

- Mix
- Normalize
- Multipole moments
- Atomic density matrices
- Construct density
- Calculate density matrix
- Pseudo density
- Density
- Calculate projections
- Send coefs to domains
- Redistribute coefs
- General diagonalize
- Blacs Orbital Layouts
- DistributedAtomicHamiltonian
- Potential matrix
- LCAO eigensolver
- SCF-cycle
- XC Correction
- Atomic
- Hartree integrate/restrict
- Poisson
- XC 3D grid
- vbar
- Hamiltonian
- LCAO WFS Initialize
- Distribute overlap matrix
- TCI: Calculate S, T, P
- Basis functions set positions
- Basic WFS set positions
- TCI: Evaluate splines
- Initialization