Range-Separated Random Phase Approximations

Bastien Mussard^{1,2} J. Toulouse¹, P. Reinhardt¹, O. Franck¹, E. Luppi¹, D. Rocca³, J. G. Ángyán³

¹ Laboratoire de Chimie Théorique, Université Pierre et Marie Curie
 ² Institut du Calcul et de la Simulation, Université Pierre et Marie Curie
 ³ Cristallographie Résonnance Magnétique et Modélisation, Université de Lorraine

bastien.mussard@upmc.fr www.lct.jussieu.fr/pagesperso/mussard/ Adiabatic connection formula for E_c

$$E_{\rm c} = \frac{1}{2} \int_0^1 d\alpha \, \operatorname{Tr} \left(\mathbb{W}_{\rm ee}.\mathbb{P}_{{\rm c},\alpha} \right)$$

$$= \int_{0}^{1} d\alpha \left(\langle \Psi_{\alpha} | \hat{W}_{ee} | \Psi_{\alpha} \rangle - \langle \Phi_{0} | \hat{W}_{ee} | \Phi_{0} \rangle \right)$$

[Langreth, Perdew PRB 15 2884 (1977)]

Fluctuation-dissipation theorem for $P_{c,\alpha}$

$$\mathbb{P}_{\mathsf{c},\alpha} = \int_{-\infty}^{\infty} \frac{-d\omega}{2\pi} \left[\mathbb{\Pi}_{\alpha}(i\omega) - \mathbb{\Pi}_{\mathsf{0}}(i\omega) \right]$$

Linear response equations for $\Pi_{\alpha}(i\omega)$ $\Pi_{\alpha}(i\omega)^{-1} = \Pi_{0}(i\omega)^{-1} - \mathbb{E}_{\mathsf{Hxc},\alpha}(i\omega)$

RPA correlation energy $f_{xc,\alpha} = 0$ or $f_{c,\alpha} = 0$

$$E_{c} = \frac{1}{2} \int_{0}^{1} d\alpha \int_{-\infty}^{\infty} \frac{d\omega}{2\pi} \operatorname{Tr} \left(\mathbb{W}_{ee} \left[\Pi_{\alpha}^{\text{RPA}}(i\omega) - \Pi_{0}(i\omega) \right] \right)$$

where $\Pi_{\alpha}^{\text{RPA}}$ is obtained from the eigenvectors of $\begin{pmatrix} \mathbf{A}_{\alpha} & \mathbf{B}_{\alpha} \\ -\mathbf{B}_{\alpha} & -\mathbf{A}_{\alpha} \end{pmatrix}$

RPA : different formalisms

$$E_{c} = \frac{1}{2} \int_{0}^{1} d\alpha \int_{-\infty}^{\infty} \frac{d\omega}{2\pi} \operatorname{Tr} \left(\mathbb{W}_{ee} \left[\Pi_{\alpha}^{\mathsf{RPA}}(i\omega) - \Pi_{0}(i\omega) \right] \right) \quad \begin{array}{l} (\mathbf{B})_{ia,jb} = \langle ab | ij \rangle \\ (\mathbf{A})_{ia,jb} = \Delta \epsilon_{ia,jb} + \langle ib | aj \rangle \end{array}$$

Analytical integration on the frequency

density matrix formulation

$$E_{\mathsf{c}} = \frac{1}{2} \int_{0}^{1} d\alpha \operatorname{Tr} \left(\mathbb{W}_{\mathsf{ee}}.\mathbb{P}_{\mathsf{c},\alpha}^{\mathsf{RPA}} \right)$$

Both analytical integration
 plasmon formulation

$$\textit{E}_{c} = \frac{1}{2} \textrm{tr} \left(\boldsymbol{\Omega}^{\mathsf{RPA}} - \boldsymbol{\Omega}^{\mathsf{TDA}} \right)$$

which is equivalent to the

rCCD formulation

 $\begin{aligned} E_{c} &= \frac{1}{2} \text{tr} \left(\textbf{B}. \textbf{T} \right) \\ \text{where } \textbf{T} \text{ are solutions of the Riccati equations} \\ \textbf{B} &+ \textbf{A}\textbf{T} + \textbf{T}\textbf{A} + \textbf{T}\textbf{B}\textbf{T} = \textbf{0} \end{aligned}$

[Scuseria, Henderson, Sorensen JCP 129 231101 (2008)]

• Analytical integration on the coupling constant dielectric matrix formulation $E_{\rm c} = \frac{1}{2} \int_{-\infty}^{\infty} \frac{d\omega}{2\pi} \operatorname{tr} \left(\log \left[\mathbf{1} - \mathbf{\Pi}_0 \mathbf{B} \right] + \mathbf{\Pi}_0 \mathbf{B} \right)$ [Jansen, BM, Rocca, Ángyán (prep.)]

RPA : different flavors

$$\begin{aligned} E_{\mathsf{c}} &= \frac{1}{2}\mathsf{tr}\left(\mathsf{B},\mathsf{T}\right) & (\mathsf{B})_{ia,jb} &= \langle ab|ij \rangle \\ \mathsf{where} \ \mathsf{B} + \mathsf{A}\mathsf{T} + \mathsf{T}\mathsf{A} + \mathsf{T}\mathsf{B}\mathsf{T} &= \mathbf{0} & (\mathsf{A})_{ia,jb} &= \Delta \epsilon_{ia,jb} + \langle ib|aj \rangle \end{aligned}$$

The Riccati equations, in this context, are derived from $\begin{pmatrix} A & B \\ -B & -A \end{pmatrix}$ which depend on a kernel \mathbf{f} that **does not or does include exchange**

► **drCCD** variant
$$\longrightarrow$$
 $\mathbb{f}_{xc} = 0 \longrightarrow$ $(\begin{array}{c} \mathsf{B}^{\mathsf{I}})_{ia,jb} = \langle ab | ij \rangle \\ (\mathsf{A}^{\mathsf{I}})_{ia,jb} = \Delta \epsilon_{ia,jb} + \langle ib | aj \rangle \end{array}$

$$\Xi_{c}^{dRPA-I} = \frac{1}{2} tr\left(\mathbf{B}^{I}.\mathbf{T}^{dRPA}\right)$$

• **rCCD** variant
$$\longrightarrow$$
 $\mathbb{F}_c = 0 \longrightarrow$

$$E_{c}^{\text{RPAx-II}} = \frac{1}{4} \text{tr} \left(\mathbf{B}^{\text{II}} \cdot \mathbf{T}^{\text{RPAx}} \right)$$

where
$$\mathbf{B}^{\mathsf{I}} + \mathbf{A}^{\mathsf{I}}\mathbf{T} + \mathbf{T}\mathbf{A}^{\mathsf{I}} + \mathbf{T}\mathbf{B}^{\mathsf{I}}\mathbf{T} = \mathbf{0}$$

$$(\mathbf{B}^{II})_{ia,jb} = \langle ab || ij \rangle \\ (\mathbf{A}^{II})_{ia,jb} = \Delta \epsilon_{ia,jb} + \langle ib || aj \rangle$$

where
$$\mathbf{B}^{II} + \mathbf{A}^{II}\mathbf{T} + \mathbf{T}\mathbf{A}^{II} + \mathbf{T}\mathbf{B}^{II}\mathbf{T} = \mathbf{0}$$

Additionnal flavors

$$E_{c}^{SOSEX} = \frac{1}{2} tr \left(\mathbf{B}^{II} \cdot \mathbf{T}^{dRPA} \right)$$
$$E_{c}^{SO2} = \frac{1}{2} tr \left(\mathbf{B}^{I} \cdot \mathbf{T}^{RPAx} \right)$$

Note that comparable additional flavors also exist in the **density matrix** formulation (they have been explored in the litterature) and in the **dielectric matrix** formulation.

Full-range DFT + RPA

DFT calculation followed by an RPA calculation using KS orbitals and energies

$$\begin{split} E^{\mathsf{DFT}} &= \min_{\Phi} \left\{ \langle \Phi | \, \hat{\mathcal{T}} + \hat{V}_{\mathsf{ne}} | \Phi \rangle + E_{\mathsf{Hxc}}[n_{\Phi}] \right\} \\ E &= \langle \Phi | \, \hat{\mathcal{T}} + \hat{V}_{\mathsf{ne}} | \Phi \rangle + E_{\mathsf{Hx}}^{\mathsf{HF}}[\Phi] + E_{\mathsf{c}}^{\mathsf{RPA}} \end{split}$$

- DFT (LDA/GGA/...) do not describe well the long-range dispersion forces, while RPA does
- Still problems with : short-range correlation energies (far too negative) strong dependence on basis size simple van der Waals dimers



Range-Separated Hybrid + RPA

▶ The idea : split the e-e interaction into a short- and a long-range parts

Hybrid **DFT** with exact HF exchange and RPA correlation, both at **long-range** : $E = \min_{\Phi} \left\{ \langle \Phi | \hat{T} + \hat{V}_{ne} + \hat{W}_{ee}^{lr} | \Phi \rangle + E_{Hxc}^{sr} [n_{\Phi}] \right\} + E_{c}^{lr,RPA}$

[Stoll,Savin DF.meth.phys. 177 (1985)][Savin IJQC 22 59 (1988)][Savin, Rec.dev. 327 (1996)]

- range separation greatly improves RPA (see srPBE+RPAx-SO2)
- basis dependence is reduced



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- range separation greatly improves RPA (see srPBE+RPAx-SO2)
- basis dependence is reduced
- ▶ for ex. : performance on the S22 dataset is very good for srPBE+RPAx-SO2



>srPBE, aug-cc-pVDZ, $\mu = 0.5$ [Toulouse,Zhu,Savin,Jansen,Ángyán 135 084119 (2011)]

Wavefunction basis convergence [Gori-Giorgi,Savin PRA 73 032506 (2006)] both partial wave and principal number expansions lead to exponential convergence of the wavefunction



Extrapolation formula for total energies We propose a **three-point extrapolation scheme** for total RSH+correlation energies $(E_X = E_{\infty} + B \exp(-\beta X))$

$$E_{\infty} = E_{XYZ} = \frac{E_Y^2 - E_X E_Z}{2E_Y - E_X - E_Z}$$

>errors (mHartree) wrt. cc-pV6Z

	He	Ne	N_2	H_2O
$\Delta E_{\rm D}$	8.488	74.523	51.581	55.850
$\Delta E_{\rm T}$	0.781	20.337	13.406	14.736
$\Delta E_{\rm Q}$	0.245	5.763	4.090	4.499
ΔE_5	0.078	0.751	0.810	0.726
$\Delta E_{\rm DTQ}$	0.205	0.401	1.083	1.105
ΔE_{TQ5}	0.002	-1.876	-0.972	-1.475

Spin-Unrestricted RPA

Implemented most of the variants with a nospinflip/spinflip block structure.

Normal distributions of errors (kcal/mol) of calculations on **AE49**

- mean errors of post-RSH calculation are better
- post-RSH calculations dist. of errors have sharper distributions
- Similar results were obtained on the **DBH24/08** dataset

We argue that **RPAx-SO2** is a good method for a wide range of applications



Study of systems with a non-integer charge

$$E^{N+\delta} = \min_{\hat{\Gamma} \to N+\delta} \operatorname{Tr} \left[\hat{\Gamma} \left(\hat{T} + \hat{V}_{ext} + \hat{V}_{ee} \right) \right]$$

$$\hat{\Gamma}^{N+\delta} = (1-\delta)\hat{\Gamma}^{N} + (\delta)\hat{\Gamma}^{N+1}$$

$$E^{N+\delta} = (1-\delta)E^{N} + (\delta)E^{N+1}$$

$$n^{N+\delta} = (1-\delta)n^{N} + (\delta)n^{N+1}$$

E is **piecewise linear** wrt *N* and has a **derivative discontinuity** at integer *N*

$$E^{N+\delta} = \min_{\hat{\Gamma} \to N+\delta} \operatorname{Tr} \left[\hat{\Gamma} \left(\hat{T} + \hat{V}_{ext} \right) + E_{Hxc}[n_{\hat{\Gamma}}] \right]$$

$$\hat{\Gamma}_{s}^{N+\delta} = (1-\delta) \hat{\Gamma}_{s}^{N,\delta} + (\delta) \hat{\Gamma}_{n}^{N+1,\delta}$$

$$n^{N+\delta}(\mathbf{r}) = \sum f_{i} |\phi_{i}^{N+\delta}(\mathbf{r})|^{2}$$

[Mori-Sánchez, Cohen, Yang JCP 125 2006]



Fractional Occupation Numbers calculations

Study of systems with a non-integer spin

 Systems with fractional occupation of degenerate spin states should have the same energy as the integer-spin states



0.5↑0.5↓

 $0.5 \uparrow 0.5$,

Constancy condition

 $E\left[\sum c_i n_i\right] = E[n] = E(N)$

[Cohen, Mori-Sánchez, Yang JCP 129 2008]



Fractional Occupation Numbers calculations [early results]







 $> \! {\rm deviation}$ from linear extrapolation for He with $1+\delta$ electrons, cc-pVDZ

PBE+RPA srPBE+lrRPA PBE+SO2 srPBE+lrSO2 PBE+SOSEX srPBE+lrSOSEX



Fractional Occupation Numbers calculations [early results]



- All (and more) is implemented in the Quantum Chemistry Package MOLPRO Forces for (RSH+)RPA energies are available and so are geometry optimizations at the (RSH+)RPA levels [BM,Szalay,Ángyán JCTC 10 1968 (2014)]
- Using the fractional occupations implementations to get IP and AE with double-hybrids
- Instabilities in the RPA problem
- at the Université Pierre et Marie Curie Odile Franck Eleanora Luppi Peter Reinhardt Julien Toulouse and, at the Université de Lorraine János Ángyán Dario Rocca