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+ FQMolecularHamiltonian()
+ FQMolecularHamiltonian()
+ FQMolecularHamiltonian()
+ kinetic()
+ nuclearAttraction()
+ coulombRepulsion()
```

```
GQCP::FQMolecularMagnetic
Hamiltonian
```

- + FQMolecularMagneticHamiltonian() + FQMolecularMagneticHamiltonian() + orbitalZeeman()
- + diamagnetic()
 + spinZeeman()