
Gilles Zérah
**Large scale parallel ab initio electronic structure
calculations with the LOBPCG method.**

CEA-DAM Ile de France
Bruyères le Châtel-91297 Arpajon Cedex
France
`gilles.zerah@cea.fr`
François Bottin
Stéphane Le Roux
Knyazev Andrew

We present an implementation in the *ab initio* plane-wave code *ABINIT*, of a parallelization scheme based on the locally optimal block preconditioned conjugate gradient *LOBPCG* method, and using an optimized three-dimensional (3D) fast Fourier transform (*FFT*).

We will first compare, for various systems, the performance of the method with the more standard eigensolvers currently used in *ABINIT*.

Next, we present the parallelization scheme, which, in addition to the standard data partitioning over processors corresponding to different *k*-points, relies upon data partitioning with respect to blocks of bands and Fourier coefficients.

Finally we analyze the performances of the whole scheme on multiprocessor machines in terms of scalability and convergence speed.