Study of nonlinear optical responses of phytochemicals of *Clitoria ternatea* by quantum mechanical approach and investigation of their anti-Alzheimer activity with in silico approach

Shradha Lakhera1 · Kamal Devlal1 · Meenakshi Rana1 · Ismail Celik2

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Abstract

*Clitoria ternatea* is a flowering plant with promising medicinal plants with a wide variety of active phytochemicals. The present study aimed at the computational investigation of the nonlinear optical (NLO) responses of the active phytochemicals of the *Clitoria ternatea*. The computational investigation of the NLO features was done by using the density functional theory (DFT) by B3LYP/6-311G+(d,p) basis set. The structural parameters, mulliken charge distribution, and molecular electrostatic potential (MEP) surface clearly show the intramolecular charge transfer within Clitorin. The NLO properties were identified by computing the polarizability parameters. As the plant has high medicinal characteristics, the inhibiting properties of its phytochemicals were also investigated to combat Alzheimer disease (AD). The systematic in silico study identifies Clitorin as the most active and inhibiting phytochemicals of the plant. The results obtained from molecular dynamics (MD) simulation tell the stability of the complex and make it a fair selection as a drug-like molecule against AD. The cardio-toxicity analysis done for the Clitorin molecule verifies that it is harmless for the heart.

Keywords *Clitoria ternatea* · Clitorin · Dipole moment · Polarizability · Molecular docking · MD simulation

Introduction

The research-based on naturally existing compounds has occupied a vast part in the development of science and technology [1]. Pharmaceutics, medicinal sciences, cosmetics, infrastructure development, health supplements, luminescent materials, cleansing agents, synthetic fibers, dyes, coatings, lasers, microfabrication, etc., are the major fields of research where naturally existing compounds have made a significant volume [2]. The reason behind their wider applications than the human-made compounds is their high reactivity, non-hazardous, and better efficiency. Due to having high reactivity, the natural compounds have many interdisciplinary applications also. Even in multimedia science, data storage, sensing devices, and nonlinear optical (NLO) devices, these compounds have grown in their applications [3]. Therefore, the development of new organic NLO materials has given birth to a new era of material science and engineering. The literature survey had shown that faster response time, higher thermal stability, better optical frequency transform, and stronger intramolecular charge transfer are observed in the organic NLO materials like l-arginine maleate dihydrate, l-methionine l-methioninium...
hydrogen maleate, and urea [4, 5]. Thus, the development of organic NLO supplements is a more convenient and followed area of research among the researchers. Numerous studies depending on the development of NLO materials from plant phytocompounds have been recorded so far. The investigation of the NLO behavior of the esterified derivative of Brassicasterol by computational approaches [6] had been reported. Some of the studies on the development of NLO candidates from plant derivatives are from Alpinia calcarata silver nanoparticles [7], Dioscorea alata [8], Mirabilis Jalapa [9], Coriandrum sativum extracts [10], chlorophyll-a extracted from Andrographis paniculata leaves [11], curcumin derivatives [12], etc.

Clitoria ternatea, a perennial flowering plant is one of the most used medicinal plants (Fig. 1) like antioxidant, hypolipidemic, anticancer, anti-inflammatory, analgesic, antipyretic, antidiabetic, antimicrobial, and gastro-intestinal antiparasitic [13, 14]. It is known by different names as Asian pigeonwings, bluebell vine, blue pea, butterfly pea, cordofan pea, Darwin pea, and many more. The leaves, roots, flowers, seeds, and even roots of this plant are beneficial. The numerous biological activities of its phytochemicals make it significant for research. The high reactivity of the phytochemicals can be interpreted as high polarizabilities. Thus, its phytochemicals can also be considered for the development of organic NLO materials. Its major phytochemicals include Cinnamic acid, Delphinidin-3,5-diglucoside, Flavanol 3-O-D-glucoside, Genistein, Kaempferol, Kaempferol-3,7-diglucoside, Clitorin, Linolenic acid, Oleic acid, Palmitic acid, Taraxerol, and Taxaxeron [15, 16]. These phytochemicals are seen to be mentioned in most of the studies done with this plant.

Since Clitoria ternatea has many active phytochemicals and medicinal property, therefore, in the present study, we have investigated the NLO properties of the phytochemicals of the Clitoria ternatea plant. The computational study was performed with the phytochemicals of this plant using density functional theory (DFT). The optimization of the structures of the phytochemicals was done reveals the quantum chemical properties of the phytochemicals. The initial screening of the reactive molecule was done by dipole moment and the molecule with the highest dipole moment was selected for the further inquiry into NLO behavior. The detection of NLO activity of the selected molecule was further done by performing reactivity analysis, spectral analysis, and calculating polarizability parameters.

Promoting the interdisciplinary side of this study, a systematic in silico study was also performed with the active phytochemicals of Clitoria ternatea. The in-silico modeling is generally used computer-aided drug designing (CADD) technique that is preferred by the chemists and researchers for the evaluation of the drug-like character of the selected systems. Thousands of research works had been reported based on CADD like Feverfew and Piper longum which were used against the Mpro and PLpro of COVID-19 [17, 18], Tridax procumbens was used against MCM7 breast cancer [19], Blumea mollis was used as an anti-fungal candidate [20], Bjerkandera adusta was used against Proteostasis Network Modules [21], and Curcuma

Fig. 1 Morphology of Clitoria ternatea plant

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longa (Turmeric) and Cymbopogon citratus (lemon grass) as lipoxygenase inhibitor [22]. Thus, motivated by this fact, in silico modeling of selected most active phytochemicals of the Clitoria ternatea plant had also been performed in this study. Many studies were seen reporting the pharmacological uses of Clitoria Ternatea derivatives in treating neurological disorders. A review of the neuropharmacological potentiality of Clitorin is explained in detail in the cited paper [23, 24]. It is also used as an anti-oxidant [25]. It is also used as a base supplement for eye drops [26]. Apart from this, this plant is also used for the treatment of various brain diseases like dementia, depression, and brain cancer (cell cycle checkpoint proteins in the cyclin/CDK pathway in cancer cells) [27, 28]. Thus, it can be said that the derivatives of the title plant can be used for medicating the brain or neuro-related sickness. Keeping this in mind, the phytochemicals of the title plant are used for inhibiting the BACE1 macromolecule of Alzheimer’s disease (AD). AD is a neurodegenerative disease that is mostly found in people over 65 years of age. Molecular docking was performed to check the extent of the binding of ligands at the reactive binding sites of the target macromolecule. Cardiotoxicity was also determined for the ligand having the best binding score. Molecular dynamics (MD) simulation was performed targeting a novel clinical candidate BACE1 receptor for the treatment of AD.

**Computational procedure and calculation**

**Chemical reactivity and NLO activity**

The PDB structures of fourteen phytochemicals of the Clitoria ternatea were downloaded from the online database “IMPPAT (Indian Medicinal Plants, Phytochemistry, And Therapeutics)” (https://cb.imsc.res.in/imppat/home) and their structures with chemical formulae are mentioned in SD 1. The software “Open Babel” (http://openbabel.org/wiki/Main_Page) was used to convert the PBD files into gif files. To investigate the NLO responses of the phytochemicals, the structures of the phytochemicals were optimized using the software Gaussian 09 [29] (https://gaussian.com/glossary/g09/). The results were analyzed using the GUI “GaussView 5.0” [30] (https://gaussian.com/gaussview6/). All the computational calculations were done for the ground state of the structures using DFT with Becke-3-Lee–Yang–Parr (B3) exchange function combined with (LYP) correlation and 6-311G basis set [31]. The optimized geometries were used for the further detection of stability and chemical reactivity checks. The energy corresponding to the frontier molecular orbitals (FMO) was also derived from the optimized geometries which were used for the calculations of global reactivity parameters. These parameters are calculated with the help of Koopman’s equations [32] given below:

\[
\Delta E = E_{LUMO} - E_{HOMO} 
\]

\[
IP = E_{HOMO}
\]

\[
EA = -E_{LUMO} 
\]

\[
CP = \frac{E_{HOMO} + E_{LUMO}}{2} 
\]

\[
\chi = \frac{(IP + EA)}{2} 
\]

\[
\eta = \frac{E_{LUMO} - E_{HOMO}}{2}, \quad S = \frac{1}{\eta} 
\]

The electronic spectra were computed using TD-DFT (time-dependent DFT) method. The vibrational spectra were also computed using the same basis set. In Raman spectra analysis, Raman intensity is also calculated corresponding to high frequency modes. It was calculated by below given expression:

\[
I = \frac{f(v_o - v_i)^4 Si}{vi^4[1 - \exp \left(\frac{hc}{kT}\right)]} 
\]

where \(I\) refers to Raman intensity of the considered mode, \(f\) is a constant with value \(10^{-12}\), \(v_o\) has value 9398.5 cm\(^{-1}\), and \(v_i\) and \(S_i\) are the vibrational wavenumber and Raman activity of selected mode respectively. \(h, c, k, T\) have their usual meanings. The UV–Vis and the Raman spectra was plotted using software “Origin 2021” (https://www.originlab.com/).

For NLO analysis, the polarizability parameters total dipole moment (\(\mu_{total}\)), total isotropic polarizability (\(\alpha_{total}\)), anisotropy of polarizability (\(\Delta \alpha\)), and first-order hyperpolarizability (\(\beta_{total}\)) were calculated by the following given formulae:

\[
\mu_{total} = \left(\mu_x^2 + \mu_y^2 + \mu_z^2\right)^{\frac{1}{2}} 
\]

\[
\alpha_{total} = \frac{1}{3}\left(\alpha_{xx} + \alpha_{yy} + \alpha_{zz}\right) 
\]

\[
\Delta \alpha = \frac{1}{\sqrt{2}} \left[\left(\alpha_{xx} - \alpha_{yy}\right)^2 + \left(\alpha_{yy} - \alpha_{zz}\right)^2 + \left(\alpha_{zz} - \alpha_{xx}\right)^2 \\
+ 6\alpha_{xz}^2 + 6\alpha_{xy}^2 + 6\alpha_{yz}^2\right]^{\frac{1}{2}} 
\]
where $\mu_x$, $\mu_y$, and $\mu_z$ are the tensor components dipole moment, $\alpha_{xx}$, $\alpha_{yy}$, and $\alpha_{zz}$ are the tensor components of polarizability, and $\beta_{xxx}$, $\beta_{yyy}$, and $\beta_{zzz}$ are the tensor components of hyperpolarizability.

**Investigation of pharmacological activity**

The in silico study was performed for the detection of the anti-Alzheimer activity of the phytochemicals of the *Clitoria ternatea*. The structure of the candidate β-Site amyloid precursor protein cleaving enzyme 1 (BACE1) inhibitor (PDB ID: 4B05) was selected as the target macromolecule and was downloaded from the online database Protein Data Bank (https://www.rcsb.org/). The BACE1 is known to produce the toxic amyloid β (Aβ) that leads to the early evolution of pathogens of AD in the human body [33]. Thus, the prevention of BACE1 will be worthwhile in blocking the development of pathogens of AD in the body. This became the major reason for considering BACE1 as a target macromolecule. The crystal structure of the BACE1 target macromolecule developed with X-ray crystallography having a resolution 1.80 Å is shown in Fig. 2.

Molecular docking was done for the selection of the phytochemical having the best binding with the target macromolecule. This will help in selecting the ligand which will have the best inhibition tendency against the AD. The software “AutoDock Vina” (https://vina.scripps.edu/) [34] was used for performing docking. The algorithm for docking was followed by the energy difference of 4 kcal/mol and the grid box centered at $(x, y, z) = (−2.273, 1.428, −15.310)$. Docking results were analyzed by the software “Biovia Discovery Studio Visualizer” (https://discover.3ds.com/discovery-studio-visualizer-download). The best pose of the ligand having the best binding affinity was selected based on the highest number of hydrogen bonds. Cardio-toxicity was also predicted for the molecule having the best binding affinity to check whether the drug-like molecule will create any cardio-related harm to the body after consumption. The cardio-toxicity was investigated by using Pred-hERG 4.2 webserver (http://predherg.labmol.com.br/) [35]. The canonical smiles were used for the prediction of the probability map of the compound. For compounds to be non-cardio-toxic, the confidence value should not exceed 0.26. Different fragments like potency, confidence, and applicability domains are given by the cardio-toxicity analysis.

MD simulation was further computed for protein–ligand complex having the best binding score. The MD simulation will be performed using software “Gromacs 2019.2 version.” The protein macromolecule was prepared using pdb2gmx with Gromos96 54a7 force field, and ligand topology was formed in the GlycoBioChem PRODRG2 Server. The complex was simulated for 100 ns in canonical (amount of substance (N), pressure (P), and temperature (T)—NPT) and isothermal-isobaric (amount of substance (N), volume (V), and equilibrium steps temperature (T)—NVT) ensembles. The parameters like root mean square deviation (RMSD) and root mean square fluctuation (RMSF), binding energy, hydrogen bond, and radius of gyration (Rg) [36, 37].

**Results and discussion**

**Structure and charge analysis**

The structure of phytochemicals Cinnamic acid, Clitorin, Delphindin-3,5-diglucoside, Flavonol 3-O-d-glucoside, Genistein, Kaempferol, Kaempferol-3,7-diglucoside, Linolenic acid, Oleic acid, Palmitic acid, Taraxerol, and Taxaxeron were optimized using DFT (SD 1). The net polarity of the molecules was obtained by optimized geometries of the structures. The values of dipole moment obtained from the optimized geometries are mentioned in SD 2. Clitorin has a dipole moment of 10.28 Debye which is highest among all other considered phytochemicals Cinnamic acid (2.68 Debye), Kaempferol-3,7-diglucoside (6.41 Debye), Flavonol 3-O-n-glucoside (2.17 Debye), Genistein (5.32 Debye),

![Fig. 2 Crystal structure of BACE1 inhibitor (PDB ID: 4B05)](image-url)
Kaempferol (6.15 Debye), Linolenic acid (1.74 Debye), Oleic acid (4.64 Debye), Taraxerol (1.69 Debye), and Taxa-
xeron (3.35 Debye). The high value of dipole moment is due
the high value of intramolecular interactions and these
interactions may lead to high polarizability [38]. The dif-
fERENCE between the dipole moment of Clitorin and other
molecules seems to be high enough to do a fair selection of
Clitorin for the NLO investigation. Based on this, Clitorin
was taken into consideration for further computational and
spectral study. The optimized structure of the Clitorin mol-
ecule is shown in Fig. 3. The Clitorin molecule is a large
molecule with 92 atoms associated with it. The structural
stability of the Clitorin molecule was studied by analyzing
bond lengths and bond angles of the molecule. The struc-
tural parameters are mentioned in SD 3 and SD 4. The total
energy of Clitorin is found at $-1,700,895.541$ kcal/mol.
A higher magnitude of bond length was observed for the
C–C bonds throughout the geometry. On the other hand, the
O–H bonds of the geometry have comparatively lower mag-
nitudes of the bond lengths that can be easily dissociated.
This shows the possibility of the dislocation of free electron
pairs from the –OH groups towards the C–C bonds of the
benzene ring. A similar possibility of the charge dislocation
can be seen by the bond angles. The angles corresponding to
the C–C–C are higher than the angles corresponding to the
C–H–O angles. The Mulliken charge distribution was also
analyzed for the Clitorin molecule and is mentioned in SD
5. The oxygen atoms contribute to the negative charge and
the hydrogen atoms contribute to the making of the positive
charge of the molecule. However, the simultaneous posi-
tive and negative charge contribution of the carbon atoms
is observed for the Clitorin molecule. The –OH groups act
as the electron-donating parts of the molecule and the free
electron charge cloud gets dislocated towards the aromatic
benzene rings.

Molecular electrostatic potential and frontier molecular orbital analysis

The molecular electrostatic potential (MEP) surface of the
Clitorin molecule is illustrated in Fig. 4. MEP surface of the
molecule indicates the nucleophilic (blue) region and the
electrophilic (red) region. The existence of these regions
indicates the possibility of intramolecular charge transfer
(ICT). The nucleophilic regions of the MEP are majorly
induced by –OH groups and the electrophilic regions are
located over the C–C bonds and C = C bonds of the benzene
rings. Thus, the ICT is taking place throughout the mole-
cule’s geometry despite transferring from functional groups.
The MEP surface is shown in Fig. 4a shows the nucleophilic
regions and Fig. 4b shows electrophilic regions.

The frontier molecular orbital (FMO) is used to establish
the stability and the chemical reactivity of Clitorin. These
parameters are considered the frontier of the electrons and
are mentioned in SD 6. The energy corresponding to the
highest occupied and lowest unoccupied molecular orbitals

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![Fig. 3 Optimized geometry of Clitorin molecule obtained using B3LYP/6-311G (+, p) basis set](image-url)
(HOMO–LUMO energies) was derived from the optimization. The molecular orbital surfaces were also computed for the Clitorin molecule and the molecular orbital surfaces are illustrated in Fig. 5. The molecular orbital surfaces are distributed only on the benzene ring made from atoms sequenc-
ing from 38C to 49 °C. However, there is not any major change in the distribution of HOMO–LUMO surfaces. The value of $\Delta E$ for Clitorin was computed as 4.343 eV. The value of $\Delta E$ for Clitorin is found to be less than the $\Delta E$ of the NLO reference materials urea (7.43 eV) and potassium dihydrogen phosphate (KDP) (6.835 eV). The moderate value of $\Delta E$ shows high excitation of electrons which shows the molecule is reactive. The high value of IP (6.597 eV) was reported for the Clitorin which shows the free-electron releasing tendency of the Clitorin electron-donating part. The low value of EA i.e., 2.254 eV was obtained which shows the electron-accepting part of the Clitorin molecule will not face and will easily attract the electron cloud. The low value of CP (-4.425 eV) and high value of $\chi$ (4.425 eV) for Clitorin show the high stability of the molecule. The chemical hardness (2.171 eV) for the Clitorin molecule was recorded as positive. This shows the Clitorin molecule stays stiff and rigid. All the computed parameters show that the Clitorin molecule is chemically reactive.

**Vibrational spectra analysis**

As the polarizing ability of any compound is proportional to the Raman intensity [39]. The Raman modes with high intensity are illustrated in Fig. 6. The details of the high-intensity modes are mentioned in SD 7. The torsional bending ($\delta$) of the C–H bonds is observed from 800 to 1300 cm$^{-1}$. The mode $\delta_{CH}$ at 855.17 cm$^{-1}$ frequency has the highest value of Raman intensity at 1808.13 cm$^{-1}$. The mode $\nu_{CC}$ with high intensity 5412.76 cm$^{-1}$ is observed with frequency 1679.64 cm$^{-1}$. The C–O bonds stretch linearly and the mode with the highest intensity of 3798.18 cm$^{-1}$ is observed with a frequency of 1771.92 cm$^{-1}$. The asymmetric stretching ($\alpha_{CH}$) of C–H bonds are observed around 2978–3149 cm$^{-1}$. The
α\textsubscript{CH} with frequency 2991.83 cm\textsuperscript{-1} has the highest Raman intensity (1037.4 cm\textsuperscript{-1}). The symmetric stretching (\(\nu\)\textsubscript{CH}) for C–H bonds have the highest mode with an intensity of 1209.95 cm\textsuperscript{-1}. The computed values of Raman intensity are extremely high. These high values are due to the presence of conjugated-\(\pi\) electrons of C–C bonds. The high values of Raman intensity give rise to the high value of polarizability of molecule \([40]\). Thus, the high value of Raman intensity was in good agreement with the high NLO activity of the Clitorin molecule. Moreover, it can be considered the cause for high reactivity of the Clitorin with macromolecules.

Absorption spectra analysis

The electronic spectra were computed by performing energy optimization of the Clitorin molecule by the TD-DFT method. UV–Vis spectra of Clitorin are illustrated in Fig. 7 and the transition details are mentioned in SD 8. The transition S\textsubscript{0} \rightarrow S\textsubscript{1} at 341.82 nm wavelength and 3.62 eV excitation energy majorly imparts in the formation of the broad absorption band of the Clitorin molecule. These transitions are induced by the presence of \(\pi \rightarrow \pi^*\) and \(n \rightarrow \pi^*\) bonds as these bonds produce strong absorption peaks around 300–400 nm and absorption boundary at 450–460 nm. The transitions S\textsubscript{0} \rightarrow S\textsubscript{2} and S\textsubscript{0} \rightarrow S\textsubscript{3} that lead to the absorption spectra are at wavelengths 327.83 and 299.42 nm with excitation energies of 3.78 eV and 4.74 eV respectively. Thus, the high excitation energies of these transitions show the high chemical reactivity of the Clitorin molecule. This leads to the high binding ability of the ligand to the active binding sites of the macromolecule \([41]\). Thus, there is a probability of Clitorin having higher binding affinity at the binding sites of the receptor among all the other ligands.

Nonlinear optical analysis

The molecular packing and intramolecular and intermolecular interactions have a straight association with the physicochemical properties of the molecules \([42]\). Thus, the quantum chemical calculations were carried out to predict the nature of electric dipole moment (\(\mu\)\textsubscript{total}), polarizability (\(\alpha\)\textsubscript{total}), and first-order hyperpolarizability (\(\beta\)\textsubscript{total}). The tensor components of \(\mu\)\textsubscript{total}, \(\Delta\alpha\), and \(\beta\)\textsubscript{total} were computed by polar calculations and are mentioned in Table 1. The value of \(\mu\)\textsubscript{total} for Clitorin was computed as 4.046 Debye. The comparison gave us that the \(\mu\)\textsubscript{total} for Clitorin was 2.5 times that of the most generally used reference NLO material, urea (1.527 Debye). Highly raised values of \(\alpha\)\textsubscript{total} (62.514 \(\times\) \(10^{-24}\) esu) and \(\Delta\alpha\) (111.87 \(\times\) \(10^{-24}\) esu) for Clitorin are reported. \(\alpha\)\textsubscript{total} of Clitorin was obtained 11 times higher than urea (5.664 \(\times\) \(10^{-24}\) esu) and \(\Delta\alpha\) was nearly 18 times that of urea (6.304 \(\times\) \(10^{-24}\) esu). The value of \(\beta\)\textsubscript{total} for Clitorin 5.415 \(\times\) \(10^{-30}\) esu was nearly 7 times higher than urea (0.781 \(\times\) \(10^{-30}\) esu). For the validation of Clitorin being a good candidate as an NLO material, the \(\beta\)\textsubscript{total} for Clitorin was also compared with some already worked compounds that were experimentally proven NLO active materials and had been used in various applications in nonlinear optics. Such compounds were mentioned in Table 2. Compounds like 3-nitroaniline, phenylurea, thiosemicarbazone derivatives, thiourea-glutaric acid, indole-7-carboxaldehyde, 2,4,6-triaminopyrimidine, and ANDIROBIN have low values of \(\beta\)\textsubscript{total} than that of Clitorin. These values were quite less than that of Clitorin which validates the good candidature of Clitorin being nonlinearly active.
In silico study

Molecular docking analysis

Molecular docking was performed to check the binding ability of the considered ligands with the BACE1 inhibitor (PDB ID: 4B05). The docking results were recorded after performing ten times docking with every ligand. The docking score of all the considered ligands is mentioned in SD 9. Clitorin, Delphinidin 3,5-diglucoside, Taraxerol, and Taraxeron were the molecules that have the highest binding affinity of $-10.4 \text{ kcal/mol}$, $-10.2 \text{ kcal/mol}$, $-10.7 \text{ kcal/mol}$, and $-10.6 \text{ kcal/mol}$ respectively. These binding scores are extremely higher than the other ligands Cinnamic acid ($-5.8 \text{ kcal/mol}$), Genistein ($-7.9 \text{ kcal/mol}$), Kaempferol ($-7.9 \text{ kcal/mol}$), Kaempferol-3,7-diglucoside ($-9.3 \text{ kcal/mol}$), Linolenic acid ($-5.8 \text{ kcal/mol}$), Octadeca-9,12-dienoic acid ($-4.9 \text{ kcal/mol}$), Oleic acid ($-5.7 \text{ kcal/mol}$), and Palmitic acid ($-4.8 \text{ kcal/mol}$).

The validity of the docking scores was supported by analyzing the number of hydrogen bonds and the dipole moment of Clitorin, Delphinidin 3,5-diglucoside, Taraxerol, and Taraxeron. The details of the number of hydrogen and hydrophobic bonds and dipole moment of these ligands are mentioned in SD 10. The binding of ligand to the binding sites of the macromolecule is considered stable by the raised value of hydrogen bonds. As hydrogen bonds involve the availability of high electronegativity, they are the strongest bonds among the other known strong intermolecular bonds. Similarly, hydrophobic bonds are the strongest among the other known weak intermolecular bonds. Thus, the more the number of hydrogen and hydrophobic bonds associated with the binding site, the more will be the strength of binding. Clitorin thus has the highest number of 7 hydrogen and 6 hydrophobic bonds associated with the binding site. The dipole moment that is the virtue of the high polarity of the molecule is also highest for Clitorin. High polarity leads to high reactivity.

### Table 1

| Component | Clitorin | Urea |
|-----------|----------|------|
| $\mu_x$  | -3.35    | 1.28 |
| $\mu_y$  | 2.082    | 0.00004 |
| $\mu_z$  | 0.902    | -0.830 |
| $\mu_{total}$ | 4.046 | 1.527 |

| Component | Clitorin | Urea |
|-----------|----------|------|
| $\alpha_{xx}$ | 435.257 | 39.260 |
| $\alpha_{xy}$ | 3.553 | 24.690 |
| $\alpha_{yy}$ | 433.886 | 38.219 |
| $\alpha_{zz}$ | -51.058 | 0.2947 |
| $\alpha_{xz}$ | -26.335 | -1.159 |
| $\alpha_{yz}$ | 396.335 | 0.454 |
| $\alpha_{zt}$ | 62.514 $\times 10^{-24}$ | 5.664 $\times 10^{-24}$ |
| $\Delta \alpha$ | 111.87 $\times 10^{-24}$ | 6.304 $\times 10^{-24}$ |

| Component | Clitorin | Urea |
|-----------|----------|------|
| $\beta_{xx}$ | -82.66 | 24.729 |
| $\beta_{xy}$ | 300.061 | 0.006 |
| $\beta_{yy}$ | 210.531 | 32.896 |
| $\beta_{zz}$ | -226.921 | -0.499 |
| $\beta_{xz}$ | -318.121 | -61.443 |
| $\beta_{yz}$ | 46.292 | 19.374 |

### Table 2

| Material | Hyperpolarizability | Reference |
|----------|---------------------|-----------|
| 3-nitroaniline | $1.347 \times 10^{-30}$ | [43] |
| Phenylurea | $2.043 \times 10^{-30}$ | [44] |
| (E/Z)-4-benzyl-1-(1-ferrocenylethyl) thiosemicarbazone | $2.1433 \times 10^{-30}$ | [45] |
| (E/Z)-4-(4-chlorobenzyl)-1-(1-ferrocenyl-ethyl) thiosemicarbazone | $1.293 \times 10^{-30}$ | [45] |
| (E/Z)-4-(2-bromo benzyl)-1-(1-ferrocenylethyl)thiosemicarbazone | $3.316 \times 10^{-30}$ | [45] |
| thiourea-glutaric acid | $3.57 \times 10^{-30}$ | [46] |
| Indole-7-carboxaldehyde | $3.96 \times 10^{-30}$ | [47] |
| 2,4,6-triaminopyrimidine | $4.73 \times 10^{-30}$ | [48] |
| methyl-2([[1k,2k,3k]-[1aS, 4S, 4aS, 8aS]-4-(furan-3-yl)-4a-methyl-8-methylene2-oxooctahydroxireno(2,3-d]isochromen-7-yl]-2,6,6-trimethyl-5-oxocyclohex-3-en-1-yl]acetate (ANDIROBIN) | $3.759 \times 10^{-30}$ | [49] |
Fig. 8  Target receptor of AD a before and b after the binding of inhibitor 4BO5

Fig. 9  2D view hydrogen bond interactions associated to protein–ligand complex after docking
Clitorin has a dipole moment equal to 3.62 Debye which is higher than Delphinidin 3,5-diglucoside (2.47 Debye), Taraxerol (1.76 Debye), and Taraxeron (3.48 Debye). Therefore, analysis of the docking score, hydrogen bonds, hydrophobic bonds, and dipole moment suggests Clitorin as the most reactive and finely bonded ligand. Binding site of the Clitorin on the receptor is shown with the help of Fig. 8. The details of the hydrogen and hydrophobic bond interactions formed by binding of Clitorin with protein were shown in SD 11 and the 2D illustration of all the interactions is shown in Fig. 9. 3D view of protein–ligand interaction within the surface due to H bonds is shown in Fig. 10. Thus, the protein–ligand complex obtained after docking was taken for MD simulation.

**MD simulation analysis**

The physical movements of the complex molecules were analyzed by MD simulation. Cocrystal ligand AZD3839 is an experimental inhibitor of BACE1 that is often used for comparing the simulation results obtained from the simulation of the protein–ligand complex. The cocrystal ligand AZD3839 and protein complexed with ligand were simulated for 100 ns of time trajectory and the results are compared. This helps in obtaining dynamic data at atomic spatial resolution. The RMSD, RMSF, binding energy, hydrogen bond, and Rg were obtained as the results of simulation and the results are illustrated in Fig. 11.

RMSD value of BACE1 with Clitorin and cocrystal ligand AZD3839 seems to fluctuate between 0.2 and 0.5 nm. The average value of the RMSD value of the complex is around 0.35 nm. The RMSD trajectory of the complex is slightly higher than the cocrystal ligand AZD3839. This variation is due to the position restraints. The energy minimization in structures results in these variations. The RMSF value of the complex shows a certain rise around residue 50, 180, and 300. However, the RMSF trajectory of BACE1 complexed with Clitorin and cocrystal ligand AZD3839 does not differ much. The average value of the RMSF of the complex is around 0.3 nm. The highest rise of the RMSF trajectory of the complex is 0.6 nm at a 50 residue number. Thus, the RMSD and RMSF values of the complex show that the ligand binds strongly to the protein binding site and protein remains unaltered due to the presence of the...
ligand. The RMSD and RMSF fluctuations are illustrated in Figs. 11a and b.

Rg shows the compactness of the protein–ligand complex. The trajectories of Rg of BACE1 complexed with Clitorin and cocrystal ligand AZD3839 have variation that shows the stability of the complex. The Rg values of the complex are lower than the values of the Rg of cocrystal ligand AZD3839. The complex value fluctuates between 2.1 and 2.2 nm during the simulation time. The trajectory reaches the highest peak of 2.2 nm five times at 37 ns, 45 ns, 76 ns, and 85 ns, and finally rises after 85 ns up to 2.25 nm. These fluctuations are less than that of cocrystal ligand AZD3839. The Rg fluctuations of both systems are illustrated in Fig. 11c. The lesser value of Rg denotes the compactness of the complex.

The hydrogen bond count imparts in stabilizing the complex. The hydrogen bond count of the complex fluctuates between 0 and 2 with an average value of 1. This value matches the hydrogen bond count of the cocrystal ligand AZD3839 and thus, imparts the stability of the complex. The trajectory tracing hydrogen bond number is illustrated in Fig. 11d.

The implementation of isothermal-isobaric ensembles results in thermodynamic parameters like Van der Waals energy, electrostatic, polar solvation, solvent accessible surface area (SASA), and binding free energy. MM-PBSA of binding free energy of BACE1 & AZD3839 and BACE1 & Clitorin complexes were computed by MD simulation and the computed values are mentioned in Table 3. The high values of Van der Waals energy (−198.863 kJ/mol) were computed for the Clitorin complexed with BACE1. The high value of Van der Waals energy shows the high binding ability of the ligand to the binding site. The computed SASA values indicate the availability of sufficient surface area for the ligand attack that enables the ligand to bind properly to the protein. The low values of SASA for both BACE1 & AZD3839 (-19.633 kJ/mol) and BACE1 & Clitorin complex (-19.110 kJ/mol) are low enough to justify the better complexion of ligand to the protein. The value of polar solvation energy for BACE1 & AZD3839 (92.651 kJ/mol) is higher than the same for BACE1 & Clitorin complex (58.163 kJ/mol). This shows the availability of short-range dispersion interactions
that are responsible for the formation of the cavity inside the macromolecule where the ligand binds. The binding free energy of the BACE1 & Clitorin (−164.643 kJ/mol) complex was higher than the BACE1 & AZD3839 (−155.552 kJ/mol). This shows the better binding of the protein–ligand complex than the cocrystal ligand and BACE1 complex.

Protein–ligand interactions from MD simulation

The position of the ligand bonded to the macromolecule at 0 ns (position after docking) and 100 ns was monitored and the position of the ligand bonded to protein at 0 ns and 100 ns of MD simulation is illustrated in Fig. 12. The amino acid residues involved in interactions like Vander Waal’s, pi-anion, pi-donor, pi-alkyl, and pi-stacked interactions of the protein–ligand complex at a 0 ns, b 50 ns, and c 100 ns of the MD simulation. The specific type of interaction is indicated by a single color that is labeled at the bottom of the image.

Fig. 13 The 2D representation of amino acid residues involved in Vander Waal’s, pi-anion, pi-donor, pi-alkyl, and pi-stacked interactions of the protein–ligand complex at a 0 ns, b 50 ns, and c 100 ns of the MD simulation. The specific type of interaction is indicated by a single color that is labeled at the bottom of the image.

Fig. 14 Probability map of Clitorin showing hERG blockage promoting regions in pink shade.
Waal’s, pi-anion, pi-donor, pi-alkyl, and pi-stacked interactions of the protein-ligand complex at 0 ns, 50 ns, and 100 ns of the MD simulation are also analyzed to justify the variation in protein-ligand interactions. A gradual rise in the number of residues involved in the binding is observed during the simulation. The two-dimensional representation of residues involved in the interactions is shown in Fig. 13. At 0 ns of the simulation, only Phe107, Asp32, Trp75, and Ile117 residues impart in forming pi-pi, pi-anion, pi-donor hydrogen bond, and pi-alkyl hydrophobic interactions (Fig. 13a). Except for Trp75 which forms the hydrogen bond, the rest of the residues form non-polar hydrophobic interactions. The number of residues increased at the 50 ns of simulation. Ala39, Tyr71, Val69, Ala99, and Leu79 (Fig. 13b) seemed to be associated with forming pi-sigma, pi-pi, and pi-stacked bonds. Thus, all the interactions at 50 ns were hydrophobic and hence non-polar. The residue count at 100 ns was increased to 15 residues shown in green color in Fig. 13c and they participate in the Vander Waals interactions. Ala39, Arg127, Ala99, and Leu79 residues make the pi-alkyl interactions and Tyr71 makes up the pi-pi stacked bonds. One pi-sigma interaction with the involvement of Val69 was also seen at 100 ns of the simulation time. Thus, the major part of the interactions at 100 ns is Vander Waal interactions that are polar. Thus, there is a change in interacting residues at 0 ns, 50 ns, and 100 ns of the simulation as well as in the chemical polarity of the complex.

**Cardiac toxicity analysis**

The probability map of Clitorin illustrated in Fig. 14 indicates that –OH group is covered from the pink color. This pink-colored region shows that –OH part of the molecule imparts in the decrement of hERG blockage region. The confidence value for the Clitorin is 0.2 which doesn’t exceed the limit. Thus, the probability map justifies that the Clitorin is an active potent against cardiotoxicity with 60% confidence value.

**Conclusion**

This study deals with the identification of a new organic NLO material, Clitorin, that is a phytochemical of the *Clitoria ternatea* plant. The computational investigation done for the identification of novel characteristics of NLO material of Clitorin better reveals the target properties. The widespread nucleophilic and electrophilic regions of the MEP surface and charge variation in –OH groups, C–C bonds, and C = C bonds validate the availability of ICT in Clitorin. The spectral features like high Raman intensity and high wavelength for the electronic transitions were also observed for Clitorin. These factors better validate the dislocation of charge from the –OH groups towards the C–C bonds and C = C bonds of benzene rings. Thus, these parameters reveal the high reactivity of Clitorin. High reactivity induces the high magnitudes of $a$ and $b$ parameters. High reactivity, although, was seemed during the docking and simulation also. Having the best binding affinity and the highest number of hydrogen bond interactions, Clitorin binds with the BACE1 target macromolecule of AD. The SASA values reveal the availability of a high interacting surface of protein towards the ligand. This leads to the high Vander Waal’s interaction energy. The Vander Waal’s interaction energy justifies the high chemical stability of the complex formed from docking. Thus, this study introduces a chemically reactive compound Clitorin that has NLO responses high enough to be used for experimental validations. Concurrently, it has high inhibiting potentiality against the BACE1 receptor of AD that gives rise to pathogens of AD in humans. Thus, it can be considered for clinical trials as an inhibitor against AD in the future.

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**Availability of data and material** Phytochemical structure: https://cb.imsc.res.in/imppat/home. Extension conversion: http://openbabel.org/wiki/Main_Page. Optimization: https://gaussian.com/. Data analysis: https://gaussian.com/gaussview6/. Graph plotting: https://www.origi- nlab.com/. Protein structure: https://www.rcsb.org/. Molecular docking: https://vina.scripps.edu/. Docking analysis: https://discover.3ds.com/discovery-studio-viewer-download. MD simulation: https://www.gromacs.org/. Cardiotoxicity analysis: http://predherg.labmol.com.br/

**Declarations**

**Conflict of interest** The authors declare no competing interests.

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