```
GQCP::FQMolecularHamiltonian
  # T
  # V
  # g
   + FQMolecularHamiltonian()
   + FQMolecularHamiltonian()
  + kinetic()
   + nuclearAttraction()
   + coulombRepulsion()
GOCP::FOMolecularMagnetic
            Hamiltonian
```

+ FQMolecularMagneticHamiltonian() + FQMolecularMagneticHamiltonian()

+ orbitalZeeman() + diamagnetic() + spinZeeman()