

# Efficient computation of sparse matrix functions for large scale electronic structure calculations: The CheSS library

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ELSI Connector Meeting 2017

# Outline

- History and motivation for CheSS
- Applicability of CheSS
- Short overview of the theory behind CheSS
- Various performance data

# Motivation for CheSS

CheSS is a “spin-off” of the linear scaling version of BigDFT.

- localized basis set leads to sparse matrices
- we have to exploit this sparsity to reach linear scaling
- we did not find a package that fits our need

⇒ we created our own sparse matrix routines within BigDFT

We have the same situation in all DFT codes with a localized basis set

⇒ we created a standalone library: CheSS

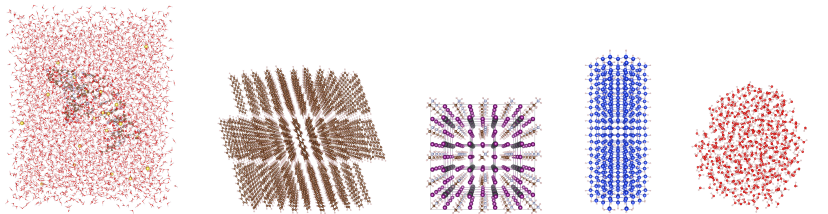
CheSS can be obtained for free from <https://launchpad.net/chess>

A paper is in review: <https://arxiv.org/abs/1704.00512>

# Applicability of CheSS

CheSS performs best for matrices exhibiting a small spectral width.

Can this be obtained in practice?



system	#atoms	S					H				
		sparsity	$\epsilon_{min}$	$\epsilon_{max}$	$\kappa$		sparsity	$\epsilon_{min}$	$\epsilon_{max}$	$\lambda$	$\Delta_{HL}$
DNA	15613	99.57%	0.72	1.65	2.29		98.46%	-29.58	19.67	49.25	2.76
bulk pentacene	6876	98.96%	0.78	1.77	2.26		97.11%	-21.83	20.47	42.30	1.03
perovskite	768	90.34%	0.70	1.50	2.15		76.47%	-20.41	26.85	47.25	2.19
Si nanowire	706	93.24%	0.72	1.54	2.16		81.61%	-16.03	25.50	41.54	2.29
water	1800	96.71%	0.83	1.30	1.57		90.06%	-26.55	11.71	38.26	9.95



# Basic idea

In CheSS we approximate matrix functions by Chebyshev polynomials:

$$p(\mathbf{M}) = \frac{c_0}{2} \mathbf{I} + \sum_{i=1}^{n_{pl}} c_i \mathbf{T}^i(\tilde{\mathbf{M}}),$$

with

$$\tilde{\mathbf{M}} = \sigma(\mathbf{M} - \tau \mathbf{I}) \quad ; \quad \sigma = \frac{2}{\epsilon_{max} - \epsilon_{min}} \quad ; \quad \tau = \frac{\epsilon_{min} + \epsilon_{max}}{2}$$

and

$$c_j = \frac{2}{n_{pl}} \sum_{k=0}^{n_{pl}-1} f \left[ \frac{1}{\sigma} \cos \left( \frac{\pi(k + \frac{1}{2})}{n_{pl}} \right) + \tau \right] \cos \left( \frac{\pi j(k + \frac{1}{2})}{n_{pl}} \right).$$

Recursion relation for the Chebyshev polynomials:

$$\mathbf{T}^0(\tilde{\mathbf{M}}) = \mathbf{I},$$

$$\mathbf{T}^1(\tilde{\mathbf{M}}) = \tilde{\mathbf{M}},$$

$$\mathbf{T}^{j+1}(\tilde{\mathbf{M}}) = 2\tilde{\mathbf{M}}\mathbf{T}^j(\tilde{\mathbf{M}}) - \mathbf{T}^{j-1}(\tilde{\mathbf{M}}).$$

Each column independent  
 $\implies$  easily parallelizable.

Strict sparsity pattern  
 $\implies$  linear scaling

# Available functions

CheSS can calculate those matrix functions needed for DFT:

- density matrix:  $f(x) = \frac{1}{1+e^{\beta(x-\mu)}}$   
(or  $f(x) = \frac{1}{2} [1 - \operatorname{erf}(\beta(\epsilon - x))]$ )
- energy density matrix:  $f(x) = \frac{x}{1+e^{\beta(x-\mu)}}$   
(or  $f(x) = \frac{x}{2} [1 - \operatorname{erf}(\beta(\epsilon - x))]$ )
- matrix powers:  $f(x) = x^a$  ( $a$  can be non-integer!)

We can calculate arbitrary functions by changing only the coefficients  $c_j$ !

Only requirement:

function  $f$  must be well representable by Chebyshev polynomials over the entire eigenvalue spectrum.

# Sparsity and truncation

CheSS works with predefined sparsity patterns.

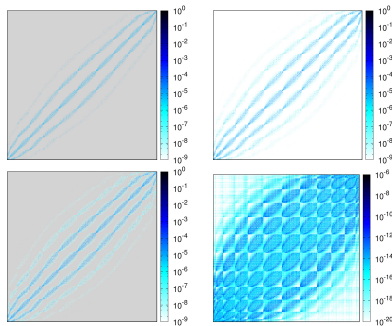
In general there are three:

- pattern for the original matrix  $\mathbf{M}$
- pattern for the matrix function  $f(\mathbf{M})$
- auxiliary pattern to perform the matrix-vector multiplications

At the moment all of them must be defined by the user.

Typically: distances atoms / basis functions

- 1 original matrix  $\mathbf{M}$
- 2 exact calculation of  $\mathbf{M}^{-1}$  without sparsity constraints
- 3 sparse calculation of  $\mathbf{M}^{-1}$  using CheSS within the sparsity pattern
- 4 difference between Fig. 2 and Fig. 3



# Accuracy – error definition

There are two possible factors affecting the accuracy of CheSS:

- error introduced due to the enforced sparsity (truncating the matrix-vector multiplications)
- error introduced by the Chebyshev fit

This also affects the definition of the “exact solution”. Two possibilities:

- 1 calculate the solution exactly and without sparsity constraints and then crop to the sparsity pattern. Shortcoming: violates in general the identity  $f^{-1}(f(\mathbf{M})) = \mathbf{M}$
- 2 calculate the solution within the sparsity pattern, and define as exact one that which fulfills  $\hat{f}^{-1}(\hat{f}(\mathbf{M})) = \mathbf{M}$

According error definitions:

$$1 \quad w_{\hat{f}_{\text{sparse}}} = \frac{1}{|\hat{f}(\mathbf{M})|} \sqrt{\sum_{(\alpha\beta) \in \hat{f}(\mathbf{M})} \left( \hat{f}(\mathbf{M})_{\alpha\beta} - f(\mathbf{M})_{\alpha\beta} \right)^2}$$

$$2 \quad w_{\hat{f}^{-1}(\hat{f})} = \frac{1}{|\hat{f}(\mathbf{M})|} \sqrt{\sum_{(\alpha\beta) \in \hat{f}(\mathbf{M})} \left( \hat{f}^{-1}(\hat{f}(\mathbf{M}))_{\alpha\beta} - M_{\alpha\beta} \right)^2}$$

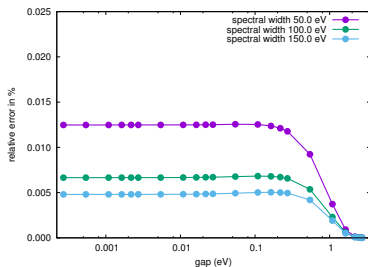
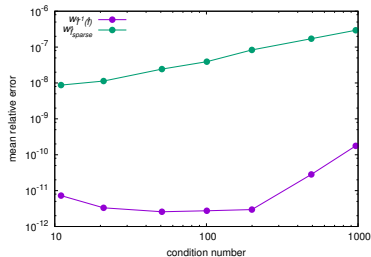
# Accuracy

Inverse:

- $w_{\hat{f}-1}(\hat{f})$ : error due to Chebyshev fit basically zero
- $w_{\hat{f}_{\text{sparse}}}$ : error due to sparsity pattern very small

Density matrix:

- energy (i.e.  $\text{Tr}(\mathbf{KH})$ ): relative error of only 0.01%
- slightly larger error for small spectral width: eigenvalues are denser, finite temperature smearing affects more

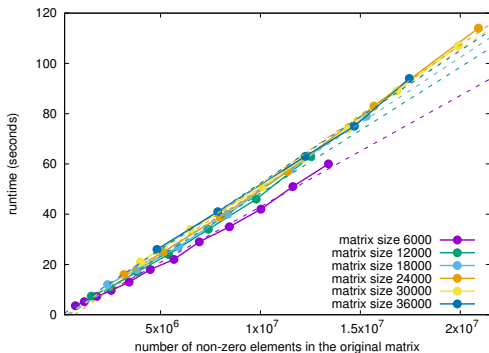


# Scaling with matrix size and sparsity

Series of matrices with the same “degree of sparsity”  
(DFT calculations of water droplet of various size).

Example: calculation of the inverse

- Runtime only depends on the number of non-zero elements of  $\mathbf{M}$
- no dependence on the total matrix size

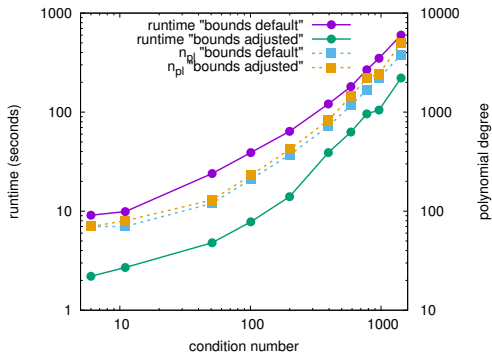


# Scaling with spectral properties

CheSS is extremely sensitive to the eigenvalue spectrum:

- required polynomial degree strongly increases with the spectral width
- as a consequence the runtime strongly increases as well
- a good input guess for the eigenvalue bounds helps a lot

Example: calculation of the inverse



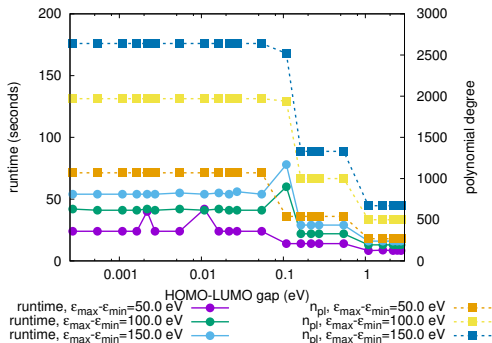
# Scaling with spectral properties

For the density matrix the performance depends on two parameters:

- spectral width (the smaller the better)
- HOMO-LUMO gap (the larger the better)

In both cases the polynomial degree can increase considerably

⇒ CheSS less efficient.



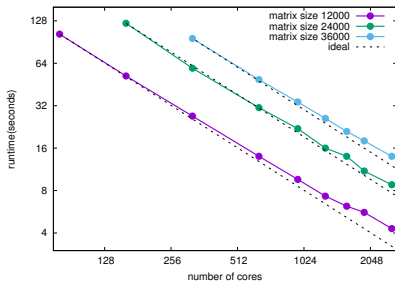
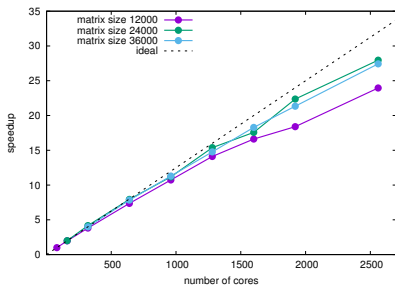


# Parallel scaling

The most compute intensive part of CheSS is the matrix-vector multiplications.

- Easily parallelizable
- identical for all operations

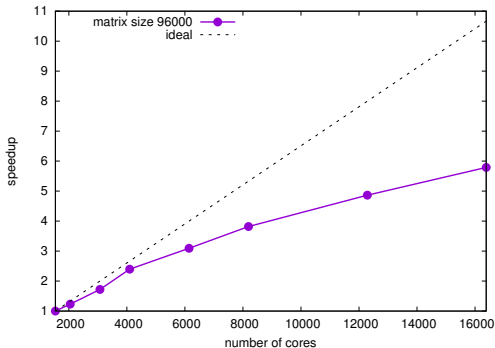
Example: Calculation of  $\mathbf{M}^{-1}$  (runs performed with 16 OpenMP threads)



# Extreme scaling

We have also performed extreme-scaling tests from 1536 to 16384 cores.

Example: Calculation of  $\mathbf{M}^{-1}$  (runs performed with 8 OpenMP threads)



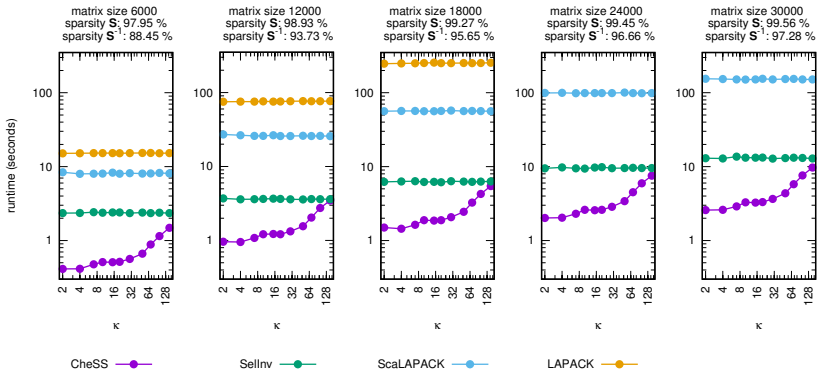
Given the small matrix (96,000) the obtained scaling is acceptable.

We will try to further improve the scaling.

# Comparison with other methods: Inverse

Comparison of the matrix inversion between:

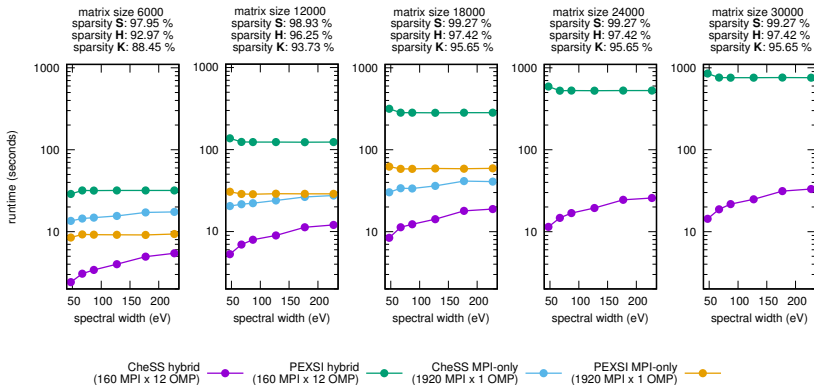
- CheSS
- SellInv
- ScaLAPACK
- LAPACK



# Comparison with other methods: Density matrix

Comparison of the density matrix calculation between:

- CheSS hybrid MPI/OpenMP
- PEXSI hybrid MPI/OpenMP
- CheSS MPI-only
- PEXSI MPI-only




# Conclusions

- CheSS is a flexible tool to calculate matrix functions for DFT
- can easily be extended to further functions
- exploits sparsity of the matrices, linear scaling possible
- works best for small spectral widths of the matrices
- very good parallel scaling (both MPI and OpenMP)

Most important:

CheSS is about to be interfaced by ELSI!



**Thank you for your attention!**