Enamine Barbiturates and Thiobarbiturates as a New Class of Bacterial Urease Inhibitors

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Abstract: Urease is a therapeutic target associated with several important diseases and health problems. Based on our previous work on the inhibition of glucosidase and other enzymes and exploiting the privileged structure assigned to the (thio)barbiturate (pyrimidine) scaffold, here we tested the capacity of two (thio)barbiturate-based compound collections to inhibit urease. Several compounds showed more activity than acetohydroxamic acid as a standard tested compound. In addition, by means of a conformational study and using the Density Functional Theory (DFT) method, we identified energetically low-lying conformers. Finally, we undertook a docking study to explore the binding mechanism of these new pyrimidine derivatives as urease inhibitors.

Keywords: pyrimidine-trione; barbituric; thiobarbituric; urease inhibitors; DFT

1. Introduction

Many microorganisms use nitrogen from urease for growth (urea amidohydrolase, EC: 3.5.1.5). Urease is an enzyme that catalyzes the hydrolysis of urea into ammonia and carbon dioxide. There is a link between health complications and ammonia production by urease [1]. In animals and humans, low pH of the stomach allows microbial strains to survive, multiply, and grow, resulting in pyelonephritis, hepatic coma, gastric carcinoma, gastric lymphoma, kidney stones, and peptic ulcer complications [2,3]. The treatment of bacterial infection with therapeutics has often proven ineffective due to drug resistance. Thus, there is a clear need for alternatives or novel treatments. Given the
involvement of ureases in various diseases, pharmaceutical research has channeled considerable efforts into discovering potent and safe urease inhibitors [4–7].

Molecules with a binding site that can chelate metals are an interesting challenge and could be a promising line of action to prevent the adverse effects of ureolytic bacterial infections in humans [8]. In this regard, many types of potent urease inhibitors have been designed [9], such as dihydropyrimidines [10], urea derivatives [11], semicarbazones [12], Schiff bases [13], hydroxamic acid derivatives [14], piperazines [15], bisoumarines [16], benzimidazoles [17], and sulfonamides [18].

Enaminone molecular scaffolds have been found in many synthetic drugs and natural products [19,20]. Specially, pyrimidine-based enamines have demonstrated an array of biological activities owing to the presence of the alkenylamine moiety (R=CH–) in their structures, and are capable of strong bonding with metal ion chelates in biological systems [21,22].

The barbituric and thiobarbituric acids [(thio)pyrimidine trione analog derivatives] have been reported urease inhibitors [12]. These can be considered privileged structures because they have antifungal [23], antimicrobial [24,25], anti-adiponectin [26], anti-sclerosis [27], anti-convulsing [28,29], antiglycation [30], and α-glucosidase inhibitory [30] properties. Several compounds A, B, and C have been reported, and showed anti-urease activity comparable to acetohydroxamic acid (Figure 1) [31,32].

Figure 1. Some examples of bacterial urease inhibitors.

In this context, and as a continuation of our extended medicinal chemistry program based on the barbituric and thiobarbituric acid moieties [30,33–44], here we examined the in vitro anti-urease activity of a collection of 1,3-dimethylbarbiturate-enamine compounds and their analogs, thiobarbiturate derivatives. In addition, molecular docking studies were performed to evaluate the molecular interactions of the newly synthesized compounds with selected drug targets (PDB ID 4GY7).

2. Materials and Methods

The synthesis and the full characterization of compounds 3a, 3d, 3h, 3j–p, 4, and 5 (Table 1) have been previously described by our group [30,33]. The rest of the compounds studied were prepared following the previously described method. The yields and full characterization are provided in the Supplementary materials.

2.1. In Vitro Urease Inhibition Assay Protocol

The urease inhibition assay was performed spectrophotometrically following the manufacturer’s instructions [45–47]. The source of urease is Jack bean Urease and the full protocol provided in the supplementary materials. In the present study, urease was preincubated with the inhibitors for a period of 15 min, which proved to be sufficient in our studies.
| Compound | Structure | Urease Inhibition IC<sub>50</sub> ± SEM [µM] |
|----------|-----------|------------------------------------------|
| 3a       | ![Structure](image1.png) | 9 ± 2                                    |
| 3b       | ![Structure](image2.png) | NA                                       |
| 3c       | ![Structure](image3.png) | 26 ± 1                                   |
| 3d       | ![Structure](image4.png) | 8 ± 0.3                                  |
| 3e       | ![Structure](image5.png) | 11 ± 0.3                                 |
| 3f       | ![Structure](image6.png) | 10 ± 0.6                                 |
| 3g       | ![Structure](image7.png) | NA                                       |
| 3h       | ![Structure](image8.png) | 6.4 ± 0.3                                |
| 3i       | ![Structure](image9.png) | 66 ± 2.4                                 |
| 3j       | ![Structure](image10.png) | 10 ± 0.9                                 |
| 3k       | ![Structure](image11.png) | 11 ± 1.2                                 |
2.2. DFT Calculation

The molecular building and geometric optimization of all 18 compounds was performed by DFT using Gaussian 03W software package with basis set method B3LYP/6-31+G(d,p). The Gauss-view program was used for molecular visualization. Geometrical, energetic and electronic parameters were calculated using structures optimized at 298 K and 1 atm. The Gibbs free energy equation ($\Delta G = -RT \ln K$) was used to search conformational space.
2.3. Molecular Docking

Molecular docking studies were conducted to rationalize the binding mode of the pyrimidine derivatives using the MOE package [Molecular Operating Environment, 2016.0810; Chemical Computing Group ULC, 1010 Sherbrooke St. West, Suite #910, Montreal, QC, Canada, H3A 2R7, 2017]. The three-dimensional structure of the urease protein was retrieved from the Protein Data Bank (PDB ID 4GY7) [48]. Water molecules were removed, and the protonation step was performed. Energy minimization using default parameters was performed to achieve a stable conformation. The optimized geometries of the pyrimidine derivatives were docked to the active site of the urease protein. For docking studies, an induced fit protocol with the triangular matcher placement method and London dG scoring function were used. The interactive docking procedure was performed for all the optimized conformers of each compound in the active site. A score was assigned to each docked compound based on its fitting in the binding cavity and its binding mode (docking study protocol are provided in the Supplementary materials).

3. Results and Discussion

3.1. Synthesis

Derivatives 3a–p, 4, and 5 (Table 1) were straightforwardly synthesized by nucleophilic substitution of the corresponding amines with enaminone scaffolds 1 [49] and 2 [50] in methanolic solution, following the reported method [30,33] (Scheme 1). Due to the potential tautomerism of these families of compounds, X-ray crystallography was carried out on two compounds (3b and 3k) to confirm the structures. (1H-NMR and 13C-NMR spectra, Figures S1–S21, and X-ray data of compounds 3b and 3k are provided in the Supplementary materials).

![Scheme 1. Synthesis of 3a–p, 4, and 5.](image_url)
3.2. In Vitro Evaluation of Urease Inhibition

The capacity of synthesized derivatives of N,N-dimethylpyrimidine trione (3a-i and 4) and N,N-diethyl-thio pyrimidine dione (3j-p and 5) to inhibit urease in vitro was examined and compared with that of acetohydroxamic acid (AHA) (IC\textsubscript{50} = 20 ± 0.4 µM), as a standard compound (Table 1).

Of note, the tested compounds showed potent urease inhibition activity, twelve of them (3a, 3d, 3e, 3f, 3h, 3j, 3k, 3l, 3m, 3n, 3o, and 4) exerting greater activity than the reference acetohydroxamic acid. The most active member between this series is compound 3h which the structure attributed the pyridine moiety. Separation the pyridine ring from the enaminone functionality in compound 3j with CH\textsubscript{2} group, the activity has been decreased compared to the most active member 3h. On the other hands, compounds 3k, and 3o showed superior activity to their counterparts 3b and 3g. Indeed, 3l and 3o showed activity similar to that of 3d, and 3f. The chloro atom in para-position in compound 3m, the activity has been decreased compared to the ortho position of compounds 3f, and 3n. Compounds 3c, 3i and 3p with sulfonic, cyclic amide or sulfonamide functionalities shown less activity compared to standard drug. The cyclohexyl group in compound 3e increases the activity with 1.8 folds than the control. it was observed that in case of compounds 4 and 5 the presence of the N,N'-diethyl group and the S atom in the thiobarbituric derivative 5 make them more voluminous compared to its analogs N,N'-dimethyl and the “O” atom in the barbituric derivatives 4. Finally, compounds 3b, and 3g were not active.

However, there are some discrepancies regarding whether AHA is or is not a competitive inhibitor to the active site of the urease enzyme [51,52]. Therefore, some limitations could be applied, and we think that comparisons are important.

3.3. Docking Study

Molecular docking studies have the potential to identify novel drug-like molecules that display high binding affinity for the target protein, and they can facilitate the understanding of biological activity data with the purpose of designing new compounds with improved activity. Hence, we extended our study to explore the conformational space and binding orientation of the synthesized pyrimidine derivatives. The binding conformations of these compounds were explored by MOE-Dock module implemented in the MOE program [53]. The optimized conformers obtained from the Gaussian program were docked to the active site of urease. The docking results indicated that all the conformers were well accommodated inside the active site and were stabilized by various hydrophilic, hydrophobic and van der Waals interactions. The residues involved in these interactions were Arg439, Ala440, His492, Asp494, His593, His594, Asp633 Ala636 and Met637. Compound 3h (IC\textsubscript{50} = 6 ± 0.3 µM) showed strong inhibitory potential against urease as compared to 3b (inactive) due to methyl substitution at the para position of the pyridine ring. Compound 3h showed good interactions with the residues of the active site by coordinating with the bi-nickel center, via its carbonyl group at the pyrimidine ring and anchoring itself in a way that permitted strong interaction of the carbonyl group with the two nickel ions. Additionally, the two carbonyl groups on the pyrimidine ring acted as hydrogen bond acceptors and mediated hydrogen bond interaction with His492 (2.3 Å) and Arg439 (2.04 Å) of the binding pocket (Figure 2b).
Figure 2. Depiction of the docking results of low energy conformer: (a) 3a, (b) 3h, (c) 3d, (d) 3j, (e) 3m, (f) 3f, (g) 3p, and (h) 5. The key residues are presented as sticks models and nickel atoms are shown as green circles.
Compound 3a (IC50 = 9 ± 2 μM), another active compound of the series, established various potential interactions with the active site of urease, as well as with nickel ions. The oxygen atom of the morpholine ring was involved in the chelation process with the nickel center in the catalytic site. This compound was further stabilized by two typical hydrogen bonds with His492 (2.8 Å) and Arg609 (3.0 Å). Apart from hydrogen bond interactions, a tetramine of the compound participated in the salt bridge interaction with the negatively charged Asp494 residue (Figure 2a). Compounds 3d and 3l (IC50 = 8 ± 0.3 μM) had almost similar structures, biological activities and patterns of binding interactions with the active site residues. The only difference was the interaction with metal ions. The lone pair of electrons for sulfur (C=S) of 3l mediated strong interaction with the two nickel ions and hydrogen bond interaction with Arg609 (2.4 Å) and Met637 (2.8 Å), while in case of 3d, the carbonyl group on the pyrimidine ring was involved in the interaction with the nickel ions as depicted in Figure 2c.

Compounds 3j (IC50 =10 ± 0.9 μM) and 3k (IC50 =11 ± 1 μM), which showed good biological activity, presented similar types of interactions. The additional effectiveness of 3j compared to 3k was due to the absence of a methyl group at the meta position of the benzene ring, which allowed the compound to establish close contacts with the active site residues (Figure 2d). The ring nitrogen of the pyridine is involved in two productive hydrogen bond interaction with His492 and Met637 at 2.54 and 3.4 Å, respectively. Moreover, the C=O of pyrimidine rind mediated a potential hydrogen bond interaction with the NH of His593 at 2.59 Å, while the compound was further stabilized through hydrophobic interactions with Ala440, Cme592, and His594.

The compounds with electron-withdrawing substitution, especially halogens, showed noteworthy inhibitory potential. For 3m, 3n, and 3o, the presence of halogen atoms with their respective position on the benzene ring affected the binding pattern and orientation of the compound within the pocket (Figure 2e). Compounds 3f (o-chloro phenyl) and 3o (o-iodo phenyl) with IC50 = 10 ± 0.6 μM showed two hydrogen bond interactions with His492 and Ala440 (Figure 2f), while 3m and 3n were stabilized by a single hydrogen bond with His492. These compounds also displayed hydrophobic interaction with Ala440, His593 and Met637. Compounds 3c (IC50 = 26 ± 1 μM) and 3p (IC50 = 22 ± 0.8 μM), both with moderate biological activity, established two hydrogen bond interactions with His492 and Ala440. However, only one interaction with the metallocenter was observed (Figure 2g). The compounds with low biological activity, as compared to the standard and other pyrimidine derivatives such as 3i (IC50 = 66 ± 2.4 μM) and 5 (IC50 = 42 ± 2.3 μM), established single hydrogen bond interaction with the active site residue Arg439 while compound mostly established by hydrophobic interaction with the active site residues (Figure 2h). However, no interaction with nickel was observed for 3i.

Indeed, the urease inhibition capacity of the synthesized pyrimidine derivatives is attributed to the mutual contribution of the distinct substitutions they bear. However, interaction with the metallocenter and hydrophilic interaction with Ala440, His492, His593, and Met637 are found to be crucial for the activity of these compounds.

3.4. Density Functional Theory (DFT)

The physicochemical properties and frontier molecular orbitals (FMOs) of the new enaminone compounds play a crucial role in enhancing bioactivity. Khon-Sham’s DFT approach with the B3LYP method was used for geometry optimization [34]. The electron-donating and -withdrawing ability of a compound can be explained by its HOMO and LUMO. The higher the energy value of HOMO, the greater the electron-contributing ability of the compound. The energy difference between HOMO and LUMO is an established parameter to measure the electron conductivity or degree of intermolecular charge transfer, which also affects bioactivity [35]. In this study, to examine the urease inhibition capacity of the pyrimidine derivatives, we randomly selected compounds for comparison with DFT results.

The results of HOMO-LUMO energies were plotted against the biological activity of the pyrimidine derivatives (Table 2). Good correlation was observed between biological activity and the energies of
the LUMO orbitals. Compounds 3h, 3k, 3f, and 3o showed significant LUMO energy of −1.89, −2.24, −1.96 and −2.26 eV, respectively.

| Compound | IC_{50} (μM) | HOMO (eV) | LUMO (eV) | E_{gap} (eV) |
|----------|--------------|-----------|-----------|-------------|
| 3a       | 9 ± 2        | −5.79330622969 | −1.21770997548 | −4.57559625421 |
| 3d       | 8 ± 0.3      | −5.88881823 | −1.7037050982 | −4.76988562237 |
| 3f       | 10 ± 0.6     | −6.2202530345 | −1.96248588673 | −4.2577614777 |
| 3h       | 6 ± 0.3      | −6.24583174683 | −1.8895934518 | −4.35627240165 |
| 3k       | 11 ± 1       | −5.70922301574 | −2.24439595033 | −3.46482706541 |
| 3o       | 9 ± 0.3      | −5.75357759138 | −2.25963433214 | −3.49394325924 |

The visualization of the HOMO-LUMO orbitals of 3h reflects the localization of FMO (frontier molecular orbital) (Figure 3). The negative and positive phases of orbitals are shown in green and red, respectively. For 3h, HOMO is localized on the pyrimidine ring, distal pyridine moiety and carbonyl group, whereas LUMO is on the pi-bond adjacent to the pyrimidine ring. In contrast, for 3o, electrons are delocalized mainly on the benzene ring with iodine group as a substituent. The influence of LUMO energy on the inhibitory activity of the compounds might be due to the presence of halogen substitution and the pyridine ring. The halogen substitution at ortho and para positions made a great contribution to the urease inhibition capacity of the derivatives. Notably, neither HOMO nor LUMO were located on the methyl group, which would explain why this group contributed the least to binding with the protein.

The energy gap for the most active compound 3h of the series was −4.35. A similar energy difference was observed for 3k, 3f, and 3o. The smaller the difference in HOMO-LUMO energies, the greater the chemical reactivity. For any potential interaction, electron transfer from high-lying HOMO to low-lying LUMO is always energetically favorable. Given this consideration, 3h, 3k, 3f, and 3o possess good activity, which correlated well with urease inhibitory activity.

![Figure 3](https://example.com/figure3.png)

**Figure 3.** The optimized geometries and surfaces of the HOMO (Highest occupied molecular orbital)-LUMO (Lowest unoccupied molecular orbital) of 3h, 3k and 3o obtained at the B3LYP/6-31G (d, p) level.

Molecular electrostatic potentials (MEP) were run for compounds 3h, 3k, and 3o at B3LYP/6-31G (d, p), providing information about the sites reactive towards nucleophilic and electrophilic attack, together with hydrogen-bond interactions. The potential of electrostatic interaction at the surface is shown by blue, red and green, representing the sites for positive, negative and no electrostatic potential, respectively (Figure 4). Moreover, the positive and negative regions of the maps were responsible for the nucleophilic and electrophilic reactivity of the compounds.
4. Conclusions

Based on our previous work addressing the inhibition of glucosidase and taking advantage of the privileged structure associated with the (thio)barbiturate (pyrimidiene) scaffold, here we have identified compounds with high in vitro anti-urease activity. All conformations of the pyrimidine derivatives were optimized with the B3LYP/6–31G method. The DFT results for the compounds correlated well with the experimental data, which indicated that the presence of the pyridine ring and electron-withdrawing groups, especially halogens (compounds 3m, 5n, and 3o), play an important role in conferring urease inhibition activity. Moreover, the molecular docking studies further helped to explain the experimental results. Because of our findings, compounds 3a, 3d, 3h, 3k, and 3o emerge as potential leads for the development of urease inhibitors. In addition, the obtained results confirm that the use of privileged structures for medicinal chemistry programs is an excellent strategy for identifying hits and leads. Further investigation will be carried out in the future for urease enzyme competitive inhibition.

Supplementary Materials: The following are available online at http://www.mdpi.com/2076-3417/10/10/3523/s1. The details of the synthesis and characterization of the studied compounds; X-ray data of compounds 3b and 3k; biological activity assay; docking study protocol and copies of NMR spectra (Figures S1–S21) of the synthesized target compounds are provided as Supplementary materials.

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