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# Lawsonone isomers, lawsonone ether and bilawsonone for dye-sensitized solar cells applications: DFT and UV–Vis studies

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## Abstract

Structural and optoelectronic properties of lawsonone (L), lawsonone ether (LE) and bilawsonone (BL) were studied theoretically using the DFT and time-dependent DFT methods with hybrid functional B3LYP5 and 6-311G (2d,p) basis set. For the monomer lawsonone molecule, isomerization reaction between two rotational isomers was analyzed based on a thermodynamic approach. The electronic spectra of the dyes molecules in a vacuum and solvents (DMSO and CH<sub>2</sub>Cl<sub>2</sub>) were computed. The maximum wavelengths were found at 355–408 nm for the LE and 350–448 nm for BL that indicated bands shift to visible range compared to L (340 nm). The UV–Vis spectra of the L and BL were measured experimentally in DMSO solution. For the BL, a broad and intensive band was observed in a visible region at 452 nm that apparently would favour sensitizing ability of the dye. The optoelectronic properties of the LE and BL showed them as more promising candidates for DSSCs applications compared to the individual lawsonone dye.

## Keywords

Lawsonone isomers, Lawsonone ether, Bilawsonone, Dye-sensitized solar cells, UV–Vis/vibrational spectra, DFT/TD-DFT