

- EOM-EA/DEA describe target states by <u>electron-attaching</u> operators
- When using closed-shell reference, target states are naturally spin pure
- EOM-EA is good for certain doublet states (e.g., Na atom in the ground and excited states)
- EOM-DEA can describe diradicals (e.g., methylene) and bond-breaking
- State and transition properties can be computed



## EOM-EA: MgOH example

- Alkali-earth atoms like Mg have [core]ns<sup>2</sup> electronic configuration
- They form ionic bonds with halogen-like ligands (e.g. Mg-OH), yielding doublet radicals with an unpaired electron localized on the metal
- EOM-EA can describe its ground and excited states
- Dyson orbitals show where the unpaired electron is
- Import structure to IQmol (<u>mgoh\_structure.xyz</u>)
- Use <u>cationic reference</u> (MgOH+, charge=1, multiplicity=1) and EA\_<u>STATES</u>
- Compute 3 target states

```
20 $rem
1 method = eom-ccsd
2 ea_states = [1, 0, 1, 1]
3 basis = 6-31G*
4 ! Dyson orbitals
5 cc_do_dyson = true
6 print_general_basis = true
7 $end
```

**Note:** Largest abelian subgroup is  $C_{2v}$ : A<sub>1</sub>, A<sub>2</sub>, B<sub>1</sub>, B<sub>2</sub> irreps Use MEM\_TOTAL=4000



- Mg-CC-Mg is a diradical: it has two unpaired electrons
- It is best described by EOM-DEA
- Import structure to IQmol (mgccmg\_structure.xyz)
- Use <u>dicationic reference</u> (MgCCMg<sup>2+</sup>, charge=+2, multiplicity=1) and <u>dea\_states</u>
- Compute 3 target states in each multiplicity
- Request transition properties between the EOM states

```
17 $rem
1 method = eom-ccsd
2 dea_singlets = [1, 0, 0, 0, 0, 0, 0, 1, 1]
3 dea_triplets = [0, 0, 0, 0, 0, 0, 1, 1, 1]
4 basis = 6-31G*
5 ! transition properties
6 cc_trans_prop = 2
7 $end
```

Note: Largest abelian subgroup is  $D_{2h}$ : A<sub>g</sub>, B<sub>1g</sub>, B<sub>2g</sub>, B<sub>3g</sub>, A<sub>u</sub>, B<sub>1u</sub>, B<sub>2u</sub>, B<sub>3u</sub> irreps Use MEM\_TOTAL=4000

## CONCEPTION OF CHEMISTRY

## EOM-EA: MgOH example

463 EOMEA transition 1/A1
1 Total energy = -275.26088905 a.u. Excitation energy = -7.2303 eV.
2 R1^2 = 0.9901 R2^2 = 0.0099 Res^2 = 1.28e-06
3 Conv-d = yes

TransitionEA, eVE\_ex (eV)1/A1-7.2300.0001/B1-3.5373.6931/B2-3.5373.693



IQmol: left panel -> Surfaces





Energy difference relative to reference gives EA=E(EOM) - E(Ref) (electron affinity of MgOH+)

Energy levels in MgOH: difference between the EOM states

## EOM-DEA: Mg-CC-Mg example

State	Trans. sym	m. E_ex (eV)	Multiplicity	f
0	1/A <sub>g</sub>	0.000	singlet	
1	1/B <sub>1u</sub>	0.020	triplet	0.00
2	1/B <sub>2u</sub>	2.873	triplet	0.00
3	1/B <sub>3u</sub>	2.873	triplet	0.00
4	1/B <sub>2u</sub>	2.988	singlet	0.31
5	1/B <sub>3u</sub>	2.988	singlet	0.31

Singlets and triplets are printed separately. Look for a line like:

408 Solving for EOMDEA-CCSD Ag transitions to singlet states Transition properties: 800 State A: eomdea ccsd/rhfref/singlets: 1/Ag 1 State B: eomdea ccsd/rhfref/singlets: 1/B2u

- 2 Energy GAP = 0.1098 a.u. = 2.9885 eV 3 Transition dipole moment (a.u.): 4 A->B: 2.043799 (X 0.000000, Y 2.043799, Z 0.000000)
- 5 B->A: 2.071908 (X 0.000000, Y 2.071908, Z 0.000000)
- 6 Oscillator strength (a.u.): 0.310042

LEAP INTO THE FUTURE OF CHEMISTRY

- 7 Transition angular momentum against gauge origin (a.u.):
- 8 A->B: (X 0.000000i, Y 0.000000i, Z 0.000000i)
- 9 B->A: (X 0.000000i, Y 0.000000i, Z 0.000000i)
- 10 Norm of one-particle transition density matrix:
- 11 A->B: 0.712898; B->A: 0.709384
- 12 ||gamma^AB||\*||gamma^BA||: 0.505718

The states can be visualized by using natural transition orbitals (NTOs)





A pair of NTOs for the 1/Ag to 1/B2g singlet transition with a SV = 0.33