

### Current state of the art in TDDFT code

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#### Current state of the art in TDDFT code

Peter Koval, Olivier Coulaud, Dietrich Foerster













Biarritz 07/10/2009

### Spectroscopy versus linear response theory

- Spectroscopic properties of molecules can be explained with linear response theory
- Linear response theory: all observables can be computed with ground state wave function
- Key quantity is linear density response:

$$\delta n(\mathbf{r},t) = \int \chi(\mathbf{r},\mathbf{r}',t'-t)\delta V(\mathbf{r}',t') d^3r'dt',$$

In dipole approximation, the polarizability reads

$$P_{ik}(\omega) = \int \mathbf{r}_i \chi(\mathbf{r}, \mathbf{r}', \omega) \mathbf{r}'_k \, \mathrm{d}^3 r \mathrm{d}^3 r'.$$

### TDDFT linear response theory

If 
$$(T+V)\psi_E = E\psi_E$$

then  $\chi = \frac{\delta \mathbf{n}}{\delta V} = \sum_{\mathbf{F} \cdot \mathbf{F} < 0} (f_{\mathbf{E}} - f_{\mathbf{F}}) \frac{\psi_{\mathbf{E}}(\mathbf{r}) \psi_{\mathbf{F}}(\mathbf{r}) \psi_{\mathbf{F}}(\mathbf{r}') \psi_{\mathbf{E}}(\mathbf{r}')}{\omega - (\mathbf{E} - \mathbf{F}) + \mathrm{i}\varepsilon}$ 



Erwin Schrödinger

$$V = V_{\text{ext}}$$



Walter Kohn

$$V = V_{\text{eff}} = V_{\text{ext}} + V_{\text{Hxc}}$$

### TDDFT linear response theory

Kohn-Sham response  $\chi_0$  and interacting response  $\chi$  are connected<sup>1</sup>

$$\chi_0 = \frac{\delta n}{\delta V_{\text{eff}}}$$

$$\chi = \frac{\delta n}{\delta V_{\text{ext}}}$$

$$V_{
m eff} = V_{
m ext} + V_{
m Hxc} \ \Rightarrow \ rac{\delta V_{
m eff}}{\delta n} = rac{\delta V_{
m ext}}{\delta n} + rac{\delta V_{
m Hxc}}{\delta n} \Rightarrow$$

$$\chi_0^{-1} = \chi^{-1} + \Sigma \Rightarrow$$
$$\chi = \frac{1}{1 - \chi_0 \Sigma} \chi_0.$$

The problem is to effectively compute the interacting polarizability

$$P_{ik} = \langle \mathbf{r}_i | \chi | \mathbf{r}_k \rangle.$$

<sup>&</sup>lt;sup>1</sup>Petersilka, Gossmann and Gross, Phys. Rev. Lett., **76** (1996) 1212

# Suitable basis for non interacting response

Kohn-Sham density response reads

$$\chi = \frac{\delta n}{\delta V} = \sum_{E : E < 0} (f_E - f_E) \frac{\psi_E(\mathbf{r}) \psi_F(\mathbf{r}) \psi_F(\mathbf{r}') \psi_E(\mathbf{r}')}{\omega - (E - F) + i\varepsilon}$$

Note: formula contains products of eigenstates  $\psi_E(r)\psi_F(r)$ 

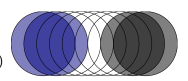
Using method of LCAO (linear combination of atomic orbitals), we get

$$\psi_{E}(\mathbf{r}) = \sum_{a} X_{a}^{E} f^{a}(\mathbf{r}) \quad \Rightarrow \quad \psi_{E}(\mathbf{r}) \psi_{F}(\mathbf{r}) = \sum_{ab} X_{a}^{E} X_{b}^{F} f^{a}(\mathbf{r}) f^{b}(\mathbf{r})$$

There are

O(N) localized products  $f^a(\mathbf{r})f^b(\mathbf{r})$ 

 $O(N^2)$  products of eigenstates  $\psi_E({m r})\psi_F({m r})$ 



# Dominant products<sup>2</sup>

Set of localized products  $f^a(\mathbf{r})f^b(\mathbf{r})$  linearly dependent  $\Rightarrow$  redundant

- ▶ find most important linear combinations of original products
- use these linear combinations as a basis

Compute a Gram matrix or metric 
$$g^{ab,cd}=\int f^a({m r})f^b({m r})f^c({m r})f^d({m r})\,d^3r$$
 Diagonalize metric  $gX^\lambda=\lambda X^\lambda$  Form orthogonal products  $F^\lambda({m r})=\sum_{ab}X^\lambda_{ab}f^a({m r})f^b({m r})$  Find  ${m vertex}\ V=X^{-1}=X^T$   $f^a({m r})f^b({m r})=\sum_{ab}V^{ab}_\lambda F^\lambda({m r})$ 

Use eigenvalue  $\lambda$  to define important **dominant products** 

<sup>&</sup>lt;sup>2</sup>D. Foerster, J. Chem. Phys. **128**, 034108 (2008)

#### Dominant products: constrains

Dominant products must preserve a simple form:

$$F^{\lambda}(\mathbf{r}) = \sum_{j} F_{j}^{\lambda}(\mathbf{r}) Y_{jm}(\mathbf{Rr})$$



- ⇒ expand the dominant products about a midpoint
- $\Rightarrow$  define the dominant products in a rotated frame R
- Metric g must be easily diagonalizable (small):
  - $\Rightarrow$  define metric g within a given atomic pair

#### Implication: vertex becomes a sparse object

$$f^a(\mathbf{r})f^b(\mathbf{r}) = \sum_{\lambda} V_{\lambda}^{ab} F^{\lambda}(\mathbf{r})$$

For a given product index  $\lambda$  only few  $V_{\lambda}^{ab}$  will be non zero.

# Construction of $\chi_0$ in $O(N^2N_{\omega})$ operations<sup>3</sup>

• Ansatz  $f^a(\mathbf{r})f^b(\mathbf{r}) = \sum_{\mu} V_{\mu}^{ab} F^{\mu}(\mathbf{r})$  leads to

$$\chi_{\mu\nu}^{0}(\omega) = \sum_{E<0,F>0} \sum_{pqrs} X_{p}^{E} X_{q}^{F} X_{r}^{E} X_{s}^{F} \frac{V_{\mu}^{pq} V_{\nu}^{rs}}{\omega - (E-F) + i\varepsilon}$$

• Spectral function  $a_{\mu\nu} = \text{Im}\chi^0_{\mu\nu}$ 

$$a_{\mu\nu}(\lambda) = \sum_{E<0,F>0} \delta(\lambda - (E-F)) \sum_{pqrs} X_q^F X_s^F X_p^E X_r^E V_{\mu}^{pq} V_{\nu}^{rs}$$

 $ightharpoonup a_{\mu\nu}(\lambda)$  via convolution

$$\begin{aligned} \textbf{\textit{a}}_{\mu\nu}(\lambda) &= \sum_{F} \sum_{pqrs} \; \rho_{-}^{qs}(-F) * \rho_{+}^{pr}(\lambda - F) V_{\mu}^{pq} V_{\nu}^{rs} \\ \text{electronic spectral function} \; \rho_{-}^{qs}(F) &= \sum_{\nu < 0} \delta(\nu - F) X_{q}^{F} X_{s}^{F} \end{aligned}$$

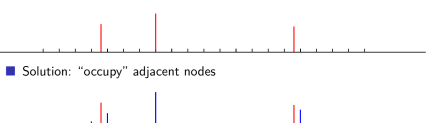
▶ Discretise  $\lambda = E - F \Rightarrow \chi^0_{\mu\nu}(\omega)$  is again convolution

$$\chi_{\mu\nu}^{0}(\omega) = \sum_{\lambda} \mathbf{a}_{\mu\nu}(\lambda) \cdot (\omega - \lambda + i\varepsilon)^{-1}$$

<sup>&</sup>lt;sup>3</sup>D. Foerster, P. Koval, J. Chem. Phys. **131**, 044103 (2009)

#### Spectral function: discretisation

► FFT needs equidistant grid, but energy differences do not fit any equidistant grid



✓ Result:  $\chi_0$  exact up to discretisation

### Spectral function: second window

- ▶ Spectral range of DFT eigenenergies is large:  $\approx 110$  eV for benzene
- ▶ Only low energy relevant  $\Rightarrow$  need  $\chi_0(\omega)$  in a small target window
- $\triangleright$   $a(\lambda)$  outside of target window contributes to  $\chi_0(\omega)$  inside

$$\chi_0(\omega) = \int_{-\infty}^{\infty} d\lambda \frac{\mathsf{a}(\lambda)}{\omega - \lambda}$$

Solution: resonant and non-resonant spectral functions

$$\chi_0(\omega) = \int_{-\infty}^{-\omega_{\max}} d\lambda \frac{b(\lambda)}{\omega - \lambda + i\varepsilon} + \int_{-\omega_{\max}}^{+\omega_{\max}} d\lambda \frac{a(\lambda)}{\omega - \lambda + i\varepsilon} + \int_{\omega_{\max}}^{+\infty} d\lambda \frac{b(\lambda)}{\omega - \lambda + i\varepsilon}$$
Nonresonant range

Resonant range

Nonresonant range

Nonresonant range

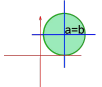
# Coulomb self energy $\Sigma_H$ : sketch of basis

▶ In the basis of dominant products  $\Sigma_H$  reads

$$\Sigma_{\mathrm{H}}^{\mu\nu} = \int d\mathbf{r}d\mathbf{r}' \;\; F^{\mu}(\mathbf{r}) \; |\mathbf{r} - \mathbf{r}'|^{-1} \; F^{\nu}(\mathbf{r}').$$

Dominant functions  $F^{\mu}(\mathbf{r})$  are either local or bilocal.





- lacksquare Generally  $F^{\mu}(m{r}) = \sum_j F^{\mu}_j(r') S_{jm}(m{R}_{\mu}m{r}'), \; m{r}' = m{r} m{C}_{\mu}.$
- ▶ Radial functions  $F_i^{\mu}(r')$  are given on a logarithmic grid.

# Coulomb self energy $\Sigma_H$ : method

 $ightharpoonup \Sigma_{
m H}^{\mu 
u}$  can be reduced to a sum over elementary Coulomb interactions

$$E_{jm,j'm'}(\boldsymbol{c},\boldsymbol{c}') = \int d\boldsymbol{r}d\boldsymbol{r}' \frac{g_{jm}(\boldsymbol{r}-\boldsymbol{c})g_{j'm'}(\boldsymbol{r}'-\boldsymbol{c}')}{|\boldsymbol{r}-\boldsymbol{r}'|}$$

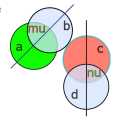
$$g_{jm}(\boldsymbol{r}) = g_{j}(r)S_{jm}(\boldsymbol{r})$$

- $ightharpoonup E_{jm,j'm'}(\boldsymbol{c},\boldsymbol{c}')=\langle g_{jm}|p^{-2}|g_{j'm'}
  angle$  in momentum space.
- Conversion to momentum space by Talman's fast Bessel transform<sup>4</sup>.

<sup>&</sup>lt;sup>4</sup>Talman J. D., Comput. Phys. Commun. **180** 332 (2009)

# Coulomb self energy $\Sigma_H$ : CPU cost

- For small molecules  $\Sigma_{
  m H}^{\mu 
  u}$  costs  $O(N^2)$
- For large molecules  $\Sigma_{\rm H}^{\mu\nu}$  costs O(N), because non overlapping  $F^{\mu}(\mathbf{r})$ ,  $F^{\nu}(\mathbf{r})$  interact via their multipoles.



✓ No 4-center integrals involved

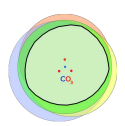
# Exchange-correlation self energy $\Sigma_{\rm xc}$

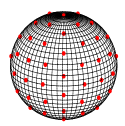
ALDA: 
$$f_{xc}(\mathbf{r}) \equiv \delta(t - t')\delta(\mathbf{r} - \mathbf{r}')\frac{dV_{xc}}{dn}(\mathbf{r})$$
  

$$\Rightarrow \Sigma_{xc}^{\mu\nu} = \int d\mathbf{r} \ F^{\mu}(\mathbf{r})f_{xc}(\mathbf{r})F^{\nu}(\mathbf{r})$$

- Spherical coordinates  $\int d\mathbf{r} = \int r^2 dr d\Omega$
- Lebedev method for solid angle  $\int d\Omega$ , Gauss-Legendre for  $\int r^2 dr$ .
- ► Organize calculation in batches.

 $\checkmark \Sigma_{xc}$  costs O(N).





#### Polarizability: CPU costs

▶ Non interacting response  $\chi_0$ 

 $O(N^2N_{\omega})$ 

▶ Self energies  $\Sigma_{\rm H}$ ,  $\Sigma_{\rm xc}$ 

` '

O(N)

▶ Solving  $\chi^{-1} = \chi_0^{-1} - \Sigma$  with direct methods

- $O(N^3N_{\omega})$
- © Fortunately  $P^{ik} = d^i \chi d^k$  can be calculated iteratively
- ► Thanks to iterative methods

$$O(N^2N_{\rm iter}N_{\omega})$$

✓ Interacting polarizability  $P^{ik}$ , total CPU

 $O(N^2N_{\omega})$ 

### Polarizability: method

$$P^{ik} = d^i \frac{1}{1 - \gamma_0 \Sigma} \chi_0 d^k \Rightarrow \text{avoid } O(N^3) \text{ inversion.}$$

▶ Bi-orthogonal Lanczos method for non-hermitian matrices

1. 
$$A = 1 - \chi_0 \Sigma \rightarrow \text{tridiagonal form } t_{nm}$$
  $A = |n\rangle t_{nm} \langle m|$ 

2. 
$$A^{-1}$$
 is easy because  $t_{nm}$  is tridiagonal  $A^{-1} = |n\rangle t_{nm}^{-1} \langle m|$ 

3. Interacting polarizability reads 
$$P = P_0 t_{11}^{-1}$$

▶ GMRES allows to control precision during iterations

1. Solve by GMRES 
$$(1-\chi_0\Sigma)x=\chi_0 d_i$$

2. Compute polarizability 
$$P^{ii} = \langle d_i | x \rangle$$

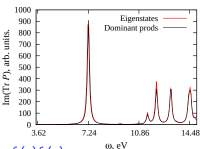
3. Mean polarizability is trace of  $P^{ii}$ .

# Dominant products basis: compression

How many dominant products  $F^{\mu}(\mathbf{r})$  needed?

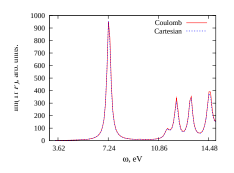
- ▶ Compare with exact result in the conventional basis.
- $\chi^0_{EF,PQ}$  is diagonal ©, but interaction kernel costs  $O(N^3)$  ©

interaction kernel K via self energy  $\Sigma$ , vertex V and density matrix D:  $K = (DV)\Sigma(V^TD)$ 



▶ Original basis 5832 products  $f_a(\mathbf{r})f_b(\mathbf{r})$ Dominant basis 1236 products  $F^{\mu}(\mathbf{r})$ .

#### Dominant products basis: Coulomb versus Cartesian metric



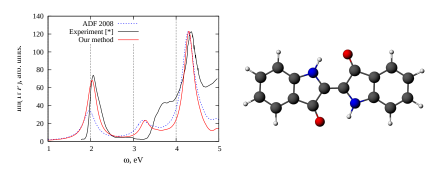


Metric	Number of products	Run time <sup>5</sup> , s	Expression for metric
Cartesian	1158	1207	$f^a(\mathbf{r})f^b(\mathbf{r})f^c(\mathbf{r})f^d(\mathbf{r})$
Coulomb	921	816.8	$f^a(\mathbf{r})f^b(\mathbf{r})f^c(\mathbf{r}')f^d(\mathbf{r}')/ \mathbf{r}-\mathbf{r}' $



<sup>&</sup>lt;sup>5</sup>(Dell laptop, one thread)

### Dominant products basis: indigo



Program	TDDFT_LR	ADF
Wall time <sup>6</sup> , hours	3.0	17

<sup>[\*]</sup> Ross Brown, IPREM unpublished (2008)

<sup>&</sup>lt;sup>6</sup>ADF with one thread on Dell laptop; TDDFT\_LR with one thread on Grid5000.

# Implementation: a bird's eye overview

▶ Input data from DFT calculation

 $H, S, f^a(\mathbf{r})$ 

▶ Diagonalize *H*, check *S* 

$$X_a^E, E$$

► Construct product basis

$$f^a(\mathbf{r})f^b(\mathbf{r}) = V_\mu^{ab}F^\mu(\mathbf{r})$$

► Compute response matrix

$$\chi_0(\mathbf{r},\mathbf{r}',\omega) = F^{\mu}(\mathbf{r})\chi^0_{\mu\nu}(\omega)F^{\nu}(\mathbf{r}')$$

...self-energies

$$\Sigma^{\mu
u} = \iint d\mathbf{r} d\mathbf{r}' F^{\mu}(\mathbf{r}) \Sigma(\mathbf{r}, \mathbf{r}') F^{
u}(\mathbf{r}')$$

...electronic excitation spectrum

$$P^{ik} = \boldsymbol{d}_i (1 - \chi_0 \Sigma)^{-1} \chi_0 \boldsymbol{d}_k$$

# Implementation: a bird's eye overview

▶ Input data from DFT calculation

 $H, S, f^a(\mathbf{r})$ 

▶ Diagonalize *H*, check *S* 

 $X_a^E, E$ 

► Construct product basis

$$f^a(\mathbf{r})f^b(\mathbf{r}) = V_\mu^{ab}F^\mu(\mathbf{r})$$

► Compute response matrix

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▶ ...self-energies

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u}(\mathbf{r}')$$

▶ ...electronic excitation spectrum

$$P^{ik} = \boldsymbol{d}_i (1 - \chi_0 \boldsymbol{\Sigma})^{-1} \chi_0 \boldsymbol{d}_k$$

# Points to mention for each step of the program

- ▶ Sketch the algorithm and implementation
- Type of parallelization chosen/implemented
- ▶ Improvements done (since June 2009)
- ▶ Wall time and speedup factors on Grid5000 / M3PEC / Dell Laptop

# Construct product basis: algorithm

#### **Algorithm 1**: Generation of bilocal vertex

- 1 forall atom pairs do
- Compute an expansion of  $f_a(\mathbf{r})f_b(\mathbf{r}) = \sum_i F_i(\mathbf{r})Y_{im}(\mathbf{r})$  (Talman)
- 3 Convert the expansion to momentum space  $F_j(r) \rightarrow F_j(p)$  (Talman)
- Build Coulomb metric g using  $F_j(p)$
- 5 Diagonalize the metric  $gX^{\lambda} = \lambda X^{\lambda}$
- 6 Build dominant products and vertex with  $\lambda=\lambda/10$
- 7 Allocate intermediate storage for dominant products and vertex
- 8 Store the generated data

#### Construct product basis: parallelization

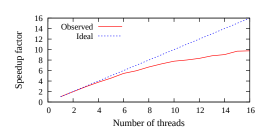
- ▶ Local vertex is a fast operation no reason to parallelize
- ightharpoonup Generation of bilocal vertices takes few seconds ( $\sim$  10 s)
- ▶ Vertices and products are needed on all nodes
- ✓ Bilocal vertices to be parallelized with OpenMP: done

#### Construct product basis: improvements

- 1. Intermediate storage introduced: storage requirements grows O(N)
- 2. Cartesian metric is implemented for testing
- 3. Loop over atom pairs is rewritten
- 4. Rotation matrices are initialized once
- 5. Talman's fast Bessel transform is reshaped in FFTW fashion
- 6. Loop over atom pairs is parallelized (OpenMP)

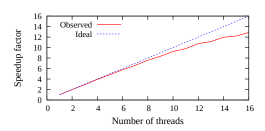
# Construct product basis: timing with benzene

OMP_NUM_THREADS	Dell laptop	Grid5000	МЗРЕС
Sequential	4.58	4.30	7.99
1	8.60	4.51	9.01
2	2.60	2.24	4.58
8	-	0.913	1.35
16	-	-	0.924
Max speedup	3.30	4.94	9.75



### Construct product basis: timing with indigo

OMP_NUM_THREADS	Grid5000	МЗРЕС
Sequential	20.7	38.8
1	21.3	42.3
2	10.8	21.3
8	2.88	5.58
16	-	3.28
Max speedup	7.40	12.9



# Compute response matrix: algorithm

#### **Algorithm 2**: Calculation of response matrix

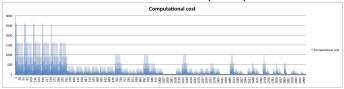
- 1 Generate Fourier transform of density matrices  $ho^\pm( au)$
- 2 forall atom quadruplets do
- 3 1. Multiply  $ho^\pm( au)$  with vertices V\_ $rho = V 
  ho^\pm( au)$
- 4 2. Find Fourier transform of spectral function  $a(\tau) = V_{rho} \cdot V_{rho}$
- 5 Do steps 1 and 2 for  $\omega_{\mathsf{max}\,2}$  compute spectral function b( au)
- 6 Nullify  $b(\tau)$  in first (target) window
- 7 Prepare second convolution  $\chi^0(\tau) = a(\tau) * FFT(\omega \lambda + i\varepsilon)$
- 8 Fourier transform of response  $\chi^0(\omega) = FFT(\chi^0(tau))$
- Interpolation of second window response  $\chi_2^0(\omega)$ , update  $\chi^0(\omega)$
- 10 Store the generated data

# Compute response matrix: parallelization

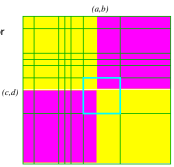
- ▶ Response matrix takes much memory (currently 11 GB for indigo)
- ▶ Although  $O(N^2N_{\omega})$ , calculation is long (3 hours for indigo)
- ► Calculation of matrix elements is independent task
- ► Available machines contain few CPU on one node (16 on M3PEC)
- Available machines consists of many nodes (hundreds)
- ► Response matrices to be parallelized with OpenMP and MPI

# Compute response matrix: parallelization

▶ Amount of work is different for atom quadruplets: load balancing



- ▶ Reorder quadruplets to obtain a good schedule for OpenMP
  - Iterative method requires matrix-vector products
  - ► There are MPI parallelized BLACS implementations: use them
  - ► Give up the "natural" blocks in favor of block-cyclic decomposition



# Compute response matrix: OpenMP parallelization

#### Algorithm 3: OpenMP parallelization of quadruplet loop

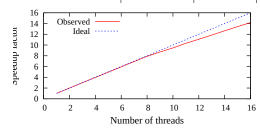
- 1 !\$OMP PARALLEL
- 2 allocate private and threadprivate variables here
- 3 !\$OMP DO SCHEDULE(DYNAMIC,10)
- 4 forall atom quadruplets do
- 5 Compute block of response matrix
- 6 Store the generated data
- 7 !SOMP END DO
- 8 deallocate private and threadprivate variables here
- 9 !\$OMP END PARALLEL
  - ✓ OpenMP parallelization is done.

# Compute response matrix: improvements

- ► Sparse data storage for vertex (less memory & more speed)
- Quadruplets of atoms introduced (half of second convolutions X)
- ► Result is saved in single precision (halves memory requirement)
- Skip unnecessary vertex-matrix multiplications in diagonal blocks X
- Automatic arrays removed
- ▶ Storage of intermediate vertex–matrix products shaped
- Single precision in vertex-matrix products
- OpenMP parallelization of the main loop
- ► Threadprivate variables for temporary arrays
- ✓ At least factor 6 speedup. OpenMP scaling on Grid5000 improved.

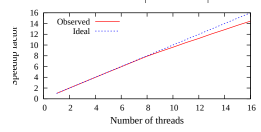
#### Compute response matrix: timing with benzene

OMP_NUM_THREADS	Dell laptop	Grid5000	МЗРЕС
Sequential	709	725	752
1	646	734	751
2	625	375	376
8	-	108	95.0
16	-	_	53.0
Max speedup	1.03	6.79	14.2



# Compute response matrix: timing with indigo

OMP_NUM_THREADS	Grid5000	МЗРЕС
Sequential	9351	10701
1	9664	10697
2	4840	5346
8	1479	1351
16	-	741
Max speedup	6.32	14.4



# Coulomb interaction matrix: algorithm

#### Algorithm 4: Calculation of Coulomb interaction matrix

- 1 Compute fast Bessel transform of all radial functions  $F_l^\lambda(p)$
- 2 Compute Wigner rotation matrices
- 3 forall atom quadruplets do
- Integrate products of radial functions with  $j_l(pR)/p^2$
- 5 Rotate contributions to Coulomb integral
- 6 Sum up the contributions from different angular momentum
  - Store the generated data

### Interaction matrices: parallelization

- ▶ Interaction matrices take less memory
- Calculation of matrix elements is independent task
- Parallelization strategy depends on speed of communication during last step (iterative construction of polarizability)

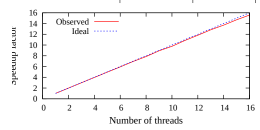
- ► Response matrices to be parallelized with OpenMP and possibly MPI
- ✓ OpenMP parallelization is done.

#### Interaction matrices: improvements

- Quadruplets of atoms introduced
- OpenMP parallelization of the main loop
- Automatic arrays removed
- Threadprivate variables for temporary arrays
- ▶ Indexing array is remove X
- ✓ Some speedup. Good scaling on Grid5000.

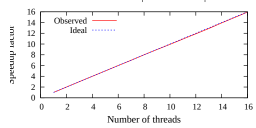
# Coulomb interaction: timing with benzene

OMP_NUM_THREADS	Dell laptop	Grid5000	МЗРЕС
Sequential	42.7	38.2	78.6
1	42.5	37.4	210
2	21.4	18.6	105
8	-	4.71	26.7
16	-	-	13.5
Max speedup	1.98	7.93	15.6



# Coulomb interaction: timing with indigo

OMP_NUM_THREADS	Grid5000	МЗРЕС
Sequential	393	757
1	365	2765
2	181	1380
8	45.6	346
16	-	174
Max speedup	8.04	15.9



# Exchange-correlation matrix: algorithm

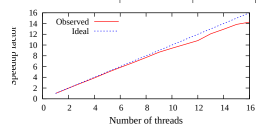
#### **Algorithm 5**: Calculation of exchange–correlation interaction matrix

- 1 Precompute rotation matrices for fast computation of  $F^{\lambda}(\mathbf{r})$
- 2 forall atom quadruplets do
- 3 Initialize batch variables; Nullify the result;
- **4 forall** *Gauss–Legendre* points  $\int dr$  **do**
- forall Lebedev points  $\int d\Omega$  do
- $\boldsymbol{\epsilon} \mid \boldsymbol{r} = (r, \Omega)$
- 7 Compute values of all dominant functions
- 8 Compute value of  $\Sigma_{x}(\mathbf{r})$  and  $\Sigma_{c}(\mathbf{r})$ 
  - Update corresponding integral
  - Store the generated data

9 10

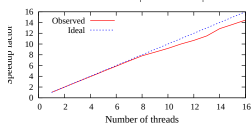
### Exchange-correlation interaction: timing with benzene

OMP_NUM_THREADS	Dell laptop	Grid5000	МЗРЕС
Sequential	79.29	97.22	141.2
1	80.45	99.20	190.4
2	41.86	49.61	96.37
8	-	13.12	24.81
16	-	-	13.34
Max speedup	1.92	7.56	14.27



# Exchange-correlation interaction: timing with indigo

OMP_NUM_THREADS	Grid5000	МЗРЕС
Sequential	365	524
1	368	583
2	186	293
8	50.9	74.7
16	-	40.3
Max speedup	7.23	13.0



# Sequential remnants in the code: indigo

Subprogram	Grid5000, 8 threads	M3PEC, 16 threads
$V_{\mu}^{ab}$	2.88	3.28
$\chi^0_{\mu u}$	1479	741
$\Sigma_{ m H}$	45.6	174
$\Sigma_{ m xc}$	50.9	40.3
$P_{ik}$	755	686
Total time	2345	1677
Remnants	11.6	32.4

#### Conclusions

- ▶ Fast algorithm allows for a  $O(N^2N_{\omega})$  scaling in true polarizability
- ✓ Most of improvements have been done in response
- ✓ Current implementation is faster than ADF in indigo molecule
- ✓ Most of the program is parallelized with OpenMP
- MPI parallelization remains to do

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- ► Mark Casida, Grenoble, France
- ▶ Ross Brown, Pau, France









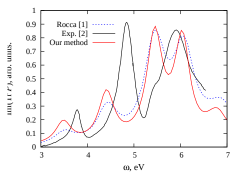




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Nouveaux Outils pour la Simulation des Solides et des Interfaces

# Largest molecule computed with our method – $C_{60}$





▶ Number of dominant products

7800

► Memory (7800\*7800/2\*8\*(128-53)/1024/1024/1024)

17 GB

▶ Walltime (Grid5000, 8 threads, 128 freq. points, for  $\chi_0$ )

7193 s

<sup>[2]</sup> Bauernschmitt et al, Am. Chem. Soc. 120, 5052 (1998) (in hexane).



<sup>[1]</sup> Dario Rocca, PhD Thesis (2007)