**Emmanuel FROMAGER – Publications (updated September 2024)**

[ResearcherID: M-4727-2017](http://www.researcherid.com/rid/M-4727-2017) [Google Scholar](https://scholar.google.fr/citations?user=6S8Or18AAAAJ&hl=fr)

[Publi. 55:](https://doi.org/10.1063/5.0229787) “Fragment quantum embedding using the Householder transformation: A multi-state extension based on ensembles”- F. Cernatic, E. Fromager, and S. Yalouz, *J. Chem. Phys.* **161**, 124107 (2024). Preprint: [arXiv:2407.14278](https://doi.org/10.48550/arXiv.2407.14278)

[Publi. 54:](https://journals.aps.org/prb/accepted/e7070O19A2d15842e4255bb4a1626a9611c402cd3) “Neutral electronic excitations and derivative discontinuities: An extended N-centered ensemble density functional theory perspective”- F. Cernatic, P.-F. Loos, B. Senjean, and E. Fromager, *Phys. Rev. B* **109**, 235113 (2024). Preprint: [arXiv:2401.04685](https://doi.org/10.48550/arXiv.2401.04685)

[Publi. 53:](https://doi.org/10.1088/2516-1075/ad45d5) “Density functional theory beyond the Born–Oppenheimer approximation: exact mapping onto an electronically non-interacting Kohn–Sham molecule”- E. Fromager and B. Lasorne, 2024 *Electron. Struct.* **6** 025002. Preprint: [arXiv:2312.15080](https://doi.org/10.48550/arXiv.2312.15080)

[Publi. 52:](https://doi.org/10.1002/jcc.27387) “Extended N-centered ensemble density functional theory of double electronic excitations”- F. Cernatic and E. Fromager, *J. Comput. Chem.* 2024;45:1945–1962. Preprint: [arXiv:2402.07161](https://doi.org/10.48550/arXiv.2402.07161)

[Publi. 51:](https://doi.org/10.1103/PhysRevB.108.155119) “Unitary transformations within density matrix embedding approaches: A perspective on the self-consistent scheme for electronic structure calculation”- Q. Marécat, B. Lasorne, E. Fromager, and M. Saubanère *Phys. Rev. B* **108**, 155119 (2023), Preprint: [arxiv:2306.07641](https://arxiv.org/abs/2306.07641)

[Publi. 50:](https://doi.org/10.1063/5.0157746) “A unified density-matrix functional construction of quantum baths in density matrix embedding theory beyond the mean-field approximation”- S. Sekaran, O. Bindech, and E. Fromager, *J. Chem. Phys.* **159**, 034107 (2023), Preprint: [arXiv:2304.14729](https://arxiv.org/abs/2304.14729)

[Publi. 49:](https://doi.org/10.1063/5.0125683) “Quantum embedding of multi-orbital fragments using the Block-Householder-transformation”- S. Yalouz, S. Sekaran, E. Fromager, and M. Saubanère, *J. Chem. Phys.* **157**, 214112 (2022), Preprint: [arXiv:2209.10302](https://arxiv.org/abs/2209.10302)

[Publi. 48:](https://doi.org/10.1039/D2CP02827A) “DFT Exchange: Sharing Perspectives on the Workhorse of Quantum Chemistry and Materials Science”- Andrew M. Teale, Trygve Helgaker, Andreas Savin, Carlo Adamo, Bálint Aradi, Alexei V. Arbuznikov, Paul W. Ayers, Evert Jan Baerends, Vincenzo Barone, Patrizia Calaminici, Eric Cancès, Emily A. Carter, Pratim Kumar Chattaraj, Henry Chermette, Ilaria Ciofini, T. Daniel Crawford, Frank De Proft, John F. Dobson, Claudia Draxl, Thomas Frauenheim,  **Emmanuel Fromager**, Patricio Fuentealba, Laura Gagliardi, Giulia Galli, Jiali Gao, Paul Geerlings, Nikitas Gidopoulos, Peter M. W. Gill, Paola Gori-Giorgi, Andreas Görling, Tim Gould, Stefan Grimme, Oleg Gritsenko, Hans Jørgen Aagaard Jensen, Erin R. Johnson, Robert O. Jones, Martin Kaupp, Andreas M. Köster, Leeor Kronik, Anna I. Krylov, Simen Kvaal, Andre Laestadius, Mel Levy, Mathieu Lewin, Shubin Liu, Pierre-François Loos, Neepa T. Maitra, Frank Neese, John P. Perdew, Katarzyna Pernal, Pascal Pernot, Piotr Piecuch, Elisa Rebolini, Lucia Reining, Pina Romaniello, Adrienn Ruzsinszky, Dennis R. Salahub, Matthias Scheffler, Peter Schwerdtfeger, Viktor N. Staroverov, Jianwei Sun, Erik Tellgren, David J. Tozer, Samuel B. Trickey, Carsten A. Ullrich, Alberto Vela, Giovanni Vignale, Tomasz A. Wesolowski, Xin Xu, and  Weitao Yang, *Phys. Chem. Chem. Phys. (2022*) **24**, 28700-28781, Advance Article, DOI: 10.1039/d2cp02827a **[2022 HOT PCCP article:** [link](https://pubs.rsc.org/en/Journals/ArticleCollectionLanding?themeId=e3d7e371-bed8-4ccd-bd58-1d9ee83cec7e&journalName=)**]**, Preprint: [10.26434/chemrxiv-2022-13j2v](https://chemrxiv.org/engage/chemrxiv/article-details/62974da519595958f0bcc339)

[Publi. 47:](https://doi.org/10.1103/PhysRevA.106.032203) “Reduced density matrix functional theory from an ab initio seniority-zero wave function: Exact and approximate formulations along adiabatic connection paths”- B. Senjean, S. Yalouz, N. Nakatani, and E. Fromager, *Phys. Rev. A* **106**, 032203 (2022), Preprint: [arXiv:2204.00699](https://arxiv.org/abs/2204.00699)

[Publi. 46:](https://doi.org/10.3390/computation10030045) “Local Potential Functional Embedding Theory: A Self-Consistent Flavor of Density Functional Theory for Lattices without Density Functionals”- S. Sekaran, M. Saubanère, and E. Fromager, *Computation* **2022**, *10*, 45. **[invited paper in the special issue of *Computation* in honour of Karlheinz Schwarz on the occasion of his 80th birthday]** Preprint: [arXiv:2202.08071](https://arxiv.org/abs/2202.08071)

[Publi. 45:](https://doi.org/10.1007/s41061-021-00359-1) “Ensemble Density Functional Theory of Neutral and Charged Excitations”- F. Cernatic, B. Senjean, V. Robert, and E. Fromager, *Top Curr Chem (Z)* **380**, 4 (2022), ***review article in the “New Horizon in Computational Chemistry Software” topical collection.*** Also published as a [book chapter](https://doi.org/10.1007/978-3-031-07658-9_8), preprint: [arXiv:2109.04943](https://arxiv.org/abs/2109.04943)

[Publi. 44:](https://doi.org/10.1103/PhysRevB.104.035121) “Householder transformed density matrix functional embedding theory”- S. Sekaran, M. Tsuchiizu, M. Saubanère, and E. Fromager, *Phys. Rev. B* **104**, 035121 (2021), Preprint: [arXiv:2103.04194](https://arxiv.org/abs/2103.04194)

[Publi. 43:](https://doi.org/10.1103/PhysRevA.103.012806) “Exact exchange-correlation potentials for calculating the fundamental gap with a fixed number of electrons”- M. J. P. Hodgson, J. Wetherell, and E. Fromager, *Phys. Rev. A* **103**, 012806 (2021), Preprint: [arXiv:2010.05642](https://arxiv.org/abs/2010.05642)

[Publi. 42:](https://doi.org/10.1039/D0FD90026E) “New approaches to study excited states in density functional theory: general discussion”- Jan Gerit Brandenburg, Kieron Burke, Emmanuel Fromager, Matteo Gatti, Sara Giarrusso, Nikitas I Gidopoulos, Paola Gori-Giorgi, Duncan Gowland, Trygve Helgaker, Matthew JP Hodgson, Lionel Lacombe, Gianluca Levi, Pierre-François Loos, Neepa T Maitra, Eduardo Maurina Morais, Nisha Mehta, Filippo Monti, Manasi R Mulay, Katarzyna Pernal, Lucia Reining, Pina Romaniello, Matthew R Ryder, Andreas Savin, Dumitru Sirbu, Andrew M Teale, Alex JW Thom, Donald G Truhlar, Jack Wetherell, Weitao Yang, *Faraday Discuss.*, 2020, **224**, 483-508.

[Publi. 41:](https://doi.org/10.1039/D0FD00059K) “Weight Dependence of Local Exchange-Correlation Functionals in Ensemble Density-Functional Theory: Double Excitations in Two-Electron Systems”- C. Marut, B. Senjean, E. Fromager, and P.-F. Loos,  *Faraday Discuss.*, 2020, **224**, 402-423, Preprint: [arXiv:2005.06159](https://arxiv.org/abs/2005.06159)

[Publi. 40:](https://doi.org/10.1039/D0FD90025G) “Strong correlation in density functional theory: general discussion”- Emmanuel Fromager, Nikitas Gidopoulos, Paola Gori-Giorgi, Trygve Helgaker, Pierre-François Loos, Thomas Malcomson, Katarzyna Pernal, Andreas Savin, Donald G Truhlar, Meilani Wibowo, Weitao Yang, *Faraday Discuss.*, 2020, **224**, 373-381.

[Publi. 39:](https://doi.org/10.1039/D0FD90023K) “New density-functional approximations and beyond: general discussion”- Jan Gerit Brandenburg, Kieron Burke, Antonio Cancio, Jannis Erhard, Emmanuel Fromager, Abhisek Ghosal, Nikitas Gidopoulos, Paola Gori-Giorgi, Trygve Helgaker, Ben Hourahine, Christoph R Jacob, Derk Kooi, Neepa Maitra, Manasi R Mulay, Katarzyna Pernal, Aurora Pribram-Jones, Lucia Reining, Pina Romaniello, Matthew R Ryder, Andreas Savin, Chris-Kriton Skylaris, Andrew M Teale, David Tozer, Donald G Truhlar, Weitao Yang, *Faraday Discuss.*, 2020, **224**, 166-200.

[Publi. 38:](https://doi.org/10.1063/5.0007388) “A weight-dependent local correlation density-functional approximation for ensembles” - P.-F. Loos and E. Fromager, *J. Chem. Phys.* **152**, 214101 (2020), **JCP Editor's Pick**, Preprint:[arXiv:2003.05553](https://arxiv.org/abs/2003.05553)

[Publi. 37:](https://link.aps.org/doi/10.1103/PhysRevLett.124.243001) “Individual Correlations in Ensemble Density Functional Theory: State- and Density-Driven Decompositions without Additional Kohn-Sham Systems” - E. Fromager, *Phys. Rev. Lett.***124**, 243001 (2020), Preprint: [arXiv:2001.08605](https://arxiv.org/abs/2001.08605) [[Supplemental Material](https://quantique.u-strasbg.fr/lib/exe/fetch.php?media=fr:pageperso:ef:supp_mat_for_revision.pdf)]

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