UV Spectra of Glycolic Acid Derivatives – A Time-Dependant Density Functional Theory Examination

Mahendiraprabu Ganesan, Selvarengan Paranthaman

Abstract: One of the serious health issues faced by people throughout the world in recent years is Skin cancer, which is caused by harmful UV radiations from the sun. To protect the human skin from direct exposure to these ultraviolet radiations, preventive measures have to be taken and once such solution is the application of suitable and safer sunscreen. It is the need of the hour to develop better and harmless sunscreens to protect the skin. One such material which plays a major role in dermatology and cosmetics industry is Glycolic acid, CH2OHCOOH, a precursor of α-hydroxy carboxylic acids. Our present study is to find out if the glycolic acid conformers and its derivatives are capable of absorbing harmful radiations in the UVB and UVA ranges. To understand this property the UV absorption spectra is calculated using Time dependant density functional theory (TD-DFT). Further, our TDDFT calculations have shown that α-Nitromandelic acid, a glycolic acid derivative shows absorption in both UVB and UVA regions. This indicates that α-Nitromandelic acid can be probably used as a UV filter in sunscreen lotions.

Keywords: UV filter, TD-DFT, Nitromandelic acid, Glycolic acid derivatives.

I. INTRODUCTION

UV radiation like UVA, UVB, and UVC radiation [1] are emitted by the sun and are harmful to human skin. We are regularly exposed to the UVB (~290–320 nm) and UVA (320–400 nm) radiations [2,3]. In general, the stratospheric region of the ozone layer filters these harmful UVC radiation [4]. But the recent reports of depletion in the ozone layer allows the harmful UV radiation reach the earth’s surface. This in turn causes skin damage like erythema (sunburn), cutaneous photaging, immune suppression, and an increased risk of skin cancer.

Need for organic sunscreens with active UV filters have increased which should not harm the skin as well as protect our skin from damages caused by these harmful radiations. Organic UV filters is one of the most common active constituents in sunscreen products used for attenuating skin photodamage [1,4,5]. When the UV radiation falls on the organic filters it excites and electron from its ground state to its excited state and this occurs because of the presence of saturated groups (n orbitals) and atoms with unpaired electrons (n orbitals). Some saturated groups that bond to this system also contribute to UV absorption. Similarly, One such material which plays a major role in dermatology and cosmetics industry is Glycolic acid, CH2OHCOOH, a precursor of α-hydroxy carboxylic acids. It also used in the sunscreen products [6-9]. However, very few studies were reported on the related systems both theoretically [10-17] as well as experimentally [11,12,18,19]. As far as we have seen from the literatures collected so far, there is no UV spectral study reported in the literature in the glycolic acid conformers and their derivatives. Therefore the purpose of this study is to find out if the glycolic acid conformers and its derivatives have the potential to absorb in the UVB and UVA ranges. Since glycolic acid is used in skin care products, it is necessary to study its UV absorption ranges.

II. COMPUTATIONAL DETAILS

The vibrational frequency calculations are performed to identify the minimum energy structure. The time dependent density functional theory (TDDFT) [20,21] calculations with DFT functional, Becke’s three parameters exact exchange functional (B3),[22] combined with the Lee–Yang–Parr gradient corrected functional (LYP)[23] with 6–311++G(d,p) basis set is used for the calculations at B3LYP/6-311++G(d,p) level of theory and is performed to understand the UV absorption ranges of the glycolic acid conformers and its derivatives. All the calculations are performed using gaussian09W program.[24]

III. RESULT AND DISCUSSION

Since Glycolic acid is an important component of dermatological and cosmetic products, it is necessary to study its UV absorption ranges. Hence, in the present study TDDFT calculations at B3LYP/6-311++G(d,p) level of theory are performed on glycolic acid conformers. Our calculations have shown that the GAC conformer of glycolic acid has maximum wavelength (221.39 nm). From our calculated results, it is understood that the glycolic acid conformers do not show absorption in the UVB/UVA ranges. Hence, we have extended the TDDFT studies on the derivatives of glycolic acid.
The glycolic acid derivatives chosen in this study are Mandelic acid, α-CyclohexyImandelic acid, p-Bromomandelic acid, Benzilic acid, p-Chloromandelic acid, p-Nitromandelic acid, m-Nitromandelic acid and o-Nitromandelic acid. Totally eight derivatives are taken for this study, to understand their ability to absorb in UVB/UVA ranges. Out of the eight derivatives of glycolic acid, three derivatives namely o-Nitro (alpha-Hydroxy-2-nitrobenzeneacetic acid), p-Nitro (alpha-Hydroxy-4-nitrobenzeneacetic acid) and m-Nitro (alpha-Hydroxy-3-nitrobenzeneacetic acid) mandelic acids show absorption in the UVB and UVA regions and these optimized structures are shown in Fig. 1. The TDDFT calculations have also been performed for the most popular UV filters such as Benzophenone-3 (BZP-3) and Urocanic acid for the comparison purpose.

![Fig. 1. The optimized structures (a) o-Nitro-, (b) p-Nitro-, (c) m-Nitromandelic acid of glycolic acid derivatives, (d) BZP-3 and (e) Urocanic acid using B3LYP method.](Image)

Further, our TDDFT studies on BZP-3 have shown that the absorption spectra for UVB and UVA regions are present in the same HOMO → LUMO transition with different wavelengths. From Table 1, it can be noted that the UVA wavelength

**Table-1: Calculated Energy (E, eV), wavelength (λ, nm), oscillator strength (f) and Molecular orbitals HOMO (H) and LUMO (L) for the glycolic acid derivatives using TD-DFT/B3LYP/6-311++G(d,p) level of theory**

<table>
<thead>
<tr>
<th>Derivatives</th>
<th>E</th>
<th>λ</th>
<th>f</th>
<th>Molecular orbitals</th>
</tr>
</thead>
<tbody>
<tr>
<td>o-Nitromandelic acid</td>
<td>3.750</td>
<td>331</td>
<td>0.017</td>
<td>H → L: 0.479</td>
</tr>
<tr>
<td></td>
<td>4.056</td>
<td>306</td>
<td>0.028</td>
<td>H → L: 0.647</td>
</tr>
<tr>
<td></td>
<td>4.264</td>
<td>291</td>
<td>0.005</td>
<td>H → L: 0.472</td>
</tr>
<tr>
<td>m-Nitromandelic acid</td>
<td>3.775</td>
<td>328</td>
<td>0.000</td>
<td>H → L: 0.683</td>
</tr>
<tr>
<td></td>
<td>4.176</td>
<td>297</td>
<td>0.017</td>
<td>H → L: 0.670</td>
</tr>
<tr>
<td></td>
<td>4.293</td>
<td>289</td>
<td>0.001</td>
<td>H → L: 0.615</td>
</tr>
<tr>
<td>p-Nitromandelic acid</td>
<td>3.242</td>
<td>382</td>
<td>0.000</td>
<td>H → L: 0.693</td>
</tr>
<tr>
<td></td>
<td>4.009</td>
<td>309</td>
<td>0.000</td>
<td>H → L: 0.690</td>
</tr>
<tr>
<td></td>
<td>4.058</td>
<td>306</td>
<td>0.016</td>
<td>H → L: 0.507</td>
</tr>
<tr>
<td></td>
<td>4.325</td>
<td>287</td>
<td>0.244</td>
<td>H → L: 0.507</td>
</tr>
<tr>
<td>Benzophenone-3</td>
<td>3.653</td>
<td>339</td>
<td>0.014</td>
<td>H → L: 0.415</td>
</tr>
<tr>
<td></td>
<td>4.078</td>
<td>304</td>
<td>0.124</td>
<td>H → L: 0.519</td>
</tr>
</tbody>
</table>

Further, our TDDFT studies on BZP-3 have shown that the absorption spectra for UVB and UVA regions are present in the same HOMO → LUMO transition with different wavelengths. From Table 1, it can be noted that the UVA wavelength

![Fig. 2. Comparison of UV spectrum of glycolic acid derivatives with some organic UV filter compounds calculated using TD-DFT/B3LYP/6-311++G(d,p) method.](Image)
is 339 nm (f=0.014) and UVB is 304 nm (f=0.124) for BZP-3. The calculated BZP-3 data agrees very well with the experimental data [13]. The UV absorption studies are also performed on cis-Urocanic acid using B3LYP/6-311++G(d,p) level of theory. Our calculations show that the Urocanic acid show absorption in UVB region. In this case, the lowest lying energy transition is HOMO → LUMO with the wavelength 305 nm (f=0.48). In both BZP-3 and cis-Urocanic acid, HOMO → LUMO transition is predominant. From the above discussion, it can be concluded that, o-Nitromandelic acid derivative is one of the best UV filter lying in both UVB and UVA region and is clearly shown in Fig. 3.

Fig. 3. The molecular orbital diagrams of (a) o-Nitromandelic acid, (b) p-Nitromandelic acid, (c) m-Nitromandelic acid, (d) Benzopene-3 and (e) Urocanic acid glycolic acid derivatives

The reason is, the molecular orbital transitions for p-Nitro and m-Nitromandelic acids lies between α-Hydroxyl group and nitrobenzeneacetic acid group, but in the case of o-Nitromandelic acid, this transition is shifted from nitrobenzene acetic acid group to α-Hydroxyl group.

IV. CONCLUSION

In the present investigation the UV absorption study for the glycolic acid derivatives is calculated using TD-DFT study. From this study, it can be concluded that the o-Nitromandelic acid could be used as UV filter. In addition to this, our theoretical studies, demonstrates the ability of using glycolic acid derivatives as UV filter.

APPENDIX

It is optional. Appendixes, if needed, appear before the acknowledgment.

ACKNOWLEDGMENT

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REFERENCES
