


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# Spectroscopic and DFT study of (1R\*,2R\*,4S\*)-5-(5,5-dimethyl-1,3-dioxan-2-yl)-8,8-dimethoxy-7-oxobicyclo[2.2.2]oct-5-en-2-yl cyanide

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# Spectroscopic and DFT study of (1R\*,2R\*,4S\*)-5-(5,5-Dimethyl-1,3-dioxan-2-yl)-8,8-dimethoxy-7-oxobicyclo[2.2.2]oct-5-en-2-yl cyanide

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**Abstract.** Proton (<sup>1</sup>H) and Carbon (<sup>13</sup>C) Nuclear magnetic resonance and Ultraviolet-Visible spectrum was simulated for (1R\*,2R\*,4S\*)-5-(5,5-Dimethyl-1,3-dioxan-2-yl)-8, 8-dimethoxy -7-oxobicyclo [2.2.2] oct-5en-2-yl Cyanide (DDO). By using Gauge Independent Atomic Orbital method, chemical shifts were generated and compared with its corresponding experimental values DDO. To understand the origin of chemical reactivity and Ultraviolet-Visible spectrum, FMO parameters measured. Non-linear optical parameters were calculated.

## INTRODUCTION

(1R\*, 2R\*, 4S\*)-5-(5,5-Dimethyl-1,3-dioxan-2-yl)-8, 8-dimethoxy -7-oxobicyclo [2.2.2]oct-5en-2-yl Cyanide (DDO) and synthesis part and experimental data reported by Santhosh et al.<sup>1</sup> This compound comes under the category of (MOBs) masked o-benzo-quinones.<sup>2</sup> A Cycloaddition reaction plays a vital role in preparation of antiglaucoma compounds<sup>3</sup> and many natural products.<sup>4,5</sup> We recently reported the results of such biologically active molecules.<sup>6-16</sup> Hence, we undertook this work with the following goals.

- calculate NMR shifts (<sup>1</sup>H and <sup>13</sup>C) and examine their relationship with measured<sup>1</sup> values,
- simulated UV-Visible spectrum and,
- NLO, MESP and FMO parameters to make the investigation comprehensive.

## COMPUTATIONAL ASPECTS

By means of DFT/6-311++G(d,p) rank of conjecture and executed with Gaussian Window 09 program<sup>17,18</sup> simulated NMR<sup>18</sup> and UV-Vis spectrum.<sup>19-21</sup> We determined Chemical reactivity<sup>22-25</sup> of selected molecule DDO.

## RESULTS

### Most stable conformer

Chosen sample is optimized with the above mentioned method. The calculated geometrical parameters compared with observed values<sup>1</sup> and are shown in Table.1. Its optimized energy:  $-2869.026 \times 10^{-3} \text{ k Jmol}^{-1}$ . DDO comes under C<sub>1</sub> symmetry point group structure and is depicted in figure 1.

## NMR SIGNALS

To verify the correlation between the calculated and experimentally observed NMR shifts, we drawn the graphs of observed chemical shifts verses computed chemical shifts for DDO. These are straight lines as shown

in figure 2 for  $^{13}\text{C}$  and  $^1\text{H}$  NMR spectra. Coefficient of association  $r^2$  is extremely close up agreement to accord for  $^1\text{H}$  and  $^{13}\text{C}$  NMR spectra of chosen molecule. It is clear that the theoretical and experimental chemical signals are well agreed, and can be evidenced from figure 2.

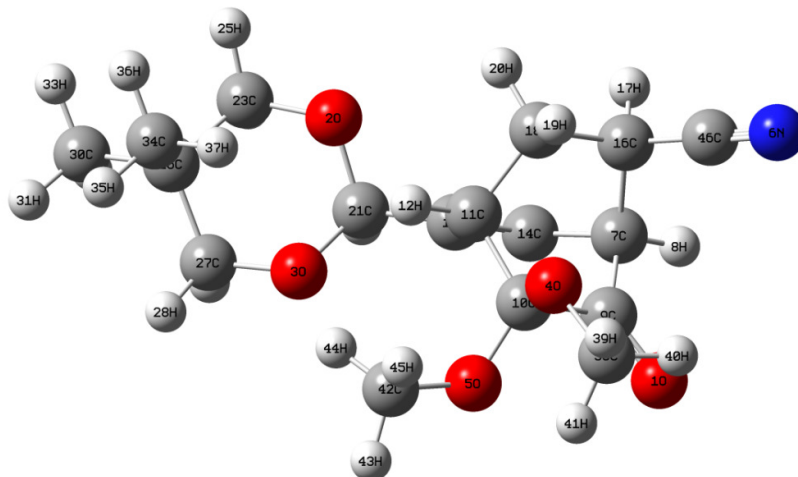


FIGURE 1: Optimized geometrical structure of DDO ( $E_{\text{DDO}} = -2869.026 \times 10^{-3} \text{ k J mol}^{-1}$ )

TABLE 1: Geometrical parameters of DDO

Geometric parameter	Calculated Value(DFT)	Expt. Value <sup>a</sup>
<b>Bond lengths ( in Å )</b>		
O1-C9	1.202	1.206
O2-C21	1.418	1.412
O2-C23	1.431	1.432
O3-C21	1.412	1.406
O4-C10	1.413	1.409
O5-C10	1.399	1.399
O5-C42	1.432	1.428
N6-C46	1.153	1.134
C7-H8	1.080	0.980
C7-C14	1.511	1.508
<b>Bond angle ( in ° )</b>		
O1-C9-C7	124.081	124.105
O1-C9-C10	123.807	123.371
O2-C21-O3	110.784	111.648
O2-C21-C13	108.003	107.569
O2-C21-H22	105.766	109.993
O2-C23-H24	109.085	109.345
O2-C23-H25	106.107	109.344
O3-C27-C26	111.728	111.182
O4-C10-O5	112.628	112.732
O5-C10-C9	105.659	104.997
<b>Dihedral angle ( in ° )</b>		
O1-C9-C7-C14	121.25	119.66
O2-C21-O3-C27	61.20	58.77
O3-C21-O2-C23	60.89	58.50
O4-C10-O5-C42	59.56	59.53
O5-C10-O4-C38	56.57	52.90
O5-C10-C9-C7	127.06	129.65
C7-C9-C10-C11	4.46	6.745
C9-C7-C14-C13	58.40	57.00
C10-C11-C18-H20	178.58	178.00
C10-C9-C7-C14	58.45	59.91

a: From reference [1]

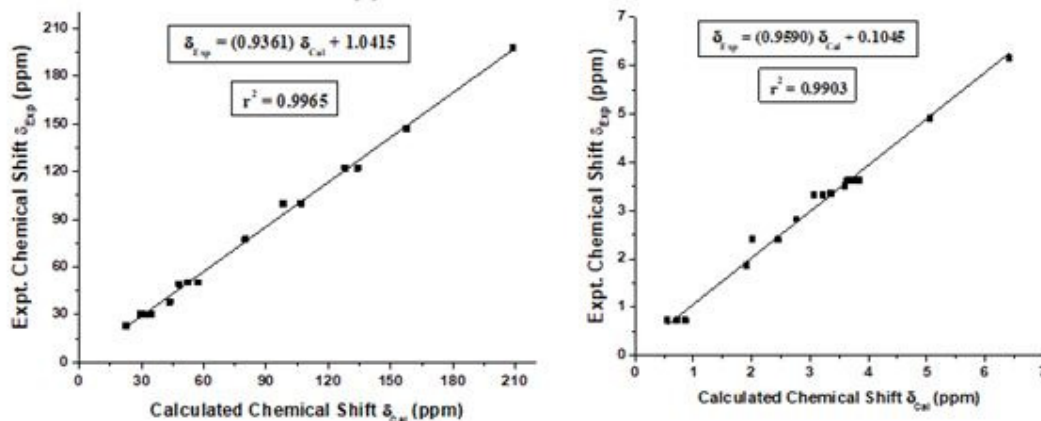


FIGURE 2: Linear regression curve for carbon and proton signals

## UV-VISIBLE PEAKS

Computed UV-Vis absorption peaks obtained in the computations are due to the electronic transitions. HOMO and LUMO determine the reactivity of the selected compound.<sup>26</sup> Electron donor is HOMO and acceptor is LUMO.<sup>27,28</sup> The calculated peaks at  $\lambda_{\max} = 318.35$  nm, its oscillator strength,  $f = 0.0060$  and another one observed at  $\lambda_{\max} = 240.84$  nm with  $f = 0.0416$  and are shown in figure 4. The origin of the signals is mainly due to the transitions of  $H \rightarrow L$  and  $H-1 \rightarrow L$  for DDO.

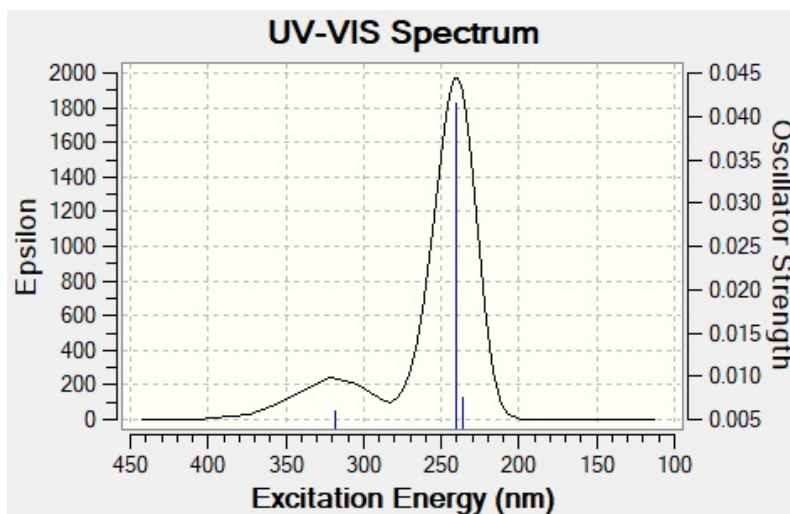


FIGURE 3: UV-Vis peaks

## CHEMICAL REACTIVITY OF THE MOLECULE DDO

Frontier molecular orbital energy gap plays vital role in understanding the chemical reactivity such as reactants kinetic characteristics and chemical reactions of the molecule. The calculated energy gap between H and L:

4.040 eV (figure 5) and the chemical potential ( $\mu = -7.238$  eV) is negative for DDO, and is chemically

stable.<sup>29,30</sup> These parameters describe aspects like drug design and toxicological behavior of eco system.

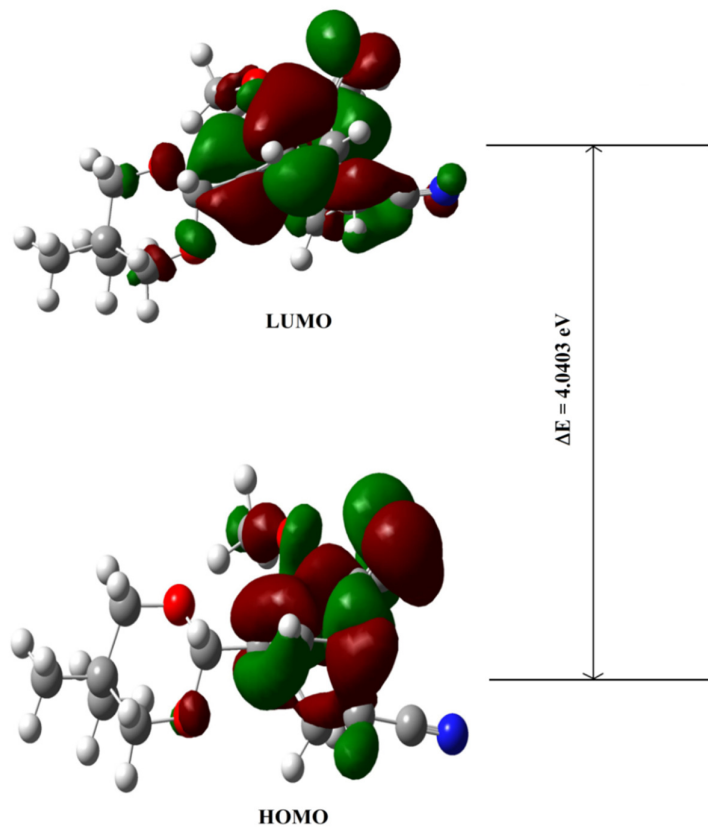


FIGURE 4: HOMO and LUMO plots of DDO

### MOLECULAR ELECTROSTATIC SURFACE POTENTIAL OF DDO

The total electron density plot of DDO (see figure.6) shows the difference between charge distributions among various parts of a given molecule<sup>31</sup>. In figure 5, relatively negative regions are shown in red and relatively positive region is shown in green. The negative region, indicated in red is primarily over the N and O atomic positions, caused by the donation of oxygen and atoms of nitrogen (lone-pair electrons), whereas the positive section designated in green is over the remaining atoms.

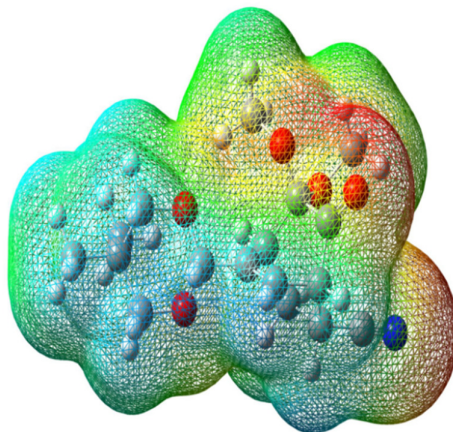


FIGURE 5: Total electron density plots of DDO

## NON-LINEAR OPTICAL (NLO) BEHAVIOUR

NLO behaviour of chosen sample confirmed by comparing the Urea values of  $\mu_t$  and  $\beta_t$  (Urea  $\mu_t$  :1.3732 Debye and  $\beta_t$  : 372.8 $\times 10^{-33}$  cm<sup>5</sup>/e.s.u). Values of DDO are  $\mu_t$  : 3.279 Debye and  $\beta_t$  : 391.279 $\times 10^{-33}$  cm<sup>5</sup>/e.s.u. Hence it can be concluded that DDO is a good NLO materials.<sup>32-36</sup>

## CONCLUSION

The following inferences are drawn from the computations:

- DDO molecule has the non-planar structure acquiring point group of C<sub>1</sub> symmetry. Theoretical computed geometric parameters of DDO are good in agreement with the values of XRD.
- Good correlation between the theoretical and experimental NMR signals.
- Theoretical UV-Vis peaks identified.
- Electron density plot was drawn and thermal energies were also estimated for DDO.
- DDO is an excellent contender for enlargement of novel NLO materials.

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