

Approximation of the Inverse Hessian in Large-Scale Variational Data Assimilation Problems

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Four-dimensional Variational Assimilation (4D-Var)

4D-Var aims to find the solution of a numerical forecast model that best fits sequences of observations distributed in space over a finite time interval.

Minimise cost function

$$J(\mathbf{v}_0) = (\mathbf{v}_0 - \mathbf{v}_0^B)^T \mathcal{B}^{-1} (\mathbf{v}_0 - \mathbf{v}_0^B) + \sum_{i=0}^n (\mathcal{H}(\mathbf{v}_i) - \mathbf{y}_i)^T \mathcal{R}^{-1} (\mathcal{H}(\mathbf{v}_i) - \mathbf{y}_i)$$

with **constraint** $\mathbf{v}_i = \mathcal{M}^{i,0}(\mathbf{v}_0)$.

analysis	\mathbf{v}_0
background (short-term forecast)	\mathbf{v}_0^B
observations	\mathbf{y}
observation operator	\mathcal{H}
model dynamics	$\mathbf{v}_{i+1} = \mathcal{M}(\mathbf{v}_i)$
background error covariance matrix	\mathcal{B}
observation error covariance matrix	\mathcal{R}

- Linearise \mathcal{H} , \mathcal{M} and solve resulting **unconstrained** optimisation problem iteratively:

$$\bar{H}_{k-1}^i \equiv \left. \frac{\partial \mathcal{H}^i}{\partial \mathbf{v}} \right|_{\mathbf{v}=\mathbf{v}_{k-1}}, \quad \bar{M}_{k-1}^{i,0} \equiv \left. \frac{\partial \mathcal{M}^{i,0}}{\partial \mathbf{v}} \right|_{\mathbf{v}=\mathbf{v}_{k-1}}$$

- Hessian** of the cost function is

$$\mathbb{H} = \mathcal{B}^{-1} + \hat{H}^T \hat{\mathcal{R}}^{-1} \hat{H}$$

where

$$\begin{aligned} \hat{H} &= [(\bar{H}^0)^T, (\bar{H}^1 \bar{M}^{1,0})^T, \dots, (\bar{H}^N \bar{M}^{N,0})^T]^T \\ \hat{\mathcal{R}} &= \text{bldiag}(\mathcal{R}_i), \quad i = 1, \dots, N. \end{aligned}$$

- Cannot store \mathbb{H} as a matrix: only action of **applying \mathbb{H} to a vector** is available.

- Why approximate \mathbb{H}^{-1} ?
 - \mathbb{H}^{-1} represents an approximation of the **Posterior Covariance Matrix** (PCM).
 - The PCM can be used to find **confidence intervals** and carry out *a posteriori* error analysis.
 - $\mathbb{H}^{-1/2}$ can be used in **ensemble forecasting**.
 - \mathbb{H}^{-1} , $\mathbb{H}^{-1/2}$ can be used for **preconditioning** in a Gauss-Newton method.
- What are the main challenges?
 - Evaluating $\mathbb{H}\mathbf{v}$ is expensive in terms of computing time and memory (involves both **forward** and **backward** model solves with a sequence of tangent linear and adjoint problems).
 - No obvious equivalent option exists for evaluating $\mathbb{H}^{-1}\mathbf{v}$.

First level preconditioning

$$\mathbb{H} = \mathcal{B}^{-1} + \hat{H}^T \hat{\mathcal{R}}^{-1} \hat{H}$$

- Precondition \mathbb{H} based on the background covariance matrix (control variable transform):

$$H = (\mathcal{B}^{1/2})^T \mathbb{H} \mathcal{B}^{1/2} = I + (\mathcal{B}^{1/2})^T \hat{H}^T \hat{\mathcal{R}}^{-1} \hat{H} \mathcal{B}^{1/2}$$

- Eigenvalues of H are bounded below by one: more details on the full **eigenspectrum** can be found in HABEN ET AL. (2011), TABEART ET AL. (2018).
- Aim here is to construct a **limited-memory approximation** to the action of H^{-1} using only matrix-vector multiplication.

Limited-memory approximation for H^{-1}

- H amenable to **limited-memory approximation**.
- Find n_e leading eigenvalues and orthonormal eigenvectors using the **Lanczos** method (needs only $H\mathbf{v}$).
- Construct approximation

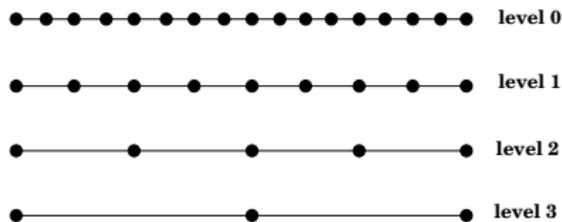
$$H \approx I + \sum_{i=1}^{n_e} (\lambda_i - 1) \mathbf{u}_i \mathbf{u}_i^T$$

- Can also use this to easily approximate matrix powers (including H^{-1} and $H^{-1/2}$):

$$H^p \approx I + \sum_{i=1}^{n_e} (\lambda_i^p - 1) \mathbf{u}_i \mathbf{u}_i^T$$

Multilevel limited-memory approximation

- Sequence of grid levels $k = 0, 1, 2, \dots$



- Matrix H_0 is available on finest grid $k = 0$.
- Construct a **multilevel** approximation to H_0^{-1} based on limited-memory approximations on a sequence of nested grids.
- Need **grid transfer operators** (more shortly).
- Identity matrix I_k on grid level k .
- $[H]_{\rightarrow k}$ means “matrix H transferred to grid level k ”.

Grid transfers for vectors

- Coarse grid level $k = c$; fine grid level $k = f$.
- Restriction matrix R ; prolongation matrix P : assume “perfect interpolation”, i.e., $RP = I_c$.
- Split fine grid vector into two parts:

$$\mathbf{v}_f = \mathbf{v}_f^{(1)} + \mathbf{v}_f^{(2)} = (I_f - PR)\mathbf{v}_f + PR\mathbf{v}_f.$$

- Restrict \mathbf{v}_f to coarse grid:

$$\mathbf{v}_c^{(1)} = R\mathbf{v}_f^{(1)} = R(I_f - PR)\mathbf{v}_f = (R - (RP)R)\mathbf{v}_f = \mathbf{0}$$

$$\mathbf{v}_c^{(2)} = R\mathbf{v}_f^{(2)} = (RP)R\mathbf{v}_f = R\mathbf{v}_f.$$

- Modes in $\mathbf{v}_f^{(1)}$ are not supported on coarse grid.

Grid transfers for matrices

- Consider action of coarse grid matrix H_c on a fine grid vector:

$$\begin{aligned}[H_c]_{\rightarrow f} \mathbf{v}_f &= \mathbf{v}_f^{(1)} + PH_c R \mathbf{v}_f^{(2)} \\ &= (I_f - PR) \mathbf{v}_f + PH_c (RP) R \mathbf{v}_f \\ &= (P(H_c - I_c)R + I_f) \mathbf{v}_f\end{aligned}$$

- This motivates matrix transfer operators
 - From coarse grid to fine grid

$$[H_c]_{\rightarrow f} = P(H_c - I_c)R + I_f$$

- From fine grid to coarse grid

$$[H_f]_{\rightarrow c} = R(H_f - I_f)P + I_c$$

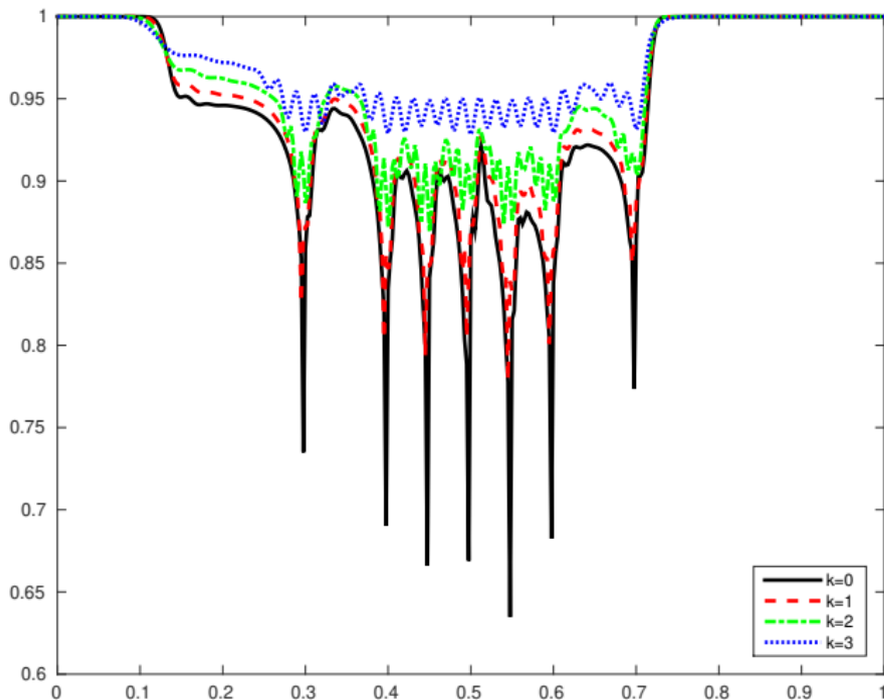
Test problem 1

- Model is 1D **Burgers' equation**.
- Discretise evolution equation on a grid with $m + 1$ nodes (level 0) to represent full Hessian H_0 .
- Grid level k contains $m_k = m/2^k + 1$ nodes.
- 1D uniform grid with 7 sensors located at 0.3, 0.4, 0.45, 0.5, 0.55, 0.6, and 0.7 in $[0, 1]$.
- Construct a multilevel approximation to H^{-1} with **four** grid levels:

k	0	1	2	3
grid points	401	201	101	51

Hessian in a multilevel framework

- Diagonal of H^{-1} on various grid levels:



Motivating idea

- Eigenvalues of $[H_c^{-1/2}]_{\rightarrow f} H_f [H_c^{-1/2}]_{\rightarrow f}$ should be clustered around 1.
- Construct an approximation to $H_c^{-1/2}$:

- Precondition H_c to obtain $\tilde{H}_c = M^T H_c M$ with eigenvalues closer to 1.
- Build \hat{H}_c , a **limited memory approximation** for \tilde{H}_c using n_c eigenvalues with the **Lanczos** method.
- Note that

$$H_c^{-1} = M\tilde{H}_c^{-1}M^T \simeq M\hat{H}_c^{-1}M^T$$

so

$$H_c^{-1/2} = M\tilde{H}_c^{-1/2} \simeq M\hat{H}_c^{-1/2}.$$

- Use $\hat{M} = [M\hat{H}_c^{-1/2}]_{\rightarrow f}$ as a preconditioner on the level above.

Outline of multilevel concept

Step 1. Start on coarsest grid level.

Step 2. Represent H_0 on grid level k as $H_k = [H_0]_{\rightarrow k}$.

Step 3. Precondition this to obtain $\tilde{H}_k = M_k^T H_k M_k$.

Step 4. Build limited memory approximation $\hat{H}_k^{-1/2}$.

Step 5. Project $\hat{M}_k = M_k \hat{H}_k^{-1/2}$ to the level above to be used as preconditioner at the next coarsest level.

Step 6. Move up one grid level and repeat from step 2.

- On coarsest grid, level $k + 1$ does not exist so set $M_k = I_k$.
- For other levels, M_k is constructed on level $k + 1$ and applied on level k .
- Preconditioners are constructed **recursively**:

$$M_k = [\hat{M}_{k+1}]_{\rightarrow k} = \left[M_{k+1} \hat{H}_{k+1}^{-1/2} \right]_{\rightarrow k}.$$

- At level 0, final inverse Hessian approximation H_{approx}^{-1} will contain eigenvalue information from **all levels**.

Algorithm in practice

- use $N_e = (n_0, n_1, \dots, n_{k_c})$ eigenvalues at each level

```
[ $\Lambda, \mathcal{U}$ ] =  $MLalg(H_0, N_e)$ 
for  $k = k_c, k_c - 1, \dots, 0$ 
  compute by the Lanczos method
     $\{\lambda_k^i, U_k^i\}, i = 1, \dots, n_k$  of  $\tilde{H}_{0 \rightarrow k}$ 
  using preconditioner  $M_k$ 
end
```

- storage:

$$\Lambda = [\lambda_0^1, \dots, \lambda_0^{n_0}, \lambda_1^1, \dots, \lambda_1^{n_1}, \dots, \lambda_{k_c}^1, \dots, \lambda_{k_c}^{n_{k_c}}],$$
$$\mathcal{U} = [U_0^1, \dots, U_0^{n_0}, U_1^1, \dots, U_1^{n_1}, \dots, U_{k_c}^1, \dots, U_{k_c}^{n_{k_c}}].$$

Assessing approximation accuracy

- **Riemannian** distance:

$$\delta(A, B) = \|\ln(B^{-1}A)\|_F = \left(\sum_{i=1}^n \ln^2 \lambda_i \right)^{1/2}$$

- Compare eigenvalues of H^{-1} and H_{approx}^{-1} on the finest grid level $k = 0$ using distance function

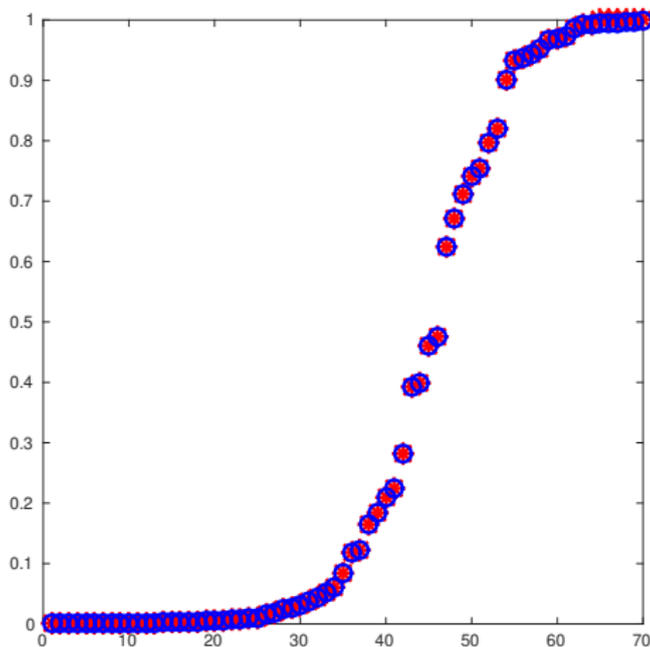
$$D = \frac{\delta(H^{-1}, H_{approx}^{-1})}{\delta(H^{-1}, I)}$$

- Vary number of eigenvalues chosen on each grid level

$$N_e = (n_0, n_1, \dots, n_{k_c})$$

Eigenvalues of the inverse Hessian

- Exact (blue circles), approximated (red stars)

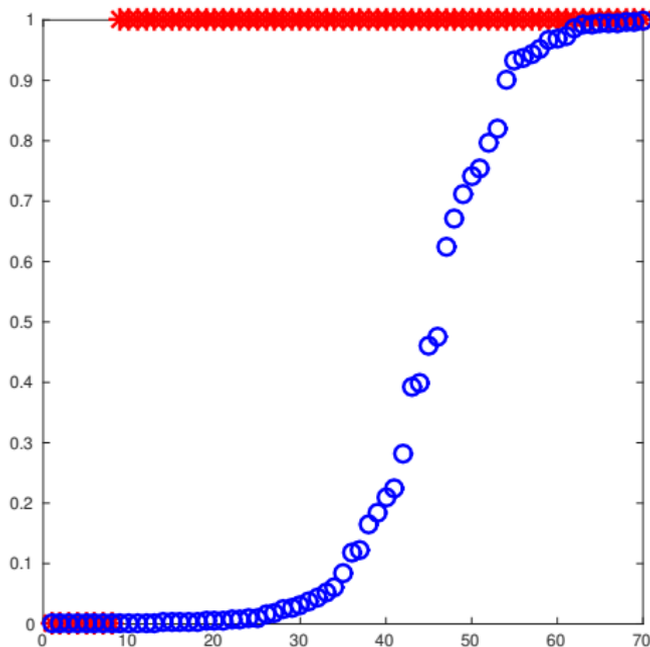


$$N_e = (64, 0, 0, 0)$$

$$D = 2.98e - 4$$

Eigenvalues of the inverse Hessian

- Exact (blue circles), approximated (red stars)

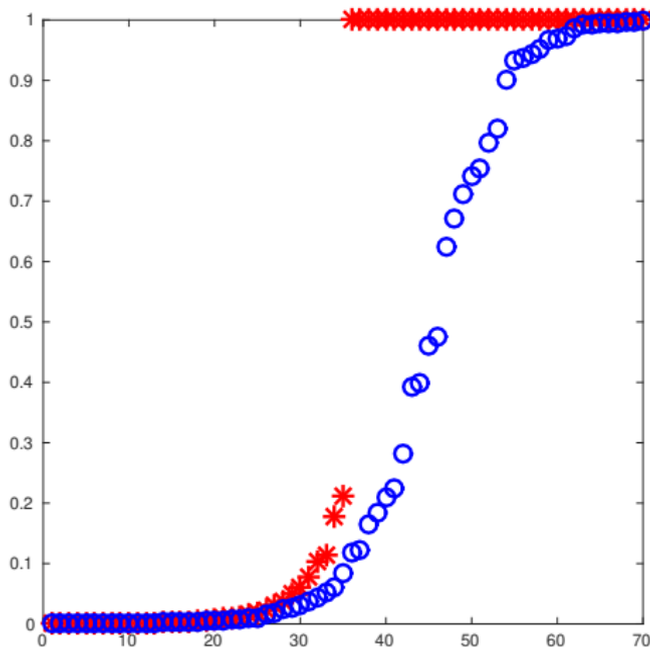


$$N_e = (8, 0, 0, 0)$$

$$D = 7.71e - 1$$

Eigenvalues of the inverse Hessian

- Exact (blue circles), approximated (red stars)

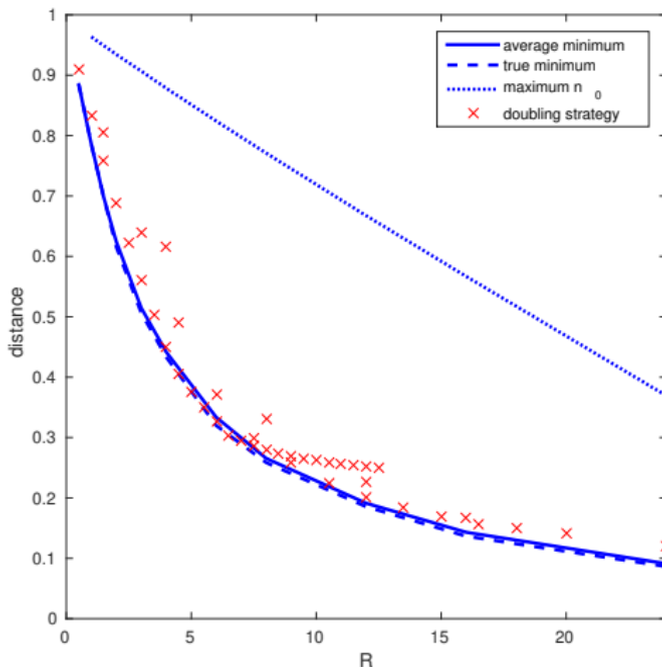


$$N_e = (0, 0, 29, 6)$$

$$D = 3.82e - 1$$

Fixed memory ratio

- Fixed memory ratio $R = \sum_{k=0}^{k_c} \frac{n_k}{2^k}$



Example: PCG iteration for one Newton step

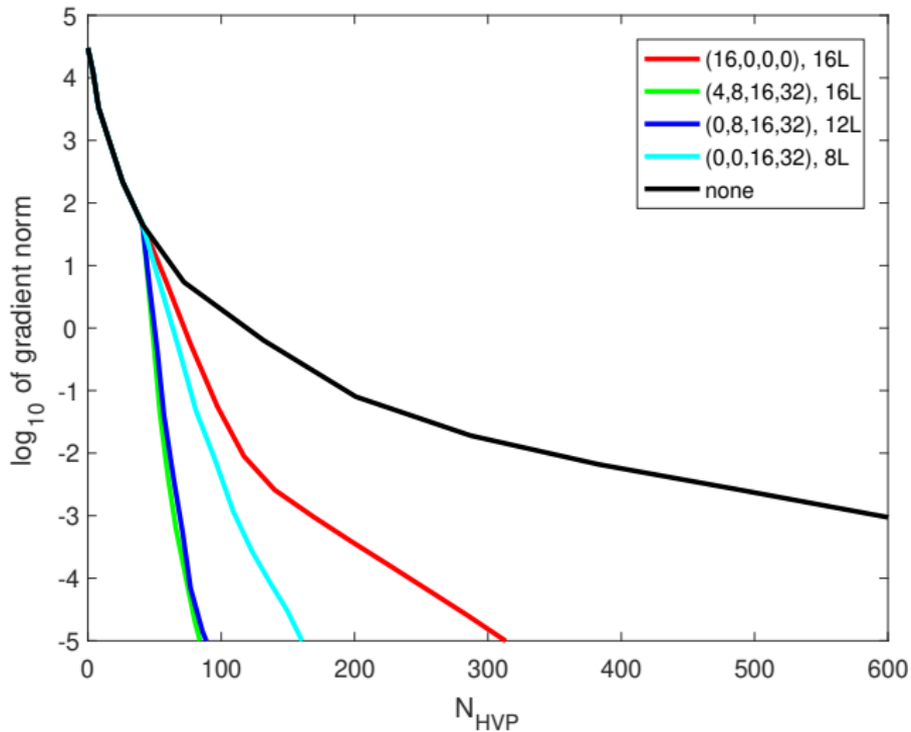
- Hessian linear system (within a Gauss-Newton method):

$$H(\mathbf{u}_k)\delta\mathbf{u}_k = G(\mathbf{u}_k)$$

- Solve using **P**reconditioned **C**onjugate **G**radient iteration (needs only $H\mathbf{v}$).
- measurement units
 - storage: length of vector on finest grid **L**
 - solve cost: cost of HVP on finest grid **HVP**

Preconditioner	# CG iterations	storage	solve cost
none	57	0 L	57 HVP
ML(400,0,0,0)	1	400 L	402 HVP
ML(4,8,16,32)	4	16 L	34 HVP
ML(0,8,16,32)	5	12 L	14 HVP
ML(0,0,16,32)	8	8 L	10 HVP

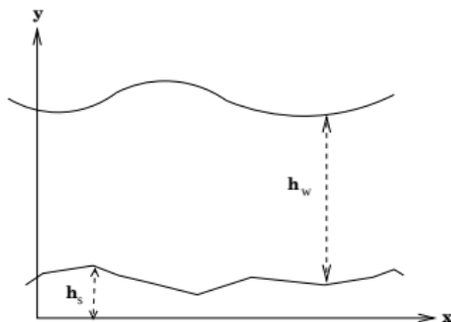
Solve cost measured in number of HVPs



Test problem 2

- Model is 1D **shallow water equations** for velocity u and geopotential $\phi = gh_w$.

$$\frac{Du}{Dt} + \frac{\partial \phi}{\partial x} = -g \frac{\partial h_s}{\partial x}$$
$$\frac{D(\ln \phi)}{Dt} + \frac{\partial u}{\partial x} = 0$$



- Uniformly spaced** sensors.
- Four grid** multilevel structure as before.

PCG iteration for one Newton step

- Background covariance matrix B constructed using a Laplacian correlation function.

	# PCG iterations			
Preconditioner	$n = 400$	$n = 800$	$n = 1600$	$n = 3200$
none	308	1302	5,879	25,085
ML(4,0,0,0)	38	34	34	47
ML(1,2,4,8)	31	29	28	37
ML(0,2,4,16)	27	26	24	32
ML(0,0,8,16)	26	25	24	30
ML(0,0,0,32)	23	19	19	24

PCG iteration for one Newton step

- Background covariance matrix \mathcal{B} constructed using a **Second-Order Auto-Regressive (SOAR)** correlation function.

	# PCG iterations			
Preconditioner	$n = 400$	$n = 800$	$n = 1600$	$n = 3200$
none	509	2,277	10,453	43,915
ML(4,0,0,0)	39	35	35	44
ML(1,2,4,8)	28	26	26	34
ML(0,2,4,16)	23	22	21	27
ML(0,0,8,16)	22	21	20	26
ML(0,0,0,32)	19	16	15	20

Practical implementation

- Algorithm based solely on repeated use of the **Lanczos** method at each level (for building limited-memory approximations).
- This calculation of largest eigenvalues and associated eigenvectors is **expensive**, and needs to be done with reasonable accuracy.
- Use of a randomised **Nyström** method looks promising.
- Major reduction in computation times comes from using a **Hessian decomposition** technique.

Hessian decomposition

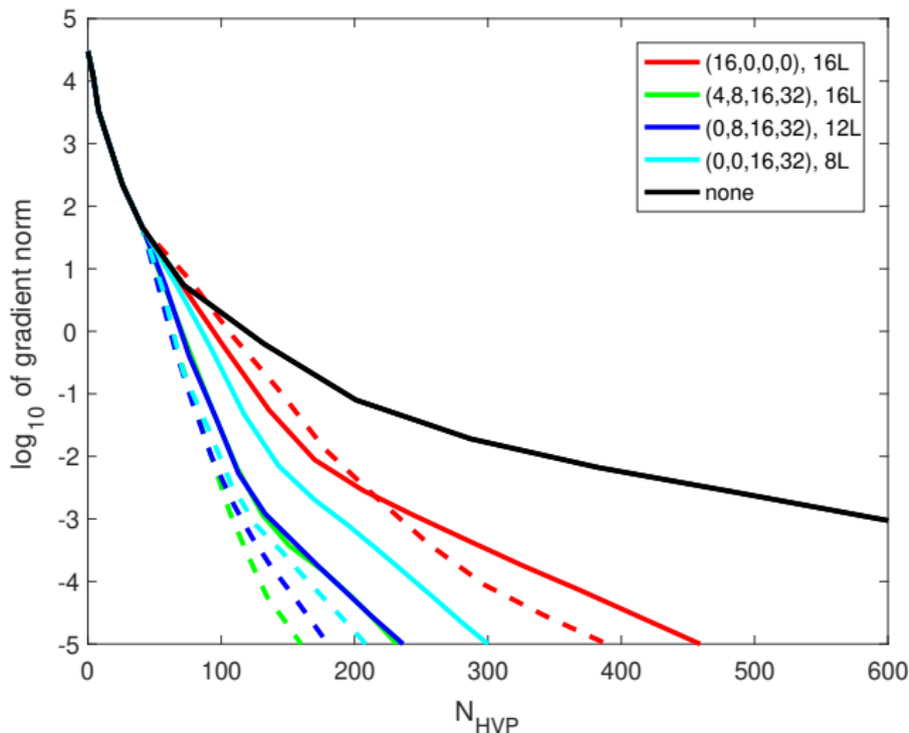
- partition domain into S subregions and compute **local Hessians** H^s such that

$$H(\mathbf{v}) = I + \sum_{s=1}^S (H^s(\mathbf{v}) - I)$$

- computational advantages of local Hessians:
 - fewer eigenvalues** required for limited-memory approximation;
 - can be calculated at a **coarser grid** level;
 - can use **local** rather than global models;
 - can be computed in **parallel**.

Sample costs including building preconditioner

- Local Hessians with 8 eigenvalues at level 0 (solid lines) or level 1 (dashed lines).



Concluding remarks

- Algorithm for building limited-memory approximations to the inverse Hessian seems promising.
- Difficult to identify the **correct number of eigenvalues** to use at each level: good rule of thumb available but analysis would be better!
- Full algorithm may not always be practical, but we have developed practical implementations based on **Hessian decompositions**.
- Currently investigating the use of **faster eigenvalue** techniques.
- Also works well for other configurations (e.g. moving sensors, different initial conditions).
- Potential for extension to higher dimensions and other applications.

Thank you!