

¹ Den2Obj: A command line tool for producing ² isosurfaces from electron density data files

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⁶ Summary

⁷ Electron density plays a foundational role in electronic structure calculations. It offers a⁸
⁹ detailed spatial representation of the distribution of electrons in molecules and materials. In¹⁰
¹¹ computational methodologies, most notably Density Functional Theory (DFT), electron density¹²
¹³ serves as a central quantity from which key properties such as total energy, electrostatic¹⁴
¹⁵ potential, and atomic forces are derived. The analysis of electron density provides significant¹⁶
¹⁷ insights into chemical bonding, reactivity, and electron redistribution, especially during bond¹⁸
¹⁹ formation and dissociation. This capability enables the prediction and interpretation of the²⁰
²¹ properties of materials and molecular systems.

²² Electron density is mathematically represented as a scalar field. It assigns a scalar value,²³
²⁴ corresponding to the probability density of finding an electron, to each point in three-dimensional²⁵
²⁶ space. Due to the complexity and high dimensionality of such data, specialized visualization²⁷
²⁸ techniques are necessary for meaningful interpretation. Common methods include contour²⁹
³⁰ plotting, isosurface generation, and volumetric rendering. These techniques facilitate the³¹
³² exploration of electronic behavior and structural characteristics.

³³ This utility, Den2Obj, is a C++ based command line program developed to generate isosurfaces³⁴
³⁵ from electron density scalar fields. It supports input formats that are commonly used in³⁶
³⁷ electronic structure calculations, including CHGCAR and PARCHG files produced by VASP, as³⁸
³⁹ well as Gaussian Cube files. Den2Obj is capable of converting these formats into a native .d2o⁴⁰
⁴¹ file format, which achieves substantial reductions in file size through efficient compression. The⁴²
⁴³ resulting isosurfaces can be exported in several widely supported geometry file formats, including⁴⁴
⁴⁵ STL (Stereolithography), PLY (Polygon File Format), and OBJ (Wavefront). These output⁴⁶
⁴⁷ formats enable convenient integration with a broad range of visualization and post-processing⁴⁸
⁴⁹ software tools.

³⁰ Statement of need

³¹ Isosurface Visualization with Den2Obj

³² Isosurfaces play a central role in scientific research and engineering applications by providing³³
³⁴ an effective means of visualizing complex scalar fields and understanding intricate physical³⁵
³⁶ phenomena. An isosurface represents a set of points in a field where a scalar quantity, known³⁷
³⁸ as the *isovalue*, remains constant. Due to their importance, many software packages support³⁹
⁴⁰ isosurface generation, including Open Data Explorer ([Thompson et al., 2004](#)), MATLAB ([The MathWorks Inc., 2022](#)), ParaView ([Ahrens et al., 2005](#)), and VESTA ([Momma & Izumi, 2011](#)).⁴¹
⁴² These tools are primarily designed for interactive use and typically rely on graphical user⁴³
⁴⁴ interfaces.

⁴⁵ In contrast, Den2Obj is a C++-based command line tool developed for automated isosurface

⁴¹ extraction from electron density and wavefunction data. It supports input formats commonly
⁴² produced by VASP ([Hafner, 2008](#)), such as CHGCAR and PARCHG, as well as Gaussian ([Frisch et](#)
⁴³ [al., 2016](#)) Cube files. The resulting isosurfaces can be exported in widely used 3D geometry
⁴⁴ formats, including Stereolithography (.stl), Polygon File Format (.ply), and Wavefront
⁴⁵ (.obj).

⁴⁶ Isosurface generation in Den20bj is carried out using either the marching cubes algorithm
⁴⁷ ([Lorensen & Cline, 1987](#)) or the marching tetrahedra algorithm ([Burke, 1994](#)), both of which
⁴⁸ can be selected via command line arguments. These algorithms are implemented with OpenMP
⁴⁹ parallelization to leverage the performance of modern multi-core CPUs. When visualizing
⁵⁰ wavefunctions rather than electron densities, it is often useful to generate separate isosurfaces
⁵¹ for positive and negative lobes. Den20bj accommodates this by enabling dual isosurface
⁵² generation through a single command line argument.

⁵³ An illustrative example is presented in [Figure 1](#), which displays the canonical molecular orbitals
⁵⁴ of the benzene molecule. These orbitals were computed using the PyQInt program. ([Filot, 2020](#))
⁵⁵ Isosurfaces were generated via the marching cubes algorithm as implemented in Den20Obj,
⁵⁶ producing .ply files that were subsequently imported into Blender ([Blender - a 3D Modelling](#)
⁵⁷ [and Rendering Package, 2018](#)), along with the atomic coordinates of benzene, for rendering.

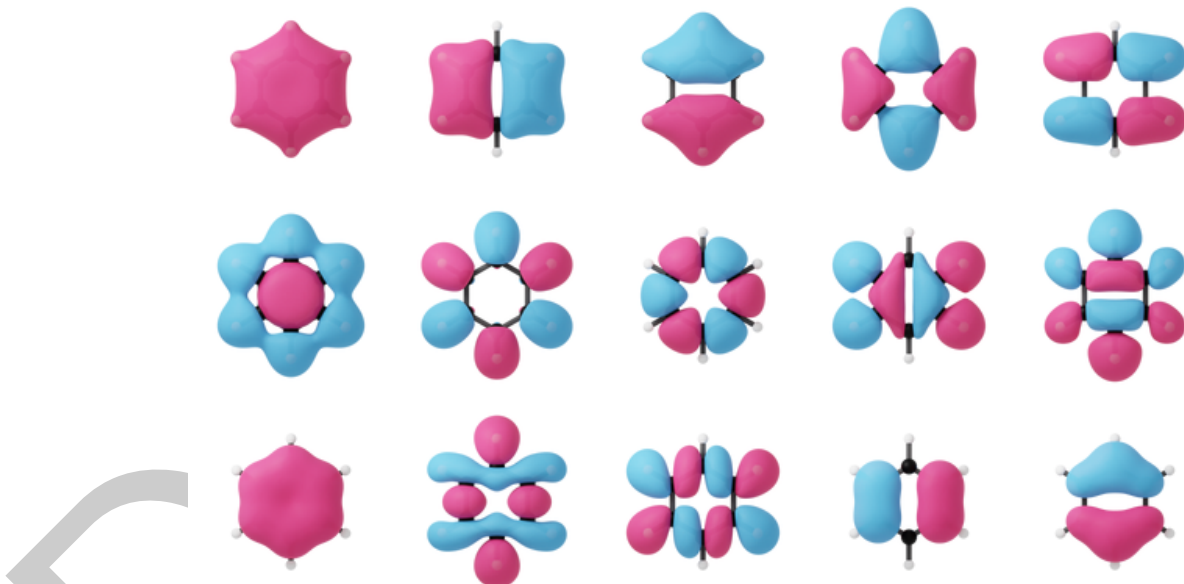


Figure 1: Isosurfaces of the first 15 canonical valence molecular orbitals of benzene.

⁵⁸ For efficient research data management purposes, Den20bj is also able to convert CHGCAR and
⁵⁹ PARCHG files to its own custom d2o format, which is a lossless format that stores the scalar field
⁶⁰ as a collection of floats utilizing compression. Upon conversion of input files to the native d2o
⁶¹ file type, the program explores various compression algorithms, i.e. lzma ([Pavlov, 1996](#)), bzip2
⁶² ([Wielard et al., 1996](#)) and gzip ([Gailly & Adler, 1992](#)), and uses the one that yields optimal
⁶³ results. In comparison to the original CHGCAR or PARCHG files, d2o files are able to achieve a
⁶⁴ compression ratio around 10%.

⁶⁵ Besides building isosurfaces, Den20bj can also produce OpenVDB ([Museth, 2013](#)) files allowing
⁶⁶ for volumetric rendering in programs such as Blender. In contrast to the rendering of isosurfaces,
⁶⁷ the main advantage of volumetric rendering is that internal details and density variations are
⁶⁸ more prominently shown, providing a comprehensive and nuanced understanding of the scalar
⁶⁹ field. In a way, volumetric rendering lies in between isosurfaces and contour plots in terms
⁷⁰ of visualizing a scalar field. An example for the molecular orbitals of benzene is provided in

⁷¹ **Figure 2.** For demonstration and testing purposes, also a scalar field generator functionality is
⁷² included that can create a number of relevant scalar fields to test the algorithms on.

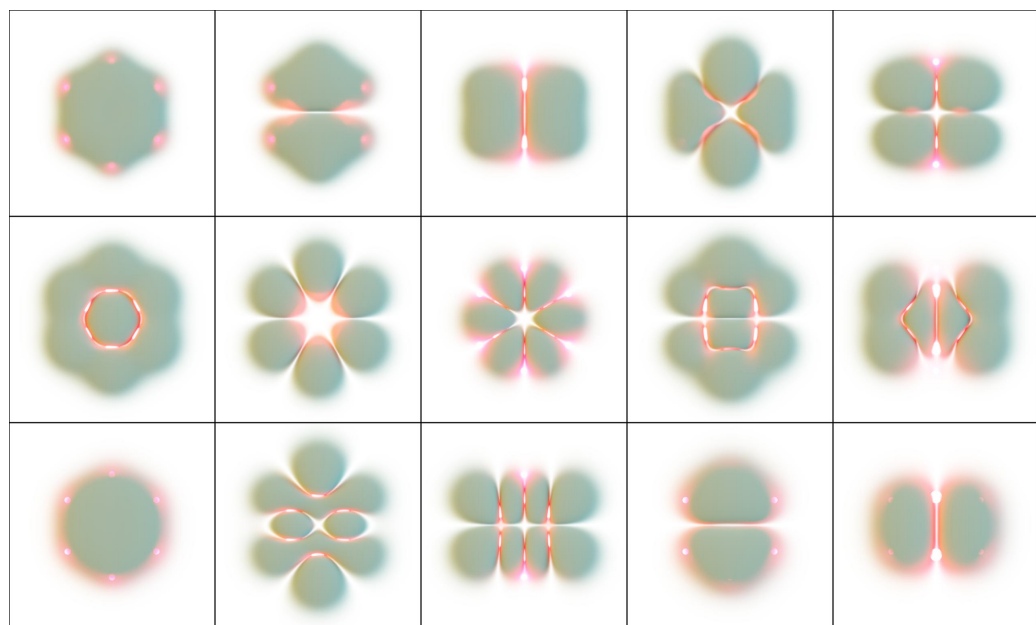


Figure 2: Volumetric rendering of the electron density associated with the first 15 canonical valence molecular orbitals of benzene using the OpenVDB format.

⁷³ Den2Obj requires a relatively small set of dependencies, being Eigen3 ([Guennebaud et al., 2010](#)),
⁷⁴ Boost ([Koranne, 2011](#)), TCLAP ([Smoot et al., 2009](#)), lzma ([Pavlov, 1996](#)), bzip2 ([Wielandt et al., 1996](#)) and gzip ([Gailly & Adler, 1992](#)). Creation of VDB files requires the presence of
⁷⁵ the OpenVDB library ([Museth, 2013](#)). The user can select during compilation whether they
⁷⁶ want to include this functionality or not. Den2Obj is designed to be used by researchers and
⁷⁷ students working in computational materials modelling using the quantum chemical software.
⁷⁸ It has already been used in a number of scientific publications. ([Filot et al., 2016](#); [Su et al., 2016, 2018](#))

⁸¹ An extensive user guide including examples, compilation instructions, tutorials (including a
⁸² rendering tutorial in Blender) and documentation of the command-line arguments, is available
⁸³ at <https://den2obj.imc-tue.nl/>.

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