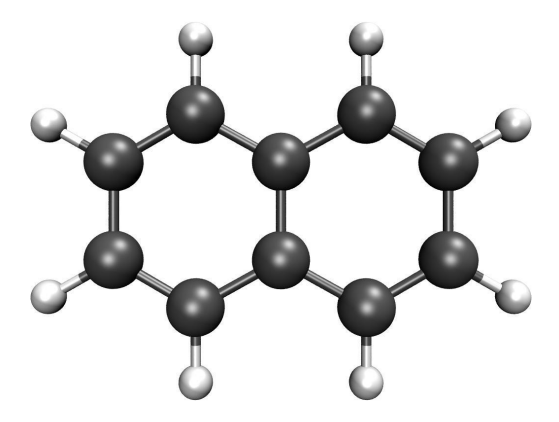
тне **Zysman-Colman** group

Calculation Report

Naphthalene

Excited States (Triplet)



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Summary of Results

Metadata

Username: osl

07/06/2022 Date:

18:50:12

Duration: 4 m, 5 s **True**

Success:

Computational

package:

Turbomole (7.5.0)

Methods: HF, MP2 Basis set: cc-pVDZ

Calculations: Excited States

Orbital spin: restricted **Multiplicity:** 1 (singlet)

SCF Energies

No. of steps: 1

Final energy: -10432.3114 eV

Final energy: -1,006,565 kJmol⁻¹

MP Energies

No. of steps: 1

Final energy: -10467.1582 eV Final energy: -1,009,927 kJmol⁻¹

Geometry

Formula: $C_{10}H_{8}$

Molar mass: 128.1705 gmol⁻¹

Alignment

method:

Minimal

X extension: 6.80 Å Y extension: 5.02 Å Z extension: 0.00 Å **Linearity ratio:** 0.26 Planarity ratio: 1.00

HOMO & LUMO

E_{HOMO,LUMO}: 10.15 eV

E_{HOMO}: -7.78 eV

E_{LUMO}: 2.37 eV

Permanent Dipole **Moment**

Total: 0.00 D X axis angle: 90.00° XY plane angle: 75.96°

Excited States

T₁ energy: 3.27 eV T₁ wavelength: 379 nm

T₁ colour: Ultraviolet

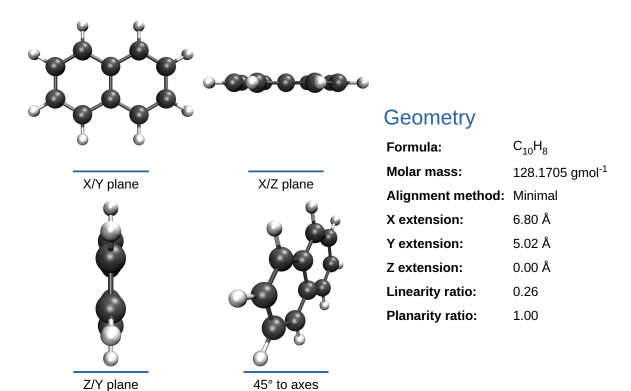
 T_1 CIE (x,y): (0.17, 0.00)

T₁ oscillator strength: 0.00

No. of triplets: 2

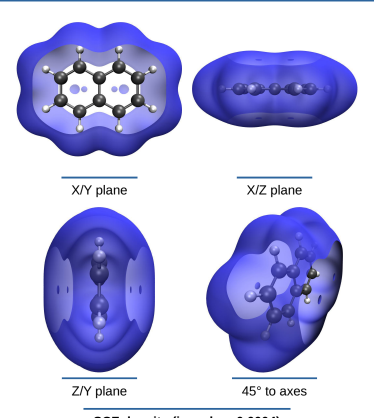
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Geometry



Aligned structure

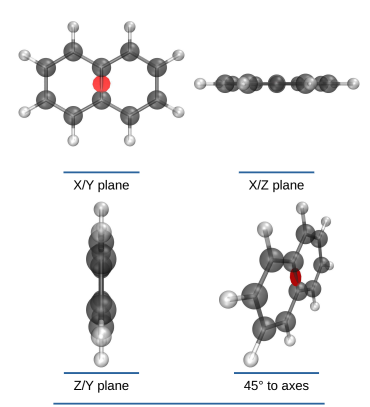
SCF Density



SCF density (isovalue: 0.0004)

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Permanent Dipole Moment



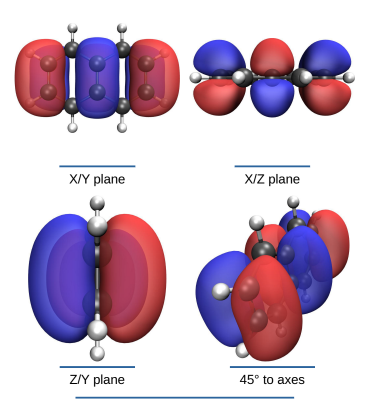
Aligned structure (dipole moment in red)

Dipole Moment

Origin X: 0.00 D Origin Y: 0.00 D Origin Z: 0.00 D **Vector X:** 0.00 D **Vector Y:** 0.00 D **Vector Z:** 0.00 D Total: 0.00 D X axis angle: 90.00° XY plane angle: 75.96 °

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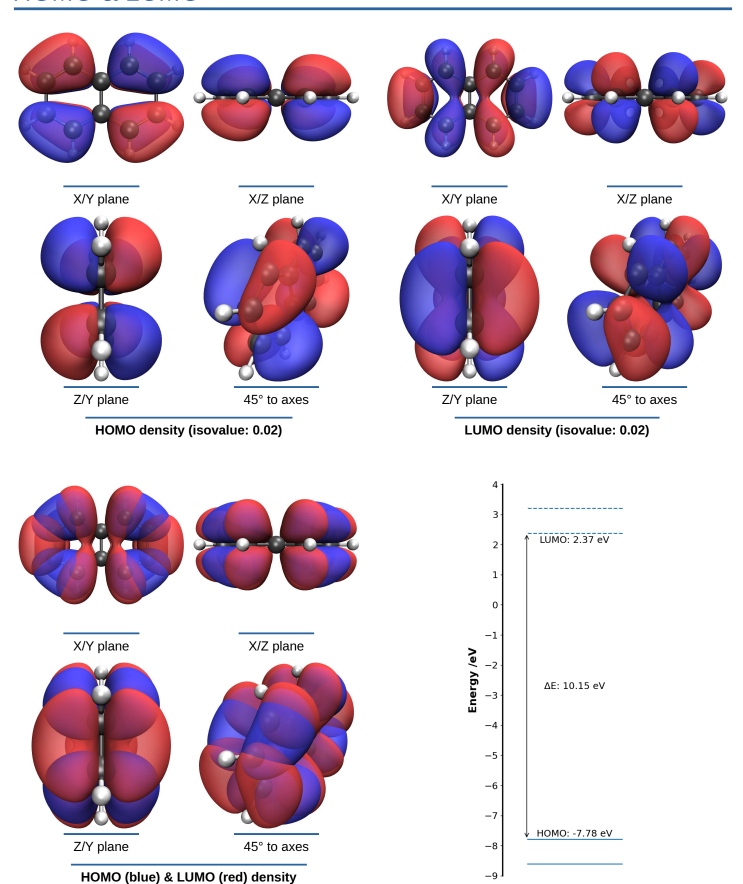
HOMO-1



HOMO-1 density (isovalue: 0.02)

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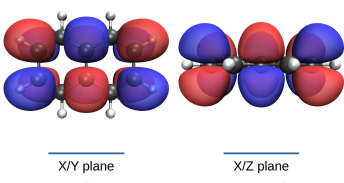
HOMO & LUMO

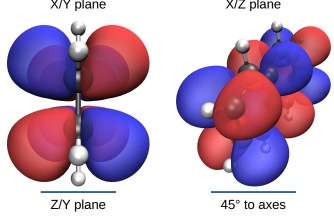


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(isovalue: 0.02)

LUMO+1

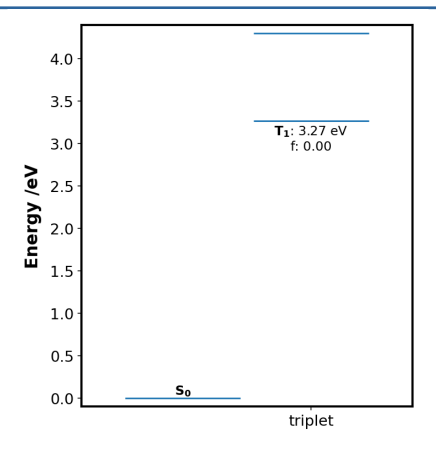




LUMO+1 density (isovalue: 0.02)

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Excited States



Excited States

T₁ energy: 3.27 eV

T₁ wavelength: 379 nm

 T_1 colour: Ultraviolet T_1 CIE (x,y): (0.17, 0.00)

T₁ oscillator strength: 0.00

No. of triplets: 2

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Table of Excited States

Level	Symbol	Symmetry	Energy /eV	Wavelength /nm	Colour, CIE (x,y)	Oscillator Strength	Transitions (probability)
1	T ₁	Triplet-A	3.2689	379.29	Ultraviolet (0.17, 0.00)	0.0000	HOMO → LUMO (0.85) HOMO-2 → LUMO+2 (0.06) HOMO-1 → LUMO+1 (0.05)
2	T ₂	Triplet-A	4.2983	288.45	Ultraviolet (0.00, 0.00)	0.0000	HOMO-1 → LUMO (0.49) HOMO → LUMO+1 (0.46) HOMO-3 → LUMO+2 (0.02)

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Table of Selected Molecular Orbitals

Level	Label	Symmetry	Energy /eV
50	LUMO+15	А	11.9600
49	LUMO+14	Α	11.7353
48	LUMO+13	Α	10.6261
47	LUMO+12	Α	10.4351
46	LUMO+11	Α	7.9455
45	LUMO+10	Α	7.4055
44	LUMO+9	Α	7.3699
43	LUMO+8	Α	6.8297
42	LUMO+7	Α	6.4869
41	LUMO+6	Α	6.3480
40	LUMO+5	Α	5.4144
39	LUMO+4	Α	5.4053
38	LUMO+3	Α	4.9896
37	LUMO+2	Α	4.7431
36	LUMO+1	Α	3.2023
35	LUMO	Α	2.3705
34	НОМО	Α	-7.7835
33	HOMO-1	Α	-8.6036
32	HOMO-2	Α	-10.3698
31	HOMO-3	Α	-12.0540
30	HOMO-4	Α	-12.9253
29	HOMO-5	Α	-13.1917
28	HOMO-6	Α	-14.1706
27	HOMO-7	Α	-14.3301
26	HOMO-8	Α	-15.2492
25	HOMO-9	Α	-15.7422
24	HOMO-10	Α	-15.7464
23	HOMO-11	Α	-16.4964
22	HOMO-12	Α	-16.8787
21	HOMO-13	A	-18.2419
20	HOMO-14	A	-18.8268
19	HOMO-15	Α	-19.1551

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Table of Atoms

Element	X Coord	Y Coord	Z Coord
С	-1.2509100	-1.4118100	-0.0000100
С	-2.4487500	-0.7132700	0.0000100
С	-2.4487500	0.7132700	0.0000100
С	-1.2509100	1.4118100	-0.0000100
С	-0.0000000	0.7179300	-0.0000200
С	-0.0000000	-0.7179400	-0.0000200
С	1.2509100	-1.4118100	-0.0000100
С	1.2509100	1.4118100	-0.0000100
С	2.4487500	0.7132700	0.0000100
С	2.4487500	-0.7132700	0.0000100
Н	-1.2480900	-2.5080700	-0.0000100
Н	-3.4000100	-1.2561900	0.0000200
Н	-3.4000100	1.2561900	0.0000300
Н	-1.2480900	2.5080700	-0.0000100
Н	1.2480900	-2.5080700	-0.0000100
Н	1.2480900	2.5080700	-0.0000200
Н	3.4000100	1.2561900	0.0000300
Н	3.4000100	-1.2561900	0.0000200

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Silico Calculation Report

Part of the silico software package

Version 1.0.0-pre.30 7 June 2022

Silico makes use of a number of 3rd party libraries and programs; please cite these appropriately in your works:

Extraction and processing of results: **cclib**^[1] Rendering of 3D images: **VMD**^[2], **Tachyon**^[3]

Rendering of graphs: Matplotlib^[4]

Calculation of CIE colour coordinates: Colour Science^[5]

Generation of reports: Mako^[6], Weasyprint^[7]

Scientific constants: SciPy^[8]

Conversion of file formats: Pybel^[9], Openbabel^[10]

Calculation of spin-orbit coupling: PySOC^[11]

Rendering of 2D structures: **RDKit**^[12]

Saving of state during submission: $Dill^{[13,14]}$

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