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Molecular Finite Element Density Functional Calculations employing a Cusp Factor to enable convergence at the nuclei

FEM calculations have been performed in Cartesian coordinates in the finite element framework FEniCS[1], using the density functional approach for a number of small molecules. In order to aid convergence of the orbitals and total energies a suitable cusp factor

$F(\mathbf{r}) = 1 + \sum_{j=1}^{N_A} c_j \exp(-2Z_j r_j)$ with $r_j = |\mathbf{r} - \mathbf{R}_j|$ was employed, such that the resulting effective potential is non-singular at all nuclei and where the coefficients c_i are obtained by solving a linear system of equations. The finite element ansatz for the pseudo orbitals leads to a sparse generalized eigenvalue problem of dimension N up to $3.6 \cdot 10^6$, which was solved employing the Jacobi-Davidson method on a High Performance SMP Machine with 32 CPUs and 512 GB of memory.

The resulting total energies and densities were compared with those obtained using the Gaussian basis set package NWChem[2] and excellent agreement was found.

[1] Martin S. Alnæs, Jan Blechta, Johan Hake, August Johansson, Benjamin Kehlet, Anders Logg, Chris Richardson, Johannes Ring, Marie E. Rognes, and Garth N. Wells. The fenics project version 1.5. *Archive of Numerical Software*, 3(100), 2015.

[2] M. Valiev, E.J. Bylaska, N. Govind, K. Kowalski, T.P. Straatsma, H.J.J. Van Dam, D. Wang, J. Nieplocha, E. Apra, T.L. Windus, and W.A. de Jong. Nwchem: A comprehensive and scalable open-source solution for large scale molecular simulations. *Computer Physics Communications*, 181(9):1477 – 1489, 2010.

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