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OUTLINE

optimisation **Orbital** for Richardson-Gaudin wavefunctions[†]

Charles-Émile Fecteau^{a,b}, Guillaume Acke^a, Patrick Bultinck^a and Paul A. Johnson^b

Do not write an abstract. That will be done when the outline has matured into a completed paper.

Introduction 1

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Orbital-optimised doubly-occupied configuration interaction (OO-DOCI) computations yield the most precise result one can expect to obtain through a geminal approach with a given orbital basis set. In other words, for a given set of orbitals, this approach gives the exact solution to the seniority-zero coulomb Hamiltonian. Incidently, its optimised orbitals are also the optimal orbitals for any geminal-based approach.

This makes this method an very useful reference tool as we develop a variationnal Richardson-Gaudin (varRG) approach based on wavefunction ansatz made from products of Richardson-Gaudin geminals 12, both as a source for the optimal orbitals (until we develop an independent orbital-optimisation routine) and as a comparison standard for the precision we can hope to achieve.

It is therefore relevant to compute OO-DOCI dissociation curves for various small systems as a reference point. We are notably interested in the following systems, for the STO-6G and cc-pVDZ orbital basis:

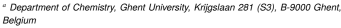
- Hydrogen polymer chains : H₄, H₆ and H₈
- Homonuclear diatomics: He2, Li2, Be2, B2, C2, N2, O2, F2 and Ne_2 (cc-pVDZ only)
- 12-electron diatomics : LiF, BeO, BN, CN⁺ and CO²⁺ (and ³⁶
- 14-electron diatomics : CN^- , CO, NO^+ and O_2^{2+} (and N_2)

Results and discussion

2.1 STO-6G

2.1.1 Hydrogen polymer chains

All hydrogen polymer chains dissociation curves were easily com- 44 30 puted and yielded smooth results.



^b Department of Chemistry, Université laval, 2325 rue de l'Université, Québec, 50 Canada, G1V 0A6

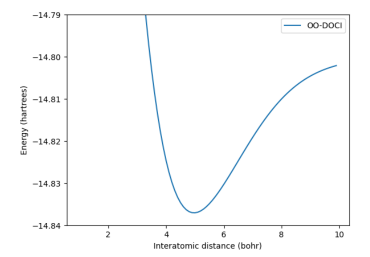


Fig. 1 OO-DOCI dissociation curve of Li_2 in the STO-6G orbital basis

2.1.2 Homonuclear diatomics

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Smooth dissociations curves have been obtained for Li_2 and F_2 (see figures 1 and 2), with preliminary results for N_2 and O_2 (see figure 4) suggesting these curves should be obtained without significant hurdles as well.

For the larger systems $(N_2, O_2 \text{ and } F_2)$, some points sometimes yielded results that layed outside the curve. However, it has generally been possible to recompute those points to get adequate results, thus these invalid results were attributed to bad convergence of the optimisation algorithm (CMA-ES, which has a stochastic nature).

The curves for Be_2 , B_2 and C_2 however, consistently produced curves that aren't completely smooth throughout dissociation (see figures 5 6 and 7). Specifically, C_2 presents an avoided crossing whereas Be presents two, and B_2 actually produces a double-well. Before moving forward, it must be determined whether these avoided crossings are to be expected for OO-DOCI results in this basis or whether they indicate a failure of the orbital optimisation to converge on the correct orbital rotation.

Various approaches have been attempted (with B_2 and C_2

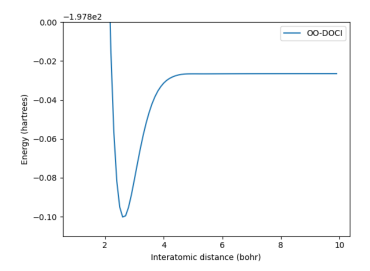


Fig. 2 OO-DOCI dissociation curve of F_2 in the STO-6G orbital basis

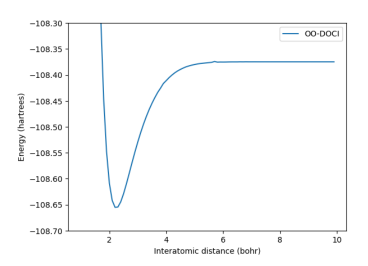


Fig. 3 OO-DOCI dissociation curve of N_2 in the STO-6G orbital basis with a badly converged point

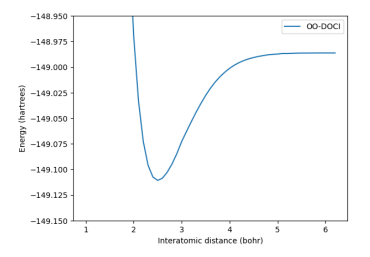


Fig. 4 Incomplete OO-DOCI dissociation curve of \mathcal{O}_2 in the STO-6G orbital basis

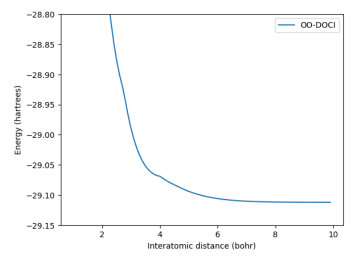


Fig. 5 OO-DOCI dissociation curve of $\it Be_2$ in the STO-6G orbital basis presenting two avoided crossings

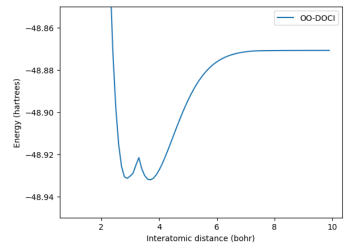


Fig. 6 OO-DOCI dissociation curve of B_2 in the STO-6G orbital basis presenting an avoided crossing in the form of a double potential well

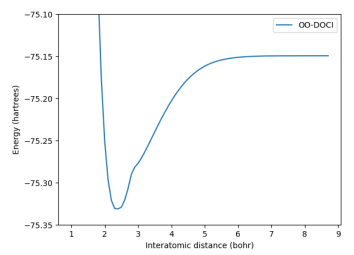


Fig. 7 OO-DOCI dissociation curve of \mathcal{C}_2 in the STO-6G orbital basis presenting an avoided crossing

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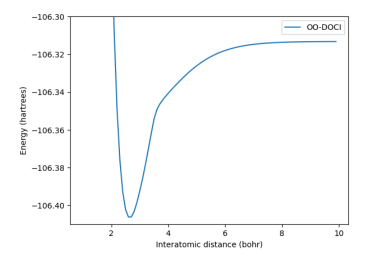


Fig. 8 OO-DOCI dissociation curve of LiF in the STO-6G orbital basis presenting an avoided crossing

specifically) in attempts to see if the optimisation's results could be improved or if the avoided crossings that were observed were indeed the optimal result (for DOCI in the given orbital basis):

- Points were computed with a significantly higher maximum number of iterations for the CMA-ES algorithm.
- Points were computed reusing the orbital rotation of the previous point in the curve, approaching from both the right and the left.
- Computations were done using DFT orbitals as an initial guess (using B3lyp, LDA, PBE and PW91 functionals).
- Computations were done using CISD natural orbitals as an initial guess.

All these approaches converged on the same results as the initial brute-force attempt. Two other approaches are being investigated before we conclude that the curves that were obtained are indeed correct:

- DOCI orbitals are being optimised based on the angles of Jacobi rotations between orbitals that share an irreductible representation within an Abelian symmetry point group of the molecule (both D_{2h} and C_{2v} for homonuclear molecules, C_{2v} for heteronuclear molecules)
- FCI natural orbitals are being computed, in order to use them as an initial guess for the DOCI orbitals.

2.1.3 12-electron diatomics

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76 Similarly to some of the homonuclear diatomics mentioned, all 12-electron dissociation curves present avoided crossings (see fig-78 ures 8, 9, 10, 11 and 12).

2.1.4 14-electron diatomics

Preliminary results indicate a smooth curve dissociation curve for 14-electron systems, although some curves present outlying points that require recomputation (see figures 13, 14, 15 and 16).

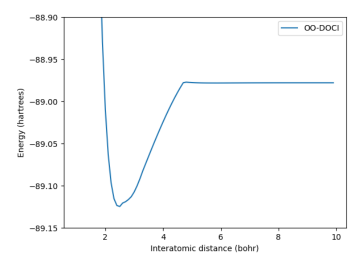


Fig. 9 OO-DOCI dissociation curve of ${\it BeO}$ in the STO-6G orbital basis presenting two avoided crossings

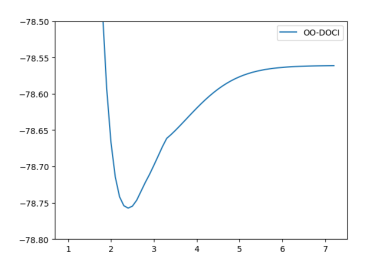


Fig. 10 Incomplete OO-DOCI dissociation curve of *BN* in the STO-6G orbital basis presenting an avoided crossing

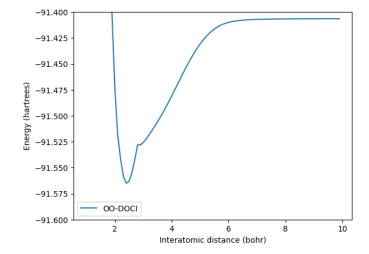


Fig. 11 OO-DOCI dissociation curve of ${\it CN}^+$ in the STO-6G orbital basis presenting an avoided crossing

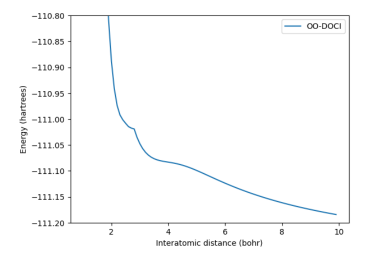


Fig. 12 OO-DOCI dissociation curve of CO^{2+} in the STO-6G orbital basis presenting an avoided crossing

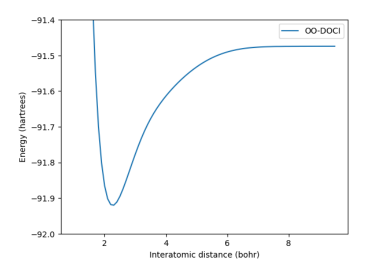


Fig. 13 Incomplete OO-DOCI dissociation curve of CN^- in the STO-6G orbital basis

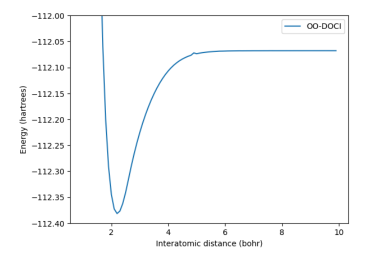


Fig. 14 OO-DOCI dissociation curve of ${\it CO}$ in the STO-6G orbital basis with a badly converged point

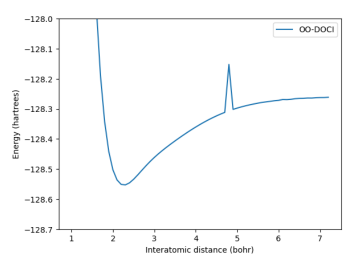


Fig. 15 Incomplete OO-DOCI dissociation curve of NO^+ in the STO-6G orbital basis with some badly converged points

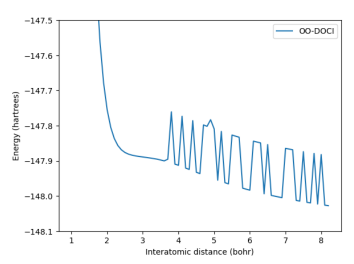


Fig. 16 Incomplete OO-DOCI dissociation curve of ${\cal O}_2^{2^+}$ in the STO-6G orbital basis with some badly converged points

83 2.2 cc-pVDZ

84 2.2.1 Hydrogen polymer chains

- Computations underway. Preliminary results seem to produce a
- 86 smooth dissociation curves.

87 2.2.2 Homonuclear diatomics

- 88 Computations underway.
- 89 2.2.3 12-electron diatomics
- 90 Computations underway.

2.2.4 14-electron diatomics

92 Computations underway.

One issue that arises when computing these dissociation curves
(as well as those for N_2 , O_2 , F_2 and Ne_2) is that the maximal
computation time allowed on some compute canada clusters is 7
days, but the computation of even a single point for systems with
14 electrons or more in the cc-pVDZ orbital basis takes more time
than that. The cedar cluster allows for longer computations, but
possibility of segmenting CMA-ES optimisation for a single
point should be investigated and implemented if possible.

101 3 Conclusions

102 4 Acknowledgements

103 5 Appendix

Notes and references

- 1 P. A. Johnson, C.-E. Fecteau, F. Berthiaume, S. Cloutier, L. Carrier, M. Gratton, P. Bultinck, S. De Baerdemacker, D. Van Neck,
 P. Limacher, and P. W. Ayers, Journal of Chemical Physics 153,
 104110 (2020).
- 2 C.-E. Fecteau, S. Cloutier, J.-D. Moisset, J. Boulay, and P. A.
 Johnson, Journal of Chemical Physics 156, 194103 (2022).