1} A_k$ , satisfies

$$\forall \mathbf{e}_{k} \exists \delta_{1} > 0 : \|S_{k}^{(2)} \mathbf{e}_{k}\|_{A}^{2} \leq \|\mathbf{e}_{k}\|_{A}^{2} - \delta_{1} \|\Pi_{k} \mathbf{e}_{k}\|_{A}^{2},$$

independently of  $e_k$  and k, then the multigrid method with  $\gamma=1$ ,  $\nu_1=0$  and  $\nu_2\geq 1$  (no pre–smoother), has a converge factor bounded above by  $\sqrt{1-\delta_1}$  with  $\delta_1\leq 1$ .

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Furthermore, if the following condition holds

$$\forall \mathbf{e}_{k} \exists \delta_{2} > 0 : \|S_{k}^{(1)} \mathbf{e}_{k}\|_{A}^{2} \leq \|\mathbf{e}_{k}\|_{A}^{2} - \delta_{2} \|\Pi_{k} S_{k}^{(1)} \mathbf{e}_{k}\|_{A}^{2},$$

independently of  $e_k$  and k, then the multigrid method based with  $\gamma=1$ ,  $\nu_1\geq 1$  and  $\nu_2=0$  (no post–smoother), has a converge factor bounded above by  $1/\sqrt{1-\delta_2}$ ,  $\delta_2\leq 1$ .

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Finally, if both estimate

$$\forall \mathbf{e}_{k} \,\exists \, \delta_{1} > 0 \, : \, \|S_{k}^{(2)} \mathbf{e}_{k}\|_{A}^{2} \leq \|\mathbf{e}_{k}\|_{A}^{2} - \delta_{1} \|\Pi_{k} \mathbf{e}_{k}\|_{A}^{2},$$

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holds, for a pre– and post–smoother an estimate of the convergence factor is given by  $\sqrt{1-\delta_1/1-\delta_2}$ .

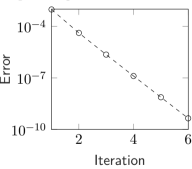
### **Concluding the example**

We test the simple *recursive* implementation from

</> E6-SimpleGMG/ex\_toepmultigrid.m

We consider the 1D Poisson problem with

- Jacobi smoother with optimal parameter,
- $nu_1 = \nu_2 = 2$  smoother steps,
- Use linear interpolation,
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Convergence result on finer meshes

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Size 127 Iteration 5 Time 5.61e-03
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#### Tests and extensions

The code contains other test problems with which you can play around. To get **better performances** you could re-implement the algorithm in a *non recursive way*.

### More general geometries

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• For structured matrices (Toeplitz, Circulant,  $\tau$ -algebra, etc.) we can discharge the problem on the properties of some functions describing the spectrum. Unfortunately, this is usually possible only when the matrix is obtained from the discretization of a PDE on a **structured** or **uniform** grid.

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- In general, our linear system could be coming from an optimization problem, being the Laplacian of a graph, being the discretization of a differential operator on an unstructured grid

### The way forward

We will reformulate the algorithm to use only purely algebraic properties of the matrix.

# Revisiting the components: *A*-convergent smoothers

To build the "source agnostic" Multigrid we start by revisiting the constitutive components.

### Theorem (A-convergent spliting)

Let A be SPD. Assume that for a given M the iteration matrix  $I-M^{-1}A$  has an A-norm less than one, or, equivalently that

$$\|I - A^{1/2}M^{-1}A^{1/2}\| < 1.$$

The symmetrization  $\overline{M} = M(M + M^T - A)^{-1}M^T$  satisfies

- (i)  $I \overline{M}^{-1}A = (I M^{-T}A)(I M^{-1}A)$ ,
- (ii)  $\overline{M} A$  is Symmetric Positive Semidefinite
- (iii)  $||I A^{1/2}M^{-1}A^{1/2}|| = ||I A^{1/2}\overline{M}^{-1}A^{1/2}||$ ,
- (iv)  $||I A^{1/2}M^{-1}A^{1/2}|| < 1 \Leftrightarrow M + M^T A \text{ SPD}$

Given  $A \in \mathbb{R}^{n \times n}$  SPD, we let J and P be two **rectangular matrices** with n rows so that we can consider the  $2 \times 2$ -block factorization:

$$A = \begin{bmatrix} \mathcal{A} & \mathcal{R} \\ \mathcal{L} & \mathcal{B} \end{bmatrix}, \quad \mathcal{A} = J^T A J, \quad \mathcal{B} = P^T A P.$$

We now suppose having two matrices  $\mathcal{M} \approx \mathcal{A}$  and  $\mathcal{D} \approx \mathcal{B}$  with  $\mathcal{D}$  SPD, where " $\approx$ " (usually) means that  $\mathcal{M}$  and  $\mathcal{D}$  are  $\mathcal{A}/\mathcal{B}$ -convergent splitting.

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Let  $\mathbf{r}_0 = \mathbf{b} - A\mathbf{u}_0$ ;

Use method  $\mathcal{M}$  for  $(J^TAJ)\mathbf{x} = J^T\mathbf{r}_0$ .

Compute the residual  $\mathbf{r}_m = b - A\mathbf{u}_m = b - A\mathbf{u}_0 - AJ\mathbf{x}_m = (I - AJ\mathcal{M}^{-1}J^T)\mathbf{r}_0$ ;

### **Algorithm 1:** Product iteration method

Given  $A \in \mathbb{R}^{n \times n}$  SPD, we let J and P be two **rectangular matrices** with n rows so that we can consider the  $2 \times 2$ -block factorization:

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### **Algorithm 2:** Product iteration method

Given  $A \in \mathbb{R}^{n \times n}$  SPD, we let J and P be two **rectangular matrices** with n rows so that we can consider the  $2 \times 2$ -block factorization:

$$A = \begin{bmatrix} \mathcal{A} & \mathcal{R} \\ \mathcal{L} & \mathcal{B} \end{bmatrix}, \quad \mathcal{A} = J^T A J, \quad \mathcal{B} = P^T A P.$$

We now suppose having two matrices  $\mathcal{M} \approx \mathcal{A}$  and  $\mathcal{D} \approx \mathcal{B}$  with  $\mathcal{D}$  SPD, where " $\approx$ " (usually) means that  $\mathcal{M}$  and  $\mathcal{D}$  are  $\mathcal{A}/\mathcal{B}$ -convergent splitting.

Let  $\mathbf{r}_0 = \mathbf{b} - A\mathbf{u}_0$ ;

Use method  $\mathcal{M}$  for  $(J^TAJ)\mathbf{x} = J^T\mathbf{r}_0$ ,;

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Use method  $\mathcal{D}$  for  $(P^TAP)\mathbf{w} = P^T\mathbf{r}_m$ ;

Compute the residual  $\mathbf{r}_w = \mathbf{b} - A\mathbf{u}_w = \mathbf{b} - A\mathbf{u}_m - AP\mathbf{w} = (I - AP\mathcal{D}^{-1}P^T)(I - AJ\mathcal{M}^{-1}J^T)\mathbf{r}_0;$ 

Use method  $\mathcal{M}$  for  $(J^T A J)\mathbf{x} = J^T \mathbf{r}_w$ ,;

The new residual is

$$\mathbf{r}_{\mathsf{new}} = \mathbf{b} - A\mathbf{u}_{\mathsf{new}} = \mathbf{b} - A\mathbf{u}_w - AJ\mathbf{x} = (I - AJ\mathcal{M}^{-T}J^T)(I - AP\mathcal{D}^{-1}P^T)(I - AJ\mathcal{M}^{-1}J^T)\mathbf{r}_0$$
**Algorithm 3:** Product iteration method

Given  $A \in \mathbb{R}^{n \times n}$  SPD, we let J and P be two **rectangular matrices** with n rows so that we can consider the  $2 \times 2$ -block factorization:

$$A = \begin{bmatrix} \mathcal{A} & \mathcal{R} \\ \mathcal{L} & \mathcal{B} \end{bmatrix}, \quad \mathcal{A} = J^T A J, \quad \mathcal{B} = P^T A P.$$

We now suppose having two matrices  $\mathcal{M} \approx \mathcal{A}$  and  $\mathcal{D} \approx \mathcal{B}$  with  $\mathcal{D}$  SPD, where " $\approx$ " (usually) means that  $\mathcal{M}$  and  $\mathcal{D}$  are  $\mathcal{A}/\mathcal{B}$ -convergent splitting.

#### Residual iteration matrix

The residual iteration  $E_r$  is therefore given by:

$$E_r = \mathbf{b} - A\mathbf{u}_{\mathsf{new}} = \mathbf{b} - A\mathbf{u}_{\mathsf{w}} - AJ\mathbf{x} = (I - AJ\mathcal{M}^{-T}J^T)(I - AP\mathcal{D}^{-1}P^T)(I - AJ\mathcal{M}^{-1}J^T),$$

thus 
$$\mathbf{u} - \mathbf{u}_0 = A^{-1}\mathbf{r}_0 \mapsto \mathbf{u} - \mathbf{u}_{\text{new}} = A^{-1}\mathbf{r}_{\text{new}}$$
 since  $AE = E_rA$ 

$$E = (I - JM^{-T}J^{T}A)(I - PD^{-1}P^{T}A)(I - JM^{-1}J^{T}A) = A^{-1}E_{r}A.$$

#### Lemma

If  $\mathcal{M}$  and  $\mathcal{D}$  are convergent smoother then  $\|E\mathbf{e}\|_A \leq \|\mathbf{e}\|_A$ .

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### Block-factorizations and product iteration methods

We implicitly define the product iteration method

$$I - B^{-1}A = (I - J\mathcal{M}^{-T}J^{T}A)(I - P\mathcal{D}^{-1}P^{T}A)(I - J\mathcal{M}^{-1}J^{T}A).$$

#### Lemma

If  $\mathcal{M}$  and  $\mathcal{D}$  are convergent smoother then  $\|E\mathbf{e}\|_A < \|\mathbf{e}\|_A$ .

### Block-factorizations and product iteration methods

We implicitly define the product iteration method

$$I - B^{-1}A = (I - J\mathcal{M}^{-T}J^{T}A)(I - P\mathcal{D}^{-1}P^{T}A)(I - J\mathcal{M}^{-1}J^{T}A).$$

#### Theorem

Let  $\overline{\mathcal{M}} = \mathcal{M}(\mathcal{M} + \mathcal{M}^T - \mathcal{A})^{-1}\mathcal{M}^T$ , given the following block-factored matrix

$$\hat{\mathcal{B}} = \begin{bmatrix} \mathcal{M} & O \\ P^T \mathcal{A} J & I \end{bmatrix} \begin{bmatrix} (\mathcal{M} + \mathcal{M}^T - \mathcal{A})^{-1} & O \\ O & \mathcal{D} \end{bmatrix} \begin{bmatrix} \mathcal{M}^T & J^T A P \\ O & I \end{bmatrix}$$

we express explicitly the operator as

$$B^{-1} = [J, P]\hat{\mathcal{B}}[J, P]^T = J\overline{\mathcal{M}}^{-1}J^T + (I - J\overline{\mathcal{M}}^{-T}J^TA)P\mathcal{D}^{-1}P^T(I - \mathcal{A}J\mathcal{M}^{-1}J^T).$$

What did we just prove?

- 1. We can build a block-factorization preconditioner  $B^{-1}$  as  $[J,P]\hat{\mathcal{B}}[J,P]^T$ ,
- 2. The matrix  $\hat{\mathcal{B}}$  is obtained from the approximate block-factorization of the two-by-two block matrix  $\hat{\mathcal{A}} = [J, P]^T A[J, P]$ ,
- 3. The stationary matrix iteration  $I B^{-1}A$  can be expressed as the product

$$(I - J\mathcal{M}^{-T}J^{T}A)(I - P\mathcal{D}^{-1}P^{T}A)(I - J\mathcal{M}^{-1}J^{T}A),$$

that act on Range(J), Range(P), and Range(J).

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that act on Range(J), Range(P), and Range(J).

We have written, using block factorization, a **method of the type we saw in our geometric example** on the 1D Laplacian. The **high** and **low frequency** spaces are then represented as the **images** and **kernels** of the J and P maps.

# Two-grid preconditioner

For J = I then  $[J, P]^T$  has full column rank since  $[I, P][I, P]^T = I + PP^T$  is SPD.

### Two-grid preconditioner

Given a smoother M for A and an interpolation matrix P, and let  $\mathcal{D}$  be an SPD approximation to  $\mathcal{B} = P^T A P$ , such that

- $M + M^T A$  is SPD (equivalently,  $||I A^{1/2}M^{-1}A^{1/2}|| < 1$ ).
- $\mathcal{D} \mathcal{B}$  is SPD.

Then, given the block matrix

$$\hat{\mathcal{B}} = \begin{bmatrix} M & O \\ P^T A & I \end{bmatrix} \begin{bmatrix} (M^T + M - A)^{-1} & O \\ O & \mathcal{D} \end{bmatrix} \begin{bmatrix} M^T & AP \\ O & I \end{bmatrix}$$

we define the preconditioner  $B_{TG}^{-1} = [I, P]\hat{\mathcal{B}}[I, P]^T$  or, equivalently,

$$B_{TG}^{-1} = \overline{M}^{-1} + (I - AM^{-1})^T P \mathcal{D}^{-1} P^T (I - M^{-1}A).$$

What can we say about the convergence properties of such method?

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### Convergence constant

We would like to estimate the best constant

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#### Theorem

Assume that J and P are such that any vector  $\mathbf{v}$  can be decomposed as  $\mathbf{v} = J\mathbf{w} + P\mathbf{x}$ . We introduce the projectors  $\pi_A = P\mathcal{B}^{-1}P^TA = P(P^TAP)^{-1}P^T$ , and let  $\widetilde{\mathcal{M}} = \mathcal{M}^T(\mathcal{M} + \mathcal{M}^T - A)^{-1}\mathcal{M}$ . The best constant K is given by

$$K = \sup_{\mathbf{v} \in \text{Range}(I - \pi_A)} \inf_{\mathbf{w} : \mathbf{v} = (I - \pi_A)J\mathbf{w}} \frac{\mathbf{w}^T \widetilde{\mathcal{M}} \mathbf{w}}{\mathbf{v}^T A \mathbf{v}} = \sup_{\mathbf{v}} \inf_{\mathbf{w} : \mathbf{v} = (I - \pi_A)J\mathbf{w}} \frac{\mathbf{w}^T \widetilde{\mathcal{M}} \mathbf{w}}{\mathbf{v}^T A (I - \pi_A) \mathbf{v}}.$$

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#### Theorem

Assume that J and P are such that any vector  $\mathbf{v}$  can be decomposed as  $\mathbf{v} = J\mathbf{w} + P\mathbf{x}$  with [J,P] invertible. We introduce the projectors  $\pi_A = P\mathcal{B}^{-1}P^TA = P(P^TAP)^{-1}P^T$ , and let  $\widetilde{\mathcal{M}} = \mathcal{M}^T(\mathcal{M} + \mathcal{M}^T - A)^{-1}\mathcal{M}$ . The best constant K is given by

$$K = \sup_{\mathbf{w}} \frac{\mathbf{w}^T \widetilde{\mathcal{M}} \mathbf{w}}{\mathbf{w}^T J^T A (I - \pi_A) J \mathbf{w}}.$$

For the **two-grid case** we have J = I,  $\mathcal{M} = M$  a smoother for A and  $\mathcal{D} = \mathcal{B} = P^T A P$ .

$$K_{TG} = \sup_{\mathbf{v}} \inf_{\mathbf{w} : \mathbf{v} = (I - \pi_A)\mathbf{w}} \frac{\mathbf{w}^T \widetilde{M} \mathbf{w}}{\mathbf{v}^T A \mathbf{v}}$$

For the **two-grid case** we have J = I,  $\mathcal{M} = M$  a *smoother* for A and  $\mathcal{D} = \mathcal{B} = P^T A P$ . We apply the previous theorem and find:

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$$= \sup_{\mathbf{v}} \frac{\inf_{\mathbf{w}} \left[ (\pi_A \mathbf{w} + (I - \pi_A) \mathbf{v})^T \widetilde{M} (\pi_A \mathbf{w} + (I - \pi_A) \mathbf{v}) \right]}{((I - \pi_A) \mathbf{v})^T A ((I - \pi_A) \mathbf{v})}$$

• We introduce  $\pi_{\widetilde{M}} = P\widetilde{M}_c^{-1}P^T\widetilde{M} = P(P^T\widetilde{M}P)^{-1}P^T\widetilde{M}$ ,

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$$K_{TG} = \sup_{\mathbf{v}} \inf_{\mathbf{w}: \mathbf{v} = (I - \pi_A)\mathbf{w}} \frac{\mathbf{w}^T M \mathbf{w}}{\mathbf{v}^T A \mathbf{v}}$$

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$$K_{TG} = \sup_{\mathbf{v}} \inf_{\mathbf{w}: \mathbf{v} = (I - \pi_{A})\mathbf{w}} \frac{\mathbf{w}^{T} \widetilde{M} \mathbf{w}}{\mathbf{v}^{T} A \mathbf{v}}$$
$$= \sup_{\mathbf{v}} \frac{\left( (I - \pi_{\widetilde{M}}) \mathbf{v} \right)^{T} \widetilde{M} \left( (I - \pi_{\widetilde{M}}) \mathbf{v} \right)}{((I - \pi_{A}) \mathbf{v})^{T} A ((I - \pi_{A}) \mathbf{v})}$$

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$$\mathcal{K}_{TG} = \sup_{\mathbf{v} = (I - \pi_A)\mathbf{v}} \frac{\left( (I - \pi_{\widetilde{M}})\mathbf{v} \right)^T \widetilde{M} \left( (I - \pi_{\widetilde{M}})\mathbf{v} \right)}{\mathbf{v}^T A \mathbf{v}} \leq \sup_{\mathbf{v}} \frac{\mathbf{v}^T \widetilde{M} (I - \pi_{\widetilde{M}})\mathbf{v}}{\mathbf{v}^T A \mathbf{v}},$$

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- We introduce  $\pi_{\widetilde{M}} = P\widetilde{M}_c^{-1}P^T\widetilde{M} = P(P^T\widetilde{M}P)^{-1}P^T\widetilde{M}$ ,
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- $\mathbf{v}^T A \mathbf{v} \geq \mathbf{v}^T A (I \pi_A) \mathbf{v}$  and thus the opposite inequality holds:

$$\sup_{\mathbf{v}} \frac{\mathbf{v}^T \widetilde{M}(I - \pi_{\widetilde{M}}) \mathbf{v}}{\mathbf{v}^T A \mathbf{v}} \leq \sup_{\mathbf{v}} \frac{\mathbf{v}^T \widetilde{M}(I - \pi_{\widetilde{M}}) \mathbf{v}}{\mathbf{v}^T A (I - \pi_A) \mathbf{v}} = K_{TG}.$$

### The first take-home message

The Theorem we have just seen shows that

$$\rho(E_{TG}) = 1 - \frac{1}{K_{TG}}, \quad K_{TG} = \begin{cases} \sup_{\mathbf{v}} \frac{\mathbf{v}^T \widetilde{M} (I - \pi_{\widetilde{M}}) \mathbf{v}}{\mathbf{v}^T A \mathbf{v}}, \\ \sup_{\mathbf{v}} \frac{\|(I - PR) \mathbf{v}\|_{\widetilde{M}}^2}{\|\mathbf{v}\|_A^2}, \end{cases} \qquad \pi_{\widetilde{M}} = P(P^T \widetilde{M} P)^{-1} P^T \widetilde{M},$$

for 
$$\widetilde{M} = M^T (M + M^T - A)^{-1} M$$
, and  $R = (P^T \widetilde{M} P)^{-1} P^T \widetilde{M}$ .

• Observe that RP is the identity on the coarse space.

Working with the symmetrized smoother  $\widetilde{M}$  is useful for proving estimates, but not so much for estimating constants.

### Corollary

Let  $\widetilde{M}$  be spectrally equivalent to an SPD matrix D, i.e., such that

$$\exists c_1, c_2 > 0 : c_1 \mathbf{v}^T D \mathbf{v} \leq \mathbf{v}^T \widetilde{\mathcal{M}} \mathbf{v} \leq c_2 \mathbf{v}^T D \mathbf{v} \quad \forall v.$$

Then, with  $\pi_D = P(P^TDP)^{-1}P^TD$  the following estimate for  $K_{TG}$  holds

$$c_1 \sup_{\mathbf{v}} \frac{\mathbf{v}^T D(I - \pi_D) \mathbf{v}}{\mathbf{v}^T A \mathbf{v}} \leq K_{TG} \leq c_2 \sup_{\mathbf{v}} \frac{\mathbf{v}^T D(I - \pi_D) \mathbf{v}}{\mathbf{v}^T A \mathbf{v}}.$$

### Example

If M is SPD and such that M-A is positive semidefinite,  $\widetilde{M}=M(2M-A)^{-1}M$  is spectrally equivalent to M such that

$$\frac{1}{2}\mathbf{v}^{\mathsf{T}} M \mathbf{v} \le \mathbf{v}^{\mathsf{T}} \widetilde{M} \mathbf{v} \le \mathbf{v}^{\mathsf{T}} M \mathbf{v},$$

thus  $c_1 = 1/2$ , and  $c_2 = 1$ .

### Example

If M is the Gauss-Seidel iteration matrix, i.e., M=D-L, then  $\widetilde{M}=(D-U)D^{-1}(D-L)$  is spectrally equivalent to D with

$$\frac{1}{4}\mathbf{v}^{\mathsf{T}} \mathsf{M} \mathbf{v} \le \mathbf{v}^{\mathsf{T}} \widetilde{\mathsf{M}} \mathbf{v} \le \kappa^2 \mathbf{v}^{\mathsf{T}} \mathsf{M} \mathbf{v},$$

thus  $c_1 = 1/4$ , and  $c_2 = \kappa$  the maximum number of nonzero entries of A per row.

We know how to estimate these quantities for both Jacobi and Gauss-Seidel type methods.

### Increasing the number of levels

We have described the two-grid method as

$$\hat{\mathcal{B}} = \begin{bmatrix} M & O \\ P^T A & I \end{bmatrix} \begin{bmatrix} (M^T + M - A)^{-1} & O \\ O & A_c \end{bmatrix} \begin{bmatrix} M^T & AP \\ O & I \end{bmatrix}, \quad \mathcal{D} = A_c = P^T A P,$$

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- For the following discussion is better to represent it by having the block triangular matrix with unit diagonal,
- Assume that we have  $\ell \geq 1$  levels and define
  - $A_0 = A$ ,
  - $P_k: V_{k+1} \equiv \mathbb{R}^{n_{k+1}} \mapsto V_k \equiv \mathbb{R}^{n_k}$  interpolation matrix  $PV_{k+1} \subset V_k$ ,
  - $A_{k+1} = P_k^T A_k P_k$  coarse-grid k+1 matrix,
  - $M_k$  a convergent smoother for  $A_k$ , i.e.,  $\|I A_k^{1/2} M_k^{-1} A_k^{1/2}\| < 1$ .

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$$\overline{B}_k = \begin{bmatrix} I & O \\ P_k^T A_k M_k^{-1} & I \end{bmatrix} \begin{bmatrix} M_k (M_k^T + M_k - A_k)^{-1} M_k^T & O \\ O & B_{k+1} \end{bmatrix} \begin{bmatrix} I & M_k^{-T} A_k P_k \\ O & I \end{bmatrix}$$

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  - $M_k$  a convergent smoother for  $A_k$ , i.e.,  $\|I A_k^{1/2} M_k^{-1} A_k^{1/2}\| < 1$ .
- With this ingredient we define a MG as a recursive  $2 \times 2$  block-factorization preconditioner  $B_{\nu}^{-1} = [I, P_k] \overline{B}_{k}^{-1} [I, P_k]^T$ .

```
At the coarsest level set B_\ell = A_\ell, the action of B_k^{-1}\mathbf{r} is given by;

Solve for M_k\mathbf{x}_k = \mathbf{r};  /* Presmooth */

Compute the residual \mathbf{d} = \mathbf{r} - A_k\mathbf{x}_k = (I - A_kM_k^{-1})\mathbf{r};

Compute \mathbf{x}_{k+1} = B_{k+1}^{-1}P_k^T(I - A_kM_k^{-1})\mathbf{r};  /* Coarse grid correction */

Update \mathbf{x}_k = \mathbf{x}_k + P\mathbf{x}_{k+1} = M_k^{-1}\mathbf{r} + P_kB_{k+1}^{-1}P_k^T(I - A_kM_k^{-1})\mathbf{r};

Solve for M_k^T\mathbf{y} = \mathbf{r} - A_k\mathbf{x}_k;  /* Postsmooth */

Set B_k^{-1}\mathbf{r} = \mathbf{x}_k + \mathbf{y}.

Algorithm 4: Generic MG Algorithm
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Algorithm 5: Generic MG Algorithm
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That is:

$$B_k^{-1}\mathbf{r} = (M_k^{-1} + M_k^{-T} - M_k^{-1}A_kM_k^{-T} + (I - M_k^{-T}A_k)P_kB_{k+1}^{-1}P_k^T(I - A_kM_k^{-1}))\mathbf{r}$$

```
At the coarsest level set B_\ell = A_\ell, the action of B_k^{-1} \mathbf{r} is given by; Solve for M_k \mathbf{x}_k = \mathbf{r}; /* Presmooth */ Compute the residual \mathbf{d} = \mathbf{r} - A_k \mathbf{x}_k = (I - A_k M_k^{-1}) \mathbf{r}; Compute \mathbf{x}_{k+1} = B_{k+1}^{-1} P_k^T (I - A_k M_k^{-1}) \mathbf{r}; /* Coarse grid correction */ Update \mathbf{x}_k = \mathbf{x}_k + P \mathbf{x}_{k+1} = M_k^{-1} \mathbf{r} + P_k B_{k+1}^{-1} P_k^T (I - A_k M_k^{-1}) \mathbf{r}; Solve for M_k^T \mathbf{y} = \mathbf{r} - A_k \mathbf{x}_k; /* Postsmooth */ Set B_k^{-1} \mathbf{r} = \mathbf{x}_k + \mathbf{y}.
```

### Algorithm 6: Generic MG Algorithm

That is:

$$B_k^{-1} = \overline{M}_k^{-1} + (I - M_k^{-T} A_k) P_k B_{k+1}^{-1} P_k^{T} (I - A_k M_k^{-1}),$$

for  $\overline{M}_k$  the symmetrized smoother and  $B_\ell = A_\ell$ .

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Solve for M_k^T\mathbf{y} = \mathbf{r} - A_k\mathbf{x}_k; /* Postsmooth */

Set B_k^{-1}\mathbf{r} = \mathbf{x}_k + \mathbf{y}.
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## Algorithm 7: Generic MG Algorithm

That is:

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for  $\overline{M}_k$  the symmetrized smoother and  $B_\ell = A_\ell$ .

### **Definition**

We call this method the symmetric V(1,1)-cycle Multigrid.

## Proposition

Under the assumption that the smoothers  $M_k$  are convergent in the  $A_k$ -norm, the symmetric V(1,1)-cycle Multigrid preconditioner  $B_k$  is such that  $B_k-A_k$  is symmetric positive semidefinite.

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## **9** Idea

We have just rebuilt the construction without investigating the "high" and "low frequency" ideas. This will be our next target.

# **Stable decompositions**

Let  $\overline{V}_k = \text{Range}(P_0, \dots, P_{k-1})$  be the kth-level coarse space.

## Stability

We say that a decomposition

$$\mathbf{v} = \sum_k \overline{\mathbf{v}}_k^f \qquad \overline{\mathbf{v}}_k^f \in \overline{V}_k,$$

is *stable* if there exists a level independent constant  $\sigma > 0$  such that

$$\sum_{k} (\overline{\mathbf{v}}_{k}^{f})^{T} A \overline{\mathbf{v}}_{k} \equiv \sum_{k} (\overline{\mathbf{v}}_{k}^{f})^{T} A_{k} \overline{\mathbf{v}}_{k}^{f} \leq \sigma \mathbf{v}^{T} A \mathbf{v}.$$

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### Complementary space

For a space  $V_j$  we define the subspace  $V_j^f \subset V_j$  complementary to the coarse space  $P_j V_{j+1}$ .

## The idea from the simple Poisson case

We select  $V_i^f$  so that the symmetrized smoother  $\overline{M}_i$  is efficient when restricted to  $V_i^f$ , i.e.,

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## Vector decomposition

$$\mathbf{v}_j = \mathbf{v}_j^f + P_j \mathbf{v}_{j+1} = [I, P_k] \begin{bmatrix} \mathbf{v}_k^f \\ \mathbf{v}_{k+1} \end{bmatrix} \text{ with } \mathbf{v}_j^f \in V_j^f \subset V_j, \ \mathbf{v}_{j+1} \in V_{j+1}, \ j = k, k+1, \dots, \ell-1.$$

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 with  $\mathbf{v}_j^f \in V_j^f \subset V_j$ ,  $\mathbf{v}_{j+1} \in V_{j+1}$ ,  $j = k, k+1, \ldots, \ell-1$ .

$$B_k^{-1} = [I, P_k] \overline{B}_k^{-1} [I, P_k]^T, \quad I = GG^T = (B_k^{1/2} [I, P_k] \overline{B}_k^{-1/2}) (B_k^{1/2} [I, P_k] \overline{B}_k^{-1/2})^T,$$

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$$\begin{split} \boldsymbol{B}_k^{-1} &= [I, P_k] \overline{\boldsymbol{B}}_k^{-1} [I, P_k]^T, \quad I = \boldsymbol{G} \boldsymbol{G}^T = (\boldsymbol{B}_k^{1/2} [I, P_k] \overline{\boldsymbol{B}}_k^{-1/2}) (\boldsymbol{B}_k^{1/2} [I, P_k] \overline{\boldsymbol{B}}_k^{-1/2})^T, \\ &\Rightarrow \|\boldsymbol{G}\|_2 < 1 \text{ and } \begin{bmatrix} \mathbf{v}_k^f \\ \mathbf{v}_{k+1} \end{bmatrix}^T [I, P_k] \boldsymbol{B}_k [I, P_k]^T \begin{bmatrix} \mathbf{v}_k^f \\ \mathbf{v}_{k+1} \end{bmatrix} \leq \begin{bmatrix} \mathbf{v}_k^f \\ \mathbf{v}_{k+1} \end{bmatrix}^T \overline{\boldsymbol{B}}_k \begin{bmatrix} \mathbf{v}_k^f \\ \mathbf{v}_{k+1} \end{bmatrix}. \end{split}$$

By using the definition of  $\overline{B}_k$  we can estimate

$$0 \leq \mathbf{v}_{k}^{T} (B_{k} - A_{k}) \mathbf{v}_{k}$$

$$\leq \sum_{j=k}^{\ell-1} \left[ \left( M_{j}^{T} \mathbf{v}_{j}^{f} + A_{j} P_{j} \mathbf{v}_{j+1} \right)^{T} (M_{j} + M_{j}^{T} - A_{j})^{-1} \left( M_{j}^{T} \mathbf{v}_{j}^{f} + A_{j} P_{j} \mathbf{v}_{j+1} \right) \right]$$

$$+ \mathbf{v}_{\ell}^{T} A_{\ell} \mathbf{v}_{\ell} - \mathbf{v}_{k}^{T} A_{k} \mathbf{v}_{k}.$$

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2. 
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3. 
$$\mathbf{v}_{\ell}^{\mathsf{T}} A_{\ell} \mathbf{v}_{\ell} \leq \sigma_{c} \mathbf{v}_{k}^{\mathsf{T}} A_{k} \mathbf{v}_{k}$$
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$$+ \mathbf{v}_{\ell}^{T} A_{\ell} \mathbf{v}_{\ell} - \mathbf{v}_{k}^{T} A_{k} \mathbf{v}_{k} \leq (\sigma_{c} + 2(\sigma + \mu) - 1) \mathbf{v}_{k}^{T} A_{k} \mathbf{v}_{k}.$$

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,

3. 
$$\mathbf{v}_{\ell}^{\mathsf{T}} A_{\ell} \mathbf{v}_{\ell} \leq \sigma_{c} \mathbf{v}_{k}^{\mathsf{T}} A_{k} \mathbf{v}_{k}$$
.

## Theorem for the optimal choice

## Theorem (Vassilevski 2008, Theorem 5.7)

Given  $A_j$ -convergent smoother  $M_j$ ,  $j=0,\ldots,\ell-1$  for the V(1,1)-cycle MG preconditioner (with  $B=B_0$  and  $A=A_0$ ). If any fine-grid vector  $\mathbf{v}=\mathbf{v}_0$  allows for a decomposition of the form  $\mathbf{v}_j^f=\mathbf{v}_j-P_j\mathbf{v}_{j+1}$ ,  $j=0,1,\ldots,\ell-1$ , such that

- A1 Stable decomposition:  $\sum_{i} (\mathbf{v}_{i}^{f})^{T} \overline{M}_{k} \mathbf{v}_{i}^{f} \leq \sigma \mathbf{v}^{T} A \mathbf{v}$ ,
- A2 Smoother scaling:  $(1 + \delta)\mathbf{v}_j^T A \mathbf{v}_j \leq \mathbf{v}_j^T (M_j^T + M_j) \mathbf{v}_j = 2\mathbf{v}_j^T M_j \mathbf{v}_j$ ,
- A3 Stable coarse component:  $\mathbf{v}_{\ell}^T A_{\ell} \mathbf{v}_{\ell} \leq \sigma_c \mathbf{v}^T A \mathbf{v}$ ,
- A4 Efficiency of the smoothers on the components of  $A_i P_i \mathbf{v}_{i+1}$  so that

$$\sum_{j} \mathbf{v}_{j+1}^{\mathsf{T}} P_{j}^{\mathsf{T}} A_{j} (M_{j} + M_{j}^{\mathsf{T}} - A_{j})^{-1} A_{j} P_{j} \mathbf{v}_{j+1} \leq \mu \mathbf{v}^{\mathsf{T}} A \mathbf{v}^{\mathsf{T}}.$$

Then, the MG preconditioner B is uniformly spectrally equivalent to A:

$$\mathbf{v}^T A \mathbf{v} \leq \mathbf{v}^T B \mathbf{v} \leq (\sigma_c + 2(\sigma + \mu) - 1) \mathbf{v}^T A \mathbf{v}.$$

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A comes from a FEM discretization of a PDE and we can use Sobolev space and grid properties to attain **stable decompositions**.



We forget about the source of A and try to build a black-box approach that enforces the needed condition

We are at a crossroad

# The algebraic idea

Given Matrix  $A \in \mathbb{R}^{n \times n}$  SPD

Wanted Iterative method *B* to precondition the CG method:

• Hierarchy of systems

$$A_I \mathbf{x}_= \mathbf{b}_I, I = 0, \dots, \ell$$

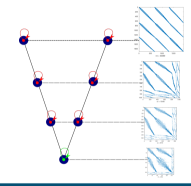
• Transfer operators:

$$P'_{l+1}: \mathbb{R}^{n_{l+1}} \to \mathbb{R}^{n_l}$$

Missing Structural/geometric infos

## Smoother: "High frequencies"

$$M_l: \mathbb{R}^{n_l} \to \mathbb{R}^{n_l}$$



Prolongator: "Low frequencies"

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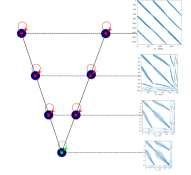
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Complementarity of Smoother and Prolongator

Let us assume that P has the form:

$$P = \begin{bmatrix} W \\ I \end{bmatrix}$$

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- $\Rightarrow \psi_{i_c} = P \delta_{i_c}$ ,  $i_c = 1, \ldots, n_c$  is a basis for the range of  $\pi_{\widetilde{M}}$ :  $\pi_{\widetilde{M}} \psi_{i_c} = \delta_{i_c}$ .

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$$K_{TG} = \sup_{\mathbf{v}} \frac{\mathbf{v}^T \widetilde{M} (I - \pi_{\widetilde{M}}) \mathbf{v}}{\mathbf{v}^T A \mathbf{v}} = \sup_{\mathbf{v}} \frac{\begin{bmatrix} \overline{\mathbf{v}}_f \\ \mathbf{0} \end{bmatrix}' \widetilde{M} (I - \pi_{\widetilde{M}}) \begin{bmatrix} \overline{\mathbf{v}}_f \\ \mathbf{0} \end{bmatrix}}{\mathbf{v}^T A \mathbf{v}}$$

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$$= \sup_{\overline{\mathbf{v}}_{f}} \sup_{\mathbf{v}_{c}} \sup_{t \in \mathbb{R}} \frac{\begin{bmatrix} \overline{\mathbf{v}}_{f}} \\ \mathbf{0} \end{bmatrix}^{T} \widetilde{M} (I - \pi_{\widetilde{M}}) \begin{bmatrix} \overline{\mathbf{v}}_{f} \\ \mathbf{0} \end{bmatrix}}{\begin{pmatrix} \begin{bmatrix} \overline{\mathbf{v}}_{f}} \\ \mathbf{0} \end{bmatrix} + t P \mathbf{v}_{c} \end{pmatrix}^{T} A \begin{pmatrix} \begin{bmatrix} \overline{\mathbf{v}}_{f}} \\ \mathbf{0} \end{bmatrix} + t P \mathbf{v}_{c} \end{pmatrix}$$

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# **Optimal prolongation**

Let us assume that *P* has the form:

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 $\Rightarrow$  Select W such that  $A_{ff}W + A_{fc} = 0$ 

### **Optimal prolongation**

Let us assume that *P* has the form:

$$P = \begin{bmatrix} W \\ I \end{bmatrix} = \begin{bmatrix} -A_{ff}^{-1}A_{fc} \\ I \end{bmatrix} \qquad A = \begin{bmatrix} A_{ff} & A_{fc} \\ A_{cf} & A_{cc} \end{bmatrix}$$

and to optimize the bound on  $K_{TG}$  we want

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Since it gives us

$$K_{TG} \leq \sup_{\overline{\mathbf{v}}_f} \frac{\begin{bmatrix} \overline{\mathbf{v}}_f \\ \mathbf{0} \end{bmatrix}^T \widetilde{M} \begin{bmatrix} \overline{\mathbf{v}}_f \\ \mathbf{0} \end{bmatrix}}{\begin{bmatrix} \overline{\mathbf{v}}_f \\ \mathbf{0} \end{bmatrix}^T A \begin{bmatrix} \overline{\mathbf{v}}_f \\ \mathbf{0} \end{bmatrix}} = \sup_{\overline{\mathbf{v}}_f} \frac{\overline{\mathbf{v}}_f^T \widetilde{M}_{ff} \overline{\mathbf{v}}_f}{\overline{\mathbf{v}}_f^T A_{ff}} = \frac{1}{\lambda_{\min}(\widetilde{M}_{ff}^{-1} A_{ff})}.$$

### **Selecting** *c* and *f* nodes

#### = The second take-home message

A reasonable guideline to construct P is to find for any coarse unit vector  $\delta_{i_c}$  in  $\mathbb{R}^{n_c}$ , an approximate solution to

$$A_{ff}\mathbf{w}_{i_c}=-A_{fc}\boldsymbol{\delta}_{i_c}$$

and build the  $i_c$ th column of P as  $\psi_{i_c} = [\mathbf{w}_{i_c}^T, \boldsymbol{\delta}_{i_c}^T]^T$ .

- We cannot solve exactly the systems for the w both for sparsity and cost reasons,
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#### Here begins the fun

What differentiates the available AMG algorithms is the procedure for identifying the coarse space through a combination of \*heuristics, heuristics, h

#### 1

#### Assumption:

Geometrically smooth functions are in the near null space of  $\boldsymbol{A}$ .

Wlog assume A s.t.  $\lambda_{\max}(A)=1$ , and let e be a small normalized eigenmode of A, i.e.,

$$A\mathbf{e} = \lambda \mathbf{e}, \|\mathbf{e}\| = 1, \lambda \ll 1$$

Thus: 
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#### Heuristic

Smooth error varies slowly in the direction of relatively large (negative) coefficients of the matrix.

#### Strong dependence (Ruge and Stüben 1987)

For a chosen tolerance  $\theta \in (0,1]$ , we say that a dof i is strongly influenced by  $j \neq i$  if

$$-a_{i,j} \geq \max_{k \neq i} (-a_{k,i}).$$

#### Define:

- $W_i = \{j \in \Omega_i : j \text{ is weakly connected to } j\}$ ,
- $S_i = \{j \in \Omega_i : j \text{ is strongly connected to } j\}$ ,
- $C_i$  set of coarse points that are allowed to interpolate i.

The  $(i, i_c)$  entry of P is given by

$$-\frac{\left(a_{i,i_c} + \sum_{i_\chi \in S_i} a_{i,i_\chi} \frac{a_{i_\chi,i_c}}{\sum_{j_c \in C_i} a_{i_\chi,j_c}}\right)}{a_{ii} + \sum_{i_\chi \in W_i} a_{i,i_\chi}}$$

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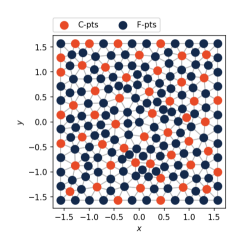
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Without delving into the details, the expression for the interpolation can be obtained by

- 1. Defining a strength matrix,  $A_s$ , obtained deleting the weak connections in A,
- First pass choosing an independent set of fine grid points based on the graph of A<sub>s</sub>,
- Second pass choosing additional points (if needed) to satisfy interpolation requirements.

To see the algorithm at work, we test it by means of the PyAMG library (Bell, Olson, and Schroder 2022) on a small problem, specifically we use it to highlight the division in Coarse and Fine dofs of a given grid.

You can run the example in **G** Google Colab by using the **Q** GitHub Gist https://bit.ly/3MToLtN.





#### Aggregation idea

The aggregation idea is using an algorithm that splits the set of vertices of the graph of A or of a re-weighted version of A (sometimes called filtered matrix) as a union of non-overlapping subsets - aggregates - each of which forms a connected sub-graph.

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#### The FEM case

For a FEM discretization of PDE on a set  $\Omega$  this corresponds to a partition of the domain:

$$\Omega = \bigcup_{i=1}^{J} \Omega_{j}, \quad \Omega_{i} \bigcap \Omega_{j} = \emptyset, i \neq j$$



**G-G** https://bit.ly/3vnAV82

Having selected aggregates

$$\{1,\ldots,n\}=\bigcup_{j=1}^J\mathcal{A}_j,\quad \mathcal{A}_i\bigcap\mathcal{A}_j=\emptyset, i\neq j,$$

The prolongator operator is then given simply by posing

$$P: \mathbb{R}^{n_c} \to \mathbb{R}^n, \quad (P\mathbf{x})_i \mapsto x_i, \quad i \in \mathcal{A}_i.$$

- Since the aggregates are mutually disjoint  $\forall i \in \{1, ..., n\}$  exist only one index  $j \in \{1, ..., n_c\}$  such that  $i \in C_i$ .
- $\P$  "the *j*th component of the vector  $\mathbf{x} \in \mathbb{R}^m$ ,  $m = |\mathcal{A}_j|$  will be mapped onto all components of the vector  $\mathbf{y} \in \mathbb{R}^n$  indices of which are in  $\mathcal{A}_i$ "
- P represents a piece-wise constant interpolation operator.

Usually *piece-wise constant* is not enough...

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• We get this Multigrid Hierarchy (that seems plausible)

Number of Levels: 7 Operator Complexity: 1.262 Grid Complexity: 1.188			
	Solver:	'pinv	
level	unknowns	nonzer	
0	250000	1248000	[79.24%]
1	41750	290584	[18.45%]
2	4704	32370	[2.06%]
3	532	3538	[0.22%]
4	70	424	[0.03%]
5	12	58	[0.00%]
6	3	9	[0.00%]

MultilevelSolver

Usually piece-wise constant is not enough...

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level	unknowns	nonzer	os
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MultilevelSolver

How can we make things better?

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- Use a procedure to **smooth out** the *basis function* induced by the aggregation procedure by using the smoother, which is, using few applications of smoothing on the prolongation matrix:

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That run for 12 iteration with last residual 8.620525e-09

#### Hierarchy for unsmoothed aggregation

MultilevelSolver

Number of Levels: 7

Operator Complexity: 1.262
Grid Complexity: 1.188
Coarse Solver: 'pinv'
level unknowns nonzeros

OB	HOHZCI	diffill will	TOVOT
[79.24%]	1248000	250000	0
[18.45%]	290584	41750	1
[2.06%]	32370	4704	2
[0.22%]	3538	532	3
[0.03%]	424	70	4
[0.00%]	58	12	5
[%00.0]	9	3	6

#### Hierarchy for smoothed aggregation

MultilevelSolver

1 0770 1

Number of Levels: 6

unlenarma

Operator Complexity: Grid Complexity:

Grid Complexity: 1.188
Coarse Solver: 'pinv'

1.337

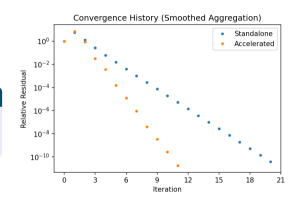
телет	unknowns	nonzer	ros
0	250000	1248000	[74.82%]
1	41750	373416	[22.39%]
2	4704	41554	[2.49%]
3	532	4526	[0.27%]

- Smoothed aggregation produces hierarchies with more nonzero entries,
- To reduce the *fill-in* filtering (dropping) strategies are usually implemented,

#### **Operator Complexity**

$$opc = \frac{\sum_{l=0}^{\ell-1} \operatorname{nnz}(A_l)}{\operatorname{nnz}(A_0)}$$

 AMG is more often used as preconditioner for the CG algorithm that as solver



You can run this example on Google Colab from https://bit.ly/30HYKPJ.



#### First comes the smoother

The approach called **compatible relaxation** consists in selecting a set of coarse degrees of freedom based solely on the smoother, the interpolation matrix is constructed later.

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From the definition of  $K_{TG}$  we have that if we find a matrix  $J_*$  such that

J1 Range
$$(J_*)$$
 = Range $(I - PR_*)$ ,  $R_* = (P^T \widetilde{M} P)^{-1} P^T \widetilde{M}$ 

we have the inequality:  $\mathbf{v}J_*^T\widetilde{M}J_*\mathbf{v} < K_{TG}\mathbf{v}^TJ^TAJ\mathbf{v}$ 



#### First comes the smoother

The approach called **compatible relaxation** consists in selecting a set of coarse degrees of freedom based solely on the smoother, the interpolation matrix is constructed later.

From the definition of  $K_{TG}$  we have that if we find a matrix  $J_*$  such that

J1 Range(
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) = Range( $I - PR_*$ ),  $R_* = (P^T \widetilde{M} P)^{-1} P^T \widetilde{M}$   
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#### 🖢 Idea

J picks a principal submatrix from A and  $\widetilde{M}$ , the inequality thus means that A has a principal submatrix that is spectrally equivalent to the same principal submatrix of  $\widetilde{M}$ .

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#### Heuristic

Fix R, then select J such that the constant  $K_{CR}$  in

$$\mathbf{v}^T J A J \mathbf{v} \leq \mathbf{v}^T J^T \widetilde{M} J \mathbf{v} \leq K_{CR} \mathbf{v}^T J^T A J \mathbf{v},$$

is close to 1.

### Compatible relaxation the idea

Let A be ans SPD matrix, M an A-convergent smoother, one can prove that

$$\|(I - \widetilde{M}^{-1}A)^m \mathbf{e}\|_A \le \frac{1}{\sqrt{m+1}} \|\mathbf{e}\|_{\widetilde{M}}$$
 (Smoothing property)

Take a **projection on the coarse space** Q being  $\widetilde{M}$ -orthogonal satisfying

$$\|(I-Q)\mathbf{e}\|_{\widetilde{M}} \leq \delta \|\mathbf{e}\|_{\mathcal{A}}$$
 (Approximation property)

Then for any e = (I - Q)e and any integer  $m \ge 1$  the following estimate holds

$$\|(I-\widetilde{M}^{-1}A)^m\mathbf{e}\|_{\widetilde{M}}\leq \frac{\delta}{\sqrt{1+m}}\|\mathbf{e}\|_{\widetilde{M}}.$$

# Compatible relaxation the algorithm

We apply our inequality for a solution of the homogeneous system  $A\mathbf{x}=\mathbf{0}$  Input:  $\mathbf{e}$  random initial iterate

$$m = 1;$$

Compute 
$$\mathbf{e}_0 = (I - Q)\mathbf{e}$$
;

Smooth 
$$e_m = (I - \widetilde{M}^{-1}A)e = (I - M^{-1}A)(I - M^{-T}A)e_{m-1};$$

if  $\|\mathbf{e}_m\|_{\widetilde{M}}/\|\mathbf{e}_0\|_{\widetilde{M}}$  is small then

The process has converged, convergence is now fast.;

#### else

Use  $\mathbf{e}_m$  to augment the coarse space, build a new Q and try again.

#### end

Since

$$\|(I-\widetilde{M}^{-1}A)^m\mathbf{e}\|_{\widetilde{M}}\leq \frac{\delta}{\sqrt{1+m}}\|\mathbf{e}\|_{\widetilde{M}},$$

an m large enough for which this procedure converge exists.

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As we have discussed implementing these methods efficiently requires some thought.

The good news is that there are several libraries that one can resort to.

hypre is a library of high performance preconditioners and solvers featuring multigrid methods for the solution of large, sparse linear systems of equations on massively parallel computers. AP https://github.com/hypre-space/hypre

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ML - Trilinos The Trilinos Project is an effort to develop algorithms and enabling technologies within an object-oriented software framework for the solution of large-scale, complex multi-physics engineering and scientific problems. An https://github.com/trilinos/Trilinos

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PSCToolkit parallel BLAS feature for sparse matrices that are capable of running on machines with thousands of high-performance cores, and construction of higher-level iterative solvers and preconditioners.

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Why all this interest in large and parallel?

Solve : 
$$A\mathbf{x} = \mathbf{b}$$
,

where

- $A \in \mathbb{R}^{n \times n}$  is a very large and sparse matrix nnz(A) = O(n),
- $\mathbf{x}, \mathbf{b} \in \mathbb{R}^n$ ,

But what does large mean?

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<sup>&</sup>quot;In a ground wire problem involving a large number of ground conductors, 13 simultaneous equations were solved..." – Dwight (1930)"

<sup>&</sup>quot;The second machine, now in operation, was designed for the direct solution of nine or fewer simultaneous equations." – Wilbur, J. B. (1936)

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<sup>&</sup>quot;Finally, though the labour of relaxation in three dimensions is prohibitively great, the future use of the new electronic calculating machines in this connexion is a distinct possibility" – Fox, L. (1947)

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"The Ferranti PEGASUS computer, with a main store of 4096 words, can solve a maximum of 86 simultaneous equations by its standard subroutine and takes about 45 minutes to complete this calculation."

- Wilson, L. B. (1959)

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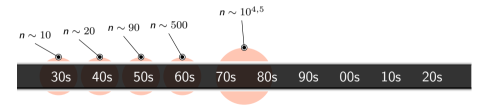
"...the bound imposed by this is  $m + n \le 474$ . In addition, this number of equations would fill one standard (1.800ft) reel of magnetic tape, and the fifty-odd hours taken in the calculation might be thought excessive."

- Barron, Swinnerton-Dyer (1960)

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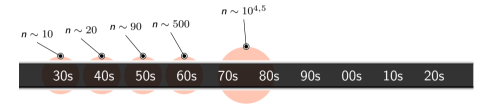


"...handling problems involving sets of simultaneous equations of two-thousandth order, and SAMIS available through "Cosmic" at the University of Georgia, which can treat up to 10,000 simultaneous equations." – Melosh, Schmele (1969)

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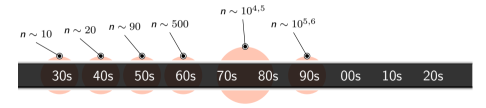


"The mini-computer cost algorithm is applied to the same complex shell problem used previously, with 9120 degrees of freedom [...]. The running times, however, are 40 and 70 hr, respectively! It would appear that improvement of mini-computer speeds is required..." – Kamel, McCabe (1978)

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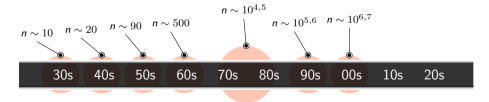


"For instance, Pomerell in 1994 reports on successful application of preconditioned Krylov methods for very ill-conditioned unstructured finite element systems of order up to 210,000 that arise in semiconductor device modeling." – Saad Y., van der Vorst, H.A. (2000)

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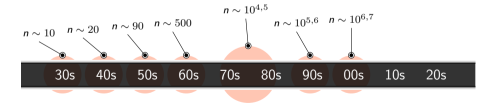


"As a second example, we show results (Table VIII) for a problem arising in ocean modeling (barotropic equation) with n = 370,000 unknowns and approximately 3.3 million nonzero entries..." – Benzi, M. (2002)

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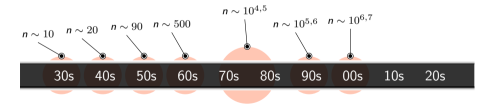


"Problem: Large, mesh size:  $180 \times 60 \times 30$ ,  $\sharp$  unknowns (in simulation): 1,010,160, Solution time 45.7 h" – Wang, de Sturler, Paulino (2006)

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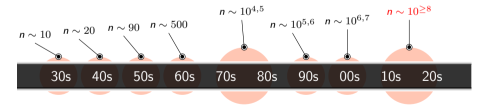


"The parallel GMRES was tested on the Tesla T10P GPU using a set of matrix data from the oil field simulation data of Conoco Phillips. The order of the system ranges from  $\sim 2000$  to  $\sim 1.1$  million." – M. Wang, H. Klie, M. Parashar, H. Sudan (2009)

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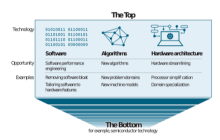
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The exascale challenge, using computer that do  $10^{15}$  Flops, targeting next-gen systems doing  $10^{18}$  Flops to solve problems with tens of billions of unknowns.

## The philosophy behind the effort



(Leiserson et al. 2020)

"As miniaturization wanes, the silicon-fabrication improvements at the Bottom will no longer provide the predictable. broad-based gains in computer performance that society has enjoyed for more than 50 years. Software performance engineering, development of algorithms, and hardware streamlining at the Top can continue to make computer applications faster in the post-Moore era."

## Where we want to solve it<sup>1</sup>

	System	Cores	Rmax (TFlops/s)
1	Fugaku	7,630,848	442,010.0
2	Summit	2,414,592	148,600.0
3	Sierra	1,572,480	94,640.0
:	:	:	:
18	Marconi-100	347,776	21,640.0
:	:	:	:
20	Piz Daint	387,872	21,230.0
:	:	:	:
74	MareNostrum	153,216	6,470.8



Marconi-100 - CINECA



Piz Daint - CSCS

- Machines with hundreds of MPI cores,
- Hybrid form of parallelism: MPI, OpenMP, CUDA/OpenCL, ...

<sup>&</sup>lt;sup>1</sup>TOP500 list, November 2021 - https://www.top500.org

### What do we ask to it?

Solve the preconditioned system:

$$B^{-1}Ax = B^{-1}b,$$

with matrix  $B^{-1} \approx A^{-1}$  (left preconditioner) such that:

Algorithmic scalability  $\max_i \lambda_i(B^{-1}A) \approx 1$  being independent of n (all the work we did on the K constant!),

Linear complexity the action of  $B^{-1}$  costs as little as possible, the best being  $\mathcal{O}(n)$  flops,

Implementation scalability in a massively parallel computer,  $B^{-1}$  should be composed of local actions, performance should depend linearly on the number of processors employed.

## PSCToolkit - psctoolkit.github.io

### Two central libraries PSBLAS and AMG4PSBLAS:

- Existing software standards:
  - MPI, OpenMP, CUDA
  - Serial sparse BLAS,

- (Par)Metis,
- AMD
- Attention to performance using modern Fortran;
- Research on new preconditioners;
- No need to delve in the data structures for the user;
- Tools for error and mesh handling beyond simple algebraic operations;
- Standard Krylov solvers





## PSCToolkit - psctoolkit.github.io

### Two central libraries PSBLAS and AMG4PSBLAS:

- Domain decomposition preconditioners
- Algebraic multigrid with aggregation schemes
  - Parallel coupled weighted matching based aggregation
  - Parallel decoupled smoothed aggregation (Vaněk, Mandel, Brezina)
- Parallel Smoothers (Block-Jacobi, DD-Schwartz, Hybrid-GS/SGS/FBGS,  $\ell_1$  variants) that can be coupled with specialized block (approximate) solvers MUMPS, SuperLU, incomplete factorizations ((H)AINV, (H)INVK/L, (H)ILU-type)
- V-Cycle, W-Cycle, K-Cycle





## PSCToolkit - psctoolkit.github.io

Two central libraries PSBLAS and AMG4PSBLAS.

• Freely available from: https://psctoolkit.github.io,

Open Source with BSD 3 Clause License.

People involved: S. Filippone, P. D'Ambra, F. Durastante.

**Contributors:** Soren Rasmussen, Zaak Beekman, Ambra Abdullahi Hassan, Alfredo Buttari, Daniela di Serafino, Michele Martone, Michele Colajanni, Fabio Cerioni, Stefano Maiolatesi, Dario Pascucci





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- The coarse solver is again a preconditioned CG method.

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- $\ell_1$ -HGS On process  $p=1,\ldots,np$  relative to the index set  $\Omega_p$  we factorize  $A_{pp}=L_{pp}+D_{pp}+L_{pp}^T$  for  $D_{pp}=\mathrm{diag}(A_{pp})$  and  $L_{pp}=\mathrm{trilu}(A_{pp})$  and select:

$$\begin{split} M_{\ell_1 - HGS} &= \mathrm{diag}((M_{\ell_1 - HGS})_p)_{p=1, \dots np}, \\ (M_{\ell_1 - HGS})_p &= L_{pp} + D_{pp} + D_{\ell_1 p}, \\ (d_{\ell_1})_{i=1}^{nb} &= \sum_{j \in \Omega_p^{nb}} |a_{ij}|. \end{split}$$

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AINV Block-Jacobi with an approximate inverse factorization on the block ⇒ suitable for GPU application!

• The prolongator P is built by dofs aggregation based on matching in the weighted (adjacency) graph of A.

Given  $\mathbf{w} \in \mathbb{R}^n$ , let  $P \in \mathbb{R}^{n \times n_c}$  and  $P_f \in \mathbb{R}^{n \times n_f}$  be a prolongator and a complementary prolongator, such that:

$$\mathbb{R}^{n} = \operatorname{Range}(P) \oplus^{\perp} \operatorname{Range}(P_f), \quad n = n_c + n_f$$

 $\mathbf{w} \in \text{Range}(P)$ : coarse space

 $Range(P_f)$ : complementary space

$$[P, P_f]^T A [P, P_f] = \begin{pmatrix} P^T A P & P^T A P_f \\ P_f^T A P & P_f^T A P_f \end{pmatrix} = \begin{pmatrix} A_c & A_{cf} \\ A_{fc} & A_f \end{pmatrix}$$

 $A_c$ : coarse matrix

 $A_f$ : hierarchical complement

### Sufficient condition for efficient coarsening

 $A_f = P_f^T A P_f$  as well conditioned as possible, i.e., Convergence rate of *compatible relaxation*:  $\rho_f = \|I - M_f^{-1} A_f\|_{A_f} \ll 1$ 

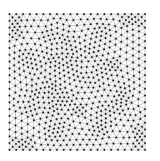
### But how we achieve it?

### Weighted graph matching

Given a graph  $G=(\mathcal{V},\mathcal{E})$  (with adjacency matrix A), and a weight vector  $\mathbf{w}$  we consider the weighted version of G obtained by considering the weight matrix  $\hat{A}$ :

$$(\hat{A})_{i,j} = \hat{a}_{i,j} = 1 - \frac{2a_{i,j}w_iw_j}{a_{i,i}w_i^2 + a_{j,j}w_j^2},$$

- a  $matching \mathcal{M}$  is a set of pairwise non-adjacent edges, containing no loops;
- a maximum product matching if it maximizes the product of the weights of the edges e<sub>i→j</sub> in it.



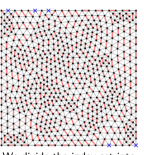
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We can formally define a prolongator.

$$P = \begin{bmatrix} \mathbf{w}_{e_{1}} & 0 & 0 \\ 0 & \ddots & 0 \\ 0 & 0 & \mathbf{w}_{e_{n_{p}}} \end{bmatrix} 2n_{p} \qquad \mathbf{0}$$

$$0 \qquad 0 \qquad \mathbf{w}_{1/|w_{1}|} \qquad 0 \qquad 0$$

$$0 \qquad 0 \qquad \ddots \qquad 0$$

$$0 \qquad 0 \qquad w_{n_{5}/|w_{n_{5}}|} \end{bmatrix} n_{s}$$

$$0 \qquad 0 \qquad 0 \qquad 0 \qquad 0$$

$$0 \qquad 0 \qquad 0$$

 $\Rightarrow$  The  $\mathcal{M}$  on  $\hat{A}$  produces  $A_f$  with diagonal entries  $\hat{a}_{ij}$  for  $(i,j) \in \mathcal{M}$  of maximal product.

We can formally define a prolongator.

$$P = \begin{bmatrix} \tilde{P} & O \\ O & W \end{bmatrix} = [\mathbf{p}_1, \dots, \mathbf{p}_J].$$

Then the preconditioner is the linear operator corresponding to the multiplicative composition of

$$I - B_{l}A_{l} = (I - (M_{l})^{-T}A_{l})(I - P_{l}B_{l+1}(P_{l})^{T}A_{l})(I - M_{l}^{-1}A_{l}) \quad \forall l < nl,$$

where 
$$A_{l+1} = (P_l)^T A_l P_l$$
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- To increase dimension reduction we can perform more than one sweep of matching per step,
- To increase regularity of  $P_l$  we can consider a smoothed prolongator by applying a Jacobi smoother,

$$P_I^s = (I - \omega D_I^{-1} A_I) P_I$$
, for  $D_I = \operatorname{diag}(A_I)$ .

We can formally define a prolongator.

$$P = \begin{bmatrix} \tilde{P} & O \\ O & W \end{bmatrix} = [\mathbf{p}_1, \dots, \mathbf{p}_J].$$

Then the preconditioner is the linear operator corresponding to the multiplicative composition of

$$I - B_{I}A_{I} = (I - (M_{I})^{-T}A_{I})(I - P_{I}B_{I+1}(P_{I})^{T}A_{I})(I - M_{I}^{-1}A_{I}) \quad \forall I < nI,$$

where  $A_{l+1} = (P_l)^T A_l P_l$  for l = 0, ..., nl - 1.

- To increase dimension reduction we can perform more than one sweep of matching per step,
- To increase regularity of  $P_l$  we can consider a smoothed prolongator by applying a Jacobi smoother,
- To increase the robustness we can use a non stationary solver as smoother.

Comparison with the preconditioners available in the Hypre, a state of the art preconditioning library from LLNL.

- ■ Run on the MareNostrum machine up to 8192 cores
- ★ Test: 3D Constant coefficient Poisson Problem with FCG
- **☞** DoF: 256k unknown × MPI core
- **T** Measures: Operator Complexity opc =  $\frac{\sum_{l=0}^{\mathsf{nl}-1} \mathrm{nnz}(A_l)}{\mathrm{nnz}(A_0)}$  and Solve Time (s).

Comparison with the preconditioners available in the Hypre, a state of the art preconditioning library from LLNL.

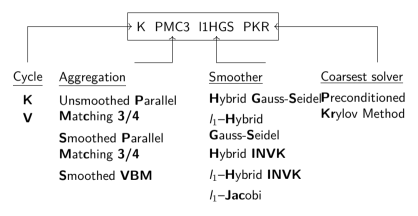
- ★ Test: 3D Constant coefficient Poisson Problem with FCG
- **T** Measures: Operator Complexity opc =  $\frac{\sum_{l=0}^{n-1} \text{nnz}(A_l)}{\text{nnz}(A_0)}$  and Solve Time (s).

#### Scaling

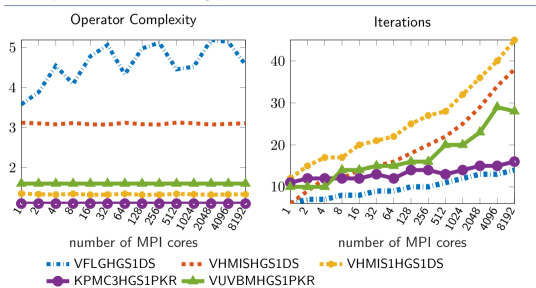
There are two common notions of scalability:

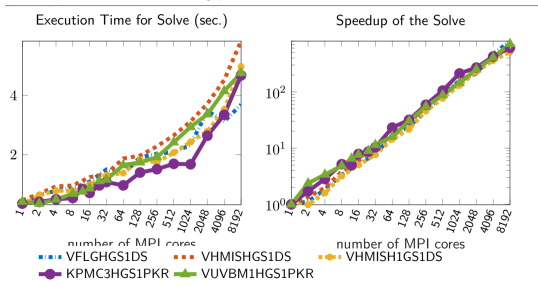
- Strong scaling is defined as how the solution time varies with the number of processors for a fixed total problem size.
- Weak scaling is defined as how the solution time varies with the number of processors for a fixed problem size per processor.

Giving a name to preconditioners with many parameters:



For Hypre we test HMIS and Falgout coarsening schemes.





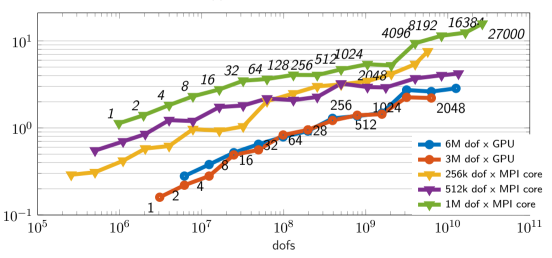
# Weak Scalability - CPU/GPU Runs - Piz Daint

The resulting performance of the multigrid preconditioner in term of implementation scalability depends also on how effective the coarsening procedure is, and on how well balanced is the distribution of the coarsest matrix.

- Run on the Piz Daint machine up to 28800 cores and 2048 GPUs
- **■** DoF: 256k/512k/1M unknown × MPI core and 3M/6M per GPUs
- Measures: execution time for solve

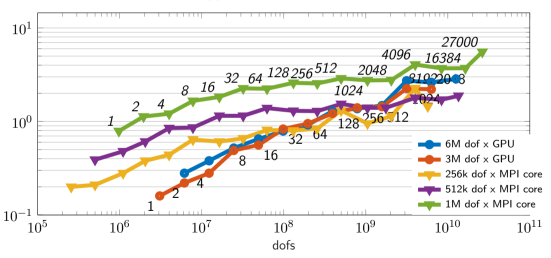
# Weak Scalability - CPU/GPU Runs - Piz Daint

Execution Time for Solve (s) - K-PMC3-HGS1-PKR vs VS-PMC3-L1JAC-PKR

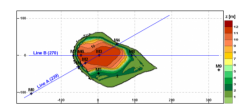


# Weak Scalability - CPU/GPU Runs - Piz Daint





#### A Large Eddy Scale simulation inside Alya



Bolund is an isolated hill situated in Roskilde Fjord, Denmark. An almost vertical escarpment in the prevailing W-SW sector ensures flow separation in the windward edge resulting in a complex flow field.

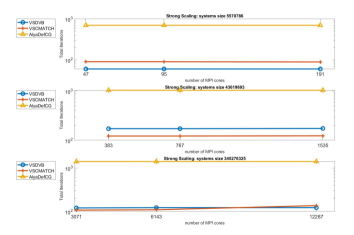
- Model: 3D incompressible unsteady Navier-Stokes equations for the Large Eddy Simulations of turbulent flows.
- **Discretization**: low-dissipation mixed FEM (linear FEM both for velocity and pressure),
- Time-Stepping: non-incremental fractional-step for pressure, explicit fourth order Runge-Kutta method for velocity.

#### Alya

Alya is a simulation code for high performance computational mechanics. It solves coupled multiphysics problems using high performance computing techniques for distributed and shared memory supercomputers, together with vectorization and optimization at the node level.

#### **Bolund Test Case - Strong Scaling - Pressure Eq.**

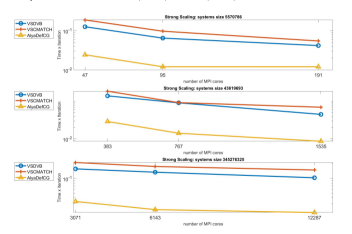
Fixed size problem with n = 5,570,786;43,619,693;345.276.325 dofs, 100 time steps



Total number of linear iterations is smaller and stable for increasing number of cores.

#### **Bolund Test Case - Strong Scaling - Pressure Eq.**

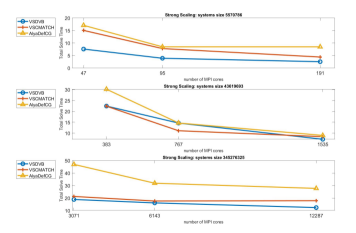
Fixed size problem with n = 5,570,786;43,619,693;345.276.325 dofs, 100 time steps



Total number of linear iterations is smaller and stable for increasing number of cores, the time needed per each iteration decreases for increasing number of cores.

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