Computational Analysis of Amiloride Analogue Inhibitors of Coxsackievirus B3 RNA Polymerase

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Abstract

Coxsackievirus B3 (CVB3) is a picornavirus that is responsible for a significant proportion of human myocarditis. However, no antiviral treatment is currently available to treat this disease or indeed any picornaviral infections. Previously it was shown that amiloride and its derivative 5-(N-ethyl-N-isopropyl)amiloride inhibit the in vitro enzymatic activity of CVB3 RNA polymerase (3Dpol). Here we measure and compare the inhibitory activity of ten amiloride analogues against CVB3 3Dpol. We show that replacement of the 3,5-diaminopyrazinyl moiety of amiloride causes loss of the inhibitory activity, whereas modifications at the 5-amino and guanidino groups increase or decrease potency. Importantly, a combination of substitutions at both the 5-amino and guanidino groups produced a compound that was more potent than its singly modified precursors. The compounds were computationally-docked into available crystal structures of CVB3 3Dpol in order to obtain a structural explanation for the activities of the analogues. To create a robust model which explained the biological activity, optimization of one of the CVB3 3Dpol crystal structures to take into account active site flexibility was necessary, together with the use of consensus docking from two different docking algorithms. This robust predictive 3D atomic model provides insights into the interactions required for inhibitor binding and provides a promising basis for the development of more potent inhibitors against this important therapeutic target.

Keywords: Coxsackievirus B3; Molecular docking; Amiloride; Drug design

Introduction

Coxsackievirus B3 (CVB3) belongs to a large family of positive-strand RNA viruses termed Picornaviridae. This family contains numerous human pathogens that cause poliomyelitis, myocarditis, meningitis, hepatitis, common cold and other diseases. Type B coxsackieviruses are responsible for 14-32% of human myocarditis cases, with CVB3 being the most common variant [1]. No antiviral treatment is currently available for any picornaviral infections.

Picornaviral genome replication is catalyzed by the viral RNA-dependent RNA polymerase 3Dpol. The three-dimensional structure of 3Dpol consists of thumb, palm and fingers domains arranged like a cupped right hand, with an active site located in the palm domain (reviewed in [2,3]). The incoming nucleotide binds directly into the active site via stacking and base pairing to a templating nucleotide, which is followed by conformational changes that reconfigure the active site for catalysis. Two Mg2+ ions, coordinated by two aspartic acid residues, are involved in the catalysis: one reduces the pKα value of the 3'-OH of the primer, allowing its deprotonation and the other orients the triphosphate of the incoming nucleotide. Viral polymerases are excellent targets for antiviral drugs, as demonstrated by the clinical success of drug treatments for human immunodeficiency virus, hepatitis B and C viruses, herpes simplex virus, human cytomegalovirus and varicella zoster virus [4].

We have previously shown that amiloride and its derivative 5-(N-ethyl-N-isopropyl)amiloride (EIPA) act as non-nucleoside inhibitors of CVB3 3Dpol [5,6]. Amiloride treatment predicted a binding site for amiloride and EIPA in 3D atomic structure of 3Dpol [5]. This site was located in close proximity to one of the Mg2+ ions and overlapped with the nucleotide binding site, thus providing a rationale for the observed competition [5]. The experimental data suggested that conformational changes are likely to occur in the process of amiloride binding to 3Dpol [5,6] and thus crystallization of such complexes may prove difficult, if not impossible. Furthermore, a mutagenesis approach to define key residues in 3Dpol that bind these inhibitors is complicated by the fact that the same residues are likely to be important in enzymatic function.

In this work we compare CVB3 3Dpol inhibitory activity of ten amiloride analogues to probe the roles of three different structural elements of amiloride: the 5-amino, guanidino and 3,5-diaminopyrazinyl groups. We show that modifications at the 5-amino and guanidino groups modulate the inhibitory activity, whereas replacement of the 3,5-diaminopyrazinyl moiety causes a complete loss with the incoming nucleotide and Mg2+ [5,6]. Computational docking predicted a binding site for amiloride and EIPA in 3D atomic structure of 3Dpol [5]. This site was located in close proximity to one of the Mg2+ ions and overlapped with the nucleotide binding site, thus providing a rationale for the observed competition [5]. The experimental data suggested that conformational changes are likely to occur in the process of amiloride binding to 3Dpol [5,6] and thus crystallization of such complexes may prove difficult, if not impossible. Furthermore, a mutagenesis approach to define key residues in 3Dpol that bind these inhibitors is complicated by the fact that the same residues are likely to be important in enzymatic function.

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of activity. We optimized one of the available 3D crystal structures to allow for active site flexibility and used two alternate docking algorithms to obtain a molecular explanation for the structure-activity relationship seen with the ten amiloride analogues. The resultant model can differentiate between active and inactive amiloride analogues.

Materials and Methods

Amiloride analogues

Amiloride, EIPA, 5-[(N,N-Dimethyl)amiloride (DMA), 5-[(N,N-Hexamethylene)amiloride (HMA), benzamil and 3,4'-dichloroben zamil (DCB) were purchased from Sigma. N-[(5-Chloro-2-hydroxybenzoyl) guanidine (CHG), N-[(5-Chloro-2-hydroxybenzoyl))-N-methyl guanidine (CHMG) and N-(5-chloro-2-methoxybenzoyl) guanidine (CMG) were purchased from Specs (Netherlands). N-Methyl-benzamil (NMB) and methyl 3-amino-5-hexamethyleneimino-6-chloropyrazinoate were synthesized as described in [7]. 5-(N,N-Hexamethylenem) benzamil (HMB) was synthesized from methyl 3-amino-5-hexamethyleneimino-6-chloropyrazinoate as follows:

Benzyl guanidine hydrochloride (1.57 g, 8.49 mmol) was added to a solution of sodium (0.17 g, 7.3 mmol) in dry isopropanol (5 mL) and the suspension was stirred for 10 min under N2. Methyl 3-amino-5-hexamethylenimino-6-chloropyrazinoate (0.484 g, 1.70 mmol) was added to the suspension and the resulting mixture was refluxed for 15 min. The mixture was cooled and water was added (25 mL). The supernatant was decanted, and the brown residue that remained was washed with water (25 mL). The residue was recrystallized from isopropanol to afford HMB as pale brown crystals (0.219 g, 32% yield) [8]. 1H NMR (400 MHz, d6-DMSO): δ 26.16 (hexamethylene C3); 28.07 (hexamethylene C2); 43.86 (benzyl CH2); 50.15 (hexamethylene C1); 117.61, 120.06 (benzyl ring); 127.11 (benzyl C4); 127.25 (benzyl C3); 128.43 (benzyl C2); 137.94 (benzyl C1); 151.05, 152.81 (pyrazine ring); 161.20 (C=O); 173.39 (guanidine C=N). HRMS (ESI+) calculated for C19H25ClN7O [M + H]+: 402.1800 (observed: 402.1799 C19H25ClN7O)

CVB3 3Dpol model preparation: CVB3 3Dpol crystal structures (PDB codes: 3CDW [8], 3DDK [9] and 3DDK [9]) were downloaded from the RCSB Protein Data Bank (http://www.pdb.org/), [10] and analyzed for docking suitability (i.e. model completeness, resolution, visual analysis for conformational changes) using Pymol (The PyMOL Molecular Graphics System, Schrödinger, LLC). All structures were aligned by their alpha carbons and for each structure, water and non-physiological ligands (i.e. detergents, salts etc.) were removed. The positions of NTP and Mg2+ ions were determined based on superposition of the structures with a closely related poliovirus 3Dpol structures as described previously [5]. Each model was then minimized sequentially (hydrogens, side-chains, backbone) under the Tripos force field for 10,000 iterations. The ligand was then removed and the optimized protein structure used to form a docking promoter for DCB as described above.

Docking of DCB proved particularly problematic due to steric clashes in all the models. This problem was overcome by firstly taking the docking result of the well behaved benzamil (into 3CDU), adding chloride ions to the inhibitor to mimic DCB, and minimizing the resultant complex sequentially (addition of hydrogens, side-chain atoms, backbone atoms) under MMFFs for 10,000 iterations. The ligand was then removed and the optimized protein structure used to form a docking promoter for DCB as described above.

The conformations of the conformationally-variable Lys61 and Lys172 residues were selected from the 3CDU [8] and 3DDK [9] crystal structures. These residues were selected as the HOT area (i.e. the area of greatest interest) for a subset minimization in Pymol (Certara L.P, St. Louis, MO, USA), using the Tripos force field [11] until consensus solutions were found.

Ligand docking – Surflex: The two tautomers of each ligand, along with their protonated forms, were minimized under the Tripos force field in Pymol (Certara L.P, St. Louis, MO, USA). Docking promoters were constructed for each of the alternate models of CVB3 3Dpol in Surflex-Dock (Certara L.P, St. Louis, MO, USA). Each promotor was constructed by selecting a 5Å sphere that encompassed active site residues Ala57, Ile58, Ser60, Lys61, Lys172, Ser173, Arg174, Asp238, Ser289 and Asp329. The bloat for the promotor was increased to 3Å (default 0) and the promotor threshold reduced to 0.27 (default 0.50). All ligands were docked into each promotor with 5 additional starting conformations per molecule, ring flexibility considered, the density of search increased to 9.0 (default 3.0), and hydrogen and heavy atom protein movement of the receptor allowed. All other parameters were kept as default. The top 20 solutions were analyzed visually in Pymol (Certara L.P, St. Louis, MO, USA) and Pymol (The PyMOL Molecular Graphics System, Schrödinger, LLC). A preferred docking solution was one which displayed a significant cluster (>50%) of docking poses in the same orientation. Where a preferred docking solution was found, the ligand-protein complex was minimized under the Tripos force-field for 10,000 iterations and analyzed visually in Pymol (The PyMOL Molecular Graphics System, Schrödinger, LLC), with all figures generated in Pymol. Furthermore, all putative polar interactions and hydrophobic interactions were calculated using the
default parameters in LigPlot + v.1.4.5 [13]. Namely, the maximum hydrogen bond distance (donor-acceptor) was 3.35Å, with the angle (donor-hydrogen-acceptor or donor-acceptor-acceptor antecedent) greater than 90°. For putative hydrophobic interactions, the maximum contact distance was 3.90Å.

Ligand docking – FRED: Using the atomic site detection in fred_receptor (OpenEye Scientific Software, Santa Fe, NM.), all potential binding pockets were selected and used to create a docking receptor for the CVB3 3Dpol crystal structure, 3CDU [8]. Omega v2.4.3 (Open Eye Scientific Software, Santa Fe, NM) was used to generate a multi-conformer structure database of all tautomers/protonated forms of the amiloride analogues. These analogues were then docked into the receptor using FRED v3.0.1 (Open Eye Scientific Software, Santa Fe, NM) with the top 20 poses requested. Docking was performed at High, Standard and Low resolution. All other parameters were kept at default. Solutions were analyzed in VIDA v4.1.1 (Open Eye Scientific Software, Santa Fe, NM).

Results/Discussion

Biological activity of amiloride analogues against CVB3 3Dpol

We examined the inhibitory activity of amiloride analogues bearing substitutions in three places: the 5-amino, guanidino and 3,5-diaminopyrazinyl groups (Figure 1). The inhibitory effects on the enzymatic activity of CVB3 3Dpol were assessed using an assay that measures the incorporation of a single nucleotide into an RNA primer-template by 3Dpol during the pre-steady-state burst phase of the reaction [5,14].

CVB3 3Dpol was pre-incubated with the RNA primer-template and with various concentrations of the analogues up to 500 M, in the presence or absence of Mg2+. Reactions were started by the addition of ATP, or ATP and Mg2+, respectively. Addition of Mg2+ together with or after the analogue was investigated because previous studies showed that the inhibition of 3Dpol by amiloride and EIPA was more pronounced when the compounds were allowed to bind to the enzyme in the absence of Mg2+ [5].

Substitutions at the 5-amino group had various effects on the inhibitory activity, with a 9-fold range in IC50 values between EIPA, amiloride, DMA and HMA (Table 1 and Figure S1). Substitutions at the guanidino group also led to variation in inhibitor potencies (Table 1 and Figure S1). Namely, compared to amiloride, DCB was more potent, benzamil less potent, and NMB was inactive. HMB, which bears a combination of substitutions at both the 5-amino and guanidino groups, was more potent than its individually modified precursors, HMA and benzamil (Table 1 and Figure S1). The replacement of the 3,5-diaminopyrazinyl moiety with 2-hydroxybenzoyl or 2-methoxybenzoyl produced inactive compounds (CHG, CHMG and CMG, Table 1).

The inhibitory effects of all active compounds were greater when they were allowed to bind to the enzyme prior to the addition of Mg2+ (Table 1 and Figure S1), suggesting overlapping binding sites and a similar mechanism of activity. The effect of Mg2+ on inhibitory potency varied between the compounds, with the difference in IC50 values for the addition of compound prior to Mg2+ versus simultaneously being 7-fold for amiloride and EIPA, 5-fold for DMA and benzamil, and

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Figure 1: Structures of amiloride analogues.
The structures of the compounds assayed against 3Dpol are shown. Although there are numerous tautomeric and protonation states, since the initial docking analysis showed little difference between them, this is the form which will be used throughout the rest of the manuscript. This figure was created using MarvinSketch (http://chemaxon.com/).
### Compound ID | IC₅₀ (µM) | Docking Results
--- | --- | ---
| | Mg²⁺ preincubation | No Mg²⁺ preincubation | % solutions in preferred cluster | C-Score | Chemgauss 3 | Number of Hydrophobic Interactions | Number of Hydrophilic Interactions |
| Amiloride | 136 ± 18 | 20 ± 1 | 100 | 7,13 | -57,04 | 4 | 7 |
| DMA | 424 ± 31 | 82 ± 29 | 82 | 7,12 | -53,55 | 6 | 7 |
| EIPA | 79 ± 10 | 11 ± 2 | 50 | 7,79 | -59,66 | 7 | 5 |
| HMA | >500 | 136 ± 22 | 70 | 6,96 | -61,22 | 9 | 3 |
| Benamil | 255 ± 20 | 52 ± 11 | 55 | 7,8 | -62,41 | 8 | 5 |
| DCB | 45 ± 1 | 15 ± 3 | 77 | 7,53 | -58,98 | 11 | 7 |
| HMB | >75 | 22 ± 4 | 70 | 7,59 | -58,48 | 10 | 6 |
| NMB | inactive | ND | NA | NA | NA | NA | NA |
| CHG | inactive | ND | NA | NA | NA | NA | NA |
| CHMG | inactive | ND | NA | NA | NA | NA | NA |
| CMG | inactive | ND | NA | NA | NA | NA | NA |

*Less than 50% inhibition at 500 µM

*No effect up to 500 µM concentration.

ND—not determined

NA—not applicable, i.e. no consistent binding poses were observed

The compounds were pre-incubated with 3Dpol and RNA primer/template at different concentrations (0 to 500 µM), either in the presence (left column) or absence (right column) of Mg²⁺, and then reactions were started by addition of ATP or ATP and Mg²⁺, respectively. The IC₅₀ values are mean ± standard error of the mean for 2-4 independent experiments and were determined as described in Materials and Methods. C-score is the consensus scoring function within Surflex (higher scores denote better docking). Chemgauss3 is the atypical scoring function within FRED and is based upon energy (lower values denote better docking).

### Table 1: Inhibitory potencies of amiloride analogues in an assay of enzymatic activity of CVB3 3Dpol and their scoring functions in Surflex-Dock and FRED algorithms.

**Ligand docking**

It is generally accepted that no single docking algorithm performs well across all protein targets [15]. Using more than one algorithm builds a consensus that increases the reliability of the results. Two alternate docking algorithms, Surflex-Dock (Certara L.P, St. Louis, MO, USA) and FRED (Open Eye Scientific Software, Santa Fe, NM), were used in this study. The Surflex-Dock algorithm uses an active site protomol as a target for ligand docking. Numerous poses of the ligand are generated and placed into the protomol and scored using the Surflex scoring functions [16]. The FRED algorithm (Open Eye Scientific Software, Santa Fe, NM) performs a systematic and exhaustive search of all possible ligand poses, which need to be generated separately using a program such as Omega (Open Eye Scientific Software, Santa Fe, NM). It then filters and optimizes the poses using the Chemgauss4 scoring function [17]. Both programs rank the compounds based upon the calculated scoring functions.

Picornaviral 3Dpol undergoes significant conformation changes during the catalytic cycle: the incoming nucleotide binds to an “open” conformation of the enzyme-RNA template complex, its binding triggers the closure of the active site to form a catalytically competent “closed” conformation, which reopens after the nucleotide addition [9,18]. One method to enrich computational docking and to accommodate targets prone to conformational changes is to use an ensemble of protein structures [19]. Three crystal structures of CVB3 3Dpol were used in docking experiments; PDB codes: 3CDW [8], 3CDU [8], and 3DDK [9] (http://www.pdb.org [10]). All structures have comparable resolution and completeness and belong to the same space groups. The only notable differences between the three are that 3CDW and 3CDU were crystallized from the Nancy strain of Coxsackievirus, whilst 3DDK was crystallized from a different strain. Also, 3CDW was co-crystallized with a VPG primer. The recently published elongated structures [20] were not included in this venture, although visual analysis suggests they would have similar docking limitations for this project as the 3DDK structure (see below). All of the amiloride analogues were initially docked into the previously-defined amiloride binding pocket [5] of all three CVB3 3Dpol structures, in the presence and absence of Mg²⁺, using Surflex-Dock in Sybylx2.0 (Certara L.P, St. Louis, MO, USA). In the presence of Mg²⁺, no consistent cluster of solutions could be obtained for any of the ligands. When Mg²⁺ was absent in the 3CDW and 3CDU crystal structures, none of the inactive compounds showed a consistent cluster of solutions, whereas six out of seven active compounds (except DCB – see below) docked and clustered in an analogous manner in the nucleotide binding site, overlapping the Mg²⁺ binding site (Figure 2). These docking solutions were almost identical to the one previously described for amiloride and EIPA [5]. All tautomers and protonated forms of these ligands docked in an analogous way, with little difference in the scoring function (C-score) value.

For the 3DDK structure, five active compounds (except DCB and DMA) showed a consistent docking solution for over 50% of the top 20 poses, but the docking orientations were almost perpendicular compared to the solutions obtained for the 3CDU/3CDW structures. A possible reason for docking differences is revealed from a comparison of the active sites in all three crystal structures: there are prominent side-chain movements of Lys172 and Lys61 between the 3CDW/3CDU and 3DDK structures. A subset energy minimization was performed on the 3DDK and 3CDU structures where the two lysine residues were allowed to move. The side-chain of Lys172 in both minimized structures moved towards the position found in 3CDU. However, the minimization of Lys61 resulted in a range of orientations, suggesting that a large amount of movement of this side-chain is possible (Figure 3). This finding is supported by high (>0.4 Å) temperature factors (a crystallographic measure of side-chain flexibility and/or disorder) for Lys61 in both the 3CDU and 3DDK crystal structures. Analysis of the electron density maps derived from the crystallography studies (calculated from the deposited structure factors in the relevant PDB entries) is consistent with at least two alternate conformers of Lys61 in the 3DDK structure, and there is poor electron density for the
side-chain in the 3CDU structure. Most crystal structures of other \textit{Picornaviridae} 3D\textsuperscript{pol} (e.g. poliovirus), including the recent elongated Coxsackievirus complexes [20] have the side-chain of Lys61 forming the NTP tunnel wall, allowing it to engage in a salt bridge with Glu177. However, although the side-chain of Lys61 in the 3DDK structure is in a similar position to that seen with other \textit{Picornaviridae} structures, there is a rotation about the Ca-C\(\beta\) bond of Glu177, orientating the side-chain so there is no salt bridge. Based on these data and the fact that a larger number of active compounds docked in an analogous manner into the 3CDU/3CDW structures when compared to the 3DDK structure, 3CDU was chosen as a preferred docking receptor.

Surprisingly, DCB did not dock in a consistent, clustered manner although it is structurally similar to benzamil. Inspection of the benzamil-3CDU complex revealed that the phenyl group spans the full extent of the binding pocket suggesting the charging atoms on the phenyl group of DCB might clash with residues Glu177 and Ser289. Previously published experimental data suggested that amiloride is a slow inhibitor, and hence it’s binding to CVB3 3D\textsuperscript{pol} is likely to involve conformational changes [5]. In poliovirus 3D\textsuperscript{pol}, the equivalent residue to Ser289 (Ser288) is important for NTP ribose recognition and active site closure, and as such it is repositioned during the conformational changes that lead to catalysis [18, 21]. This residue has been observed in multiple conformations, in various crystal structures of \textit{Picornaviridae}.

To allow for the flexibility of Ser289 in the model, the two chlorine atoms were added to the phenyl group of benzamil in the benzamil-3CDU complex, and the resulting model was minimized. The ligand was then removed and the optimized structure was used as a docking receptor for all amiloride analogues. Each ligand bound complex was allowed to minimize for 10,000 iterations, allowing for further movement of the side-chains of the residues.

All active compounds docked to the optimized CDU structure so that they have a hydrogen bond between a nitrogen atom of the guanidino group and the backbone of Ile58, and another bond between the 3-amino group and the backbone of Leu175 (Figure 4 and Figure S2). Additional interactions for each compound are shown in Figure 3 and detailed in Table S1. Although the docking model solutions are similar to those described previously for amiloride and EIPA [5], some of the predicted interactions are now different in the optimized model. For example, the two hydrogen bonds to the backbone of Ile58 were previously reported as two hydrogen bonds to Ala57 and Ser60 [5]. Since the optimized model described here differentiates between the active and inactive amiloride analogues, we believe the new approach supersedes the previous approach and the accompanying conclusions.

Analysis of the top 20 ranked poses for each compound revealed that DCB and benzamil had a small cluster (3 out of 20) of solutions in an alternate binding mode. To analyze the potential of alternate binding modes and/or binding sites not identified by the Surflex-Dock program (Certara L.P, St. Louis, MO, USA), we docked all compounds into the refined 3CDU structure using the OE Docking suite, which includes FREDv3.0.1 (Open Eye Scientific Software, Santa Fe, NM). The results for all three FRED resolutions were identical to those described above, with no consistent cluster of binding poses observed for inactive compounds, and all of the active compounds docking to overlap the nucleotide binding site, consistent with the results obtained using Surflex-Dock. The consensus between the two docking algorithms was consistent with the predicted binding mode.

Numerous studies have examined the ability of different docking programs to predict ligand affinity to an extensive array of protein targets, with the consensus being that docking is suited to identify activity, but does not effectively rank affinities [15]. Additionally, the narrow range of potencies (differences in IC\(_50\) values of less than 10-fold) for the active compounds here precluded a meaningful correlation between the docking scores and the potencies of the active compounds (Table 1).

The docking models developed herein provide a rationale for most of the biological data. Strikingly, none of the inactive analogues

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**Figure 2: Docking of amiloride analogues.**

Our optimized CVB3 3D\textsuperscript{pol} structure (based upon PDB code: 3CDU, [8]) is shown as grey cartoon. The blue spheres denoting amiloride, show a partial overlap in the binding site to the previously reported [5] position of ATP (red spheres), and displays a distinct overlap with the previously determined position of the Mg\(^{2+}\) atoms (purple spheres, PDB code: 3OL7, [18]). The position of the RNA template (orange cartoon, PDB code: 3OL7, [18], not used in docking experiments) is also shown.

**Figure 3: Energy minimization of Lys61 and Lys172.**

Displayed is the residues within 4Å of Lys 61 and Lys172 (highlighted with sticks) from 3DDK (magenta carbons) and 3CDU (yellow carbons) plus their minimized versions shown in cyan carbons and green carbons, respectively. There is significant movement between the structures for many of the residues in this region however, only the wide variety of possible side-chain orientations for Lys61 and Lys172 seemed to significantly impact the docking.
were able to dock in a consistent manner, which can be rationalized by considering the docking orientation. For example, the additional methyl group in NMB would clash with the backbone of Ile58 and remove what appears to be an important hydrogen bond. For CHG, CHMG, and CMG, many hydrogen bonding interactions are lost with the exchange of the nitrogen atoms to carbons in the pyrazine ring and the loss of 3,5-amino groups. These interactions are obviously essential for the CVB3 3Dpol inhibitory activity of this compound class. Furthermore, the boundaries of the binding pocket have been probed with larger active compounds such as DCB and HMB. Also, the 5-ethylamino group of EIPA, when compared to the 5-methylamino group of DMA, allows for extra hydrophobic interactions with Lys159 and Ile176 (Figure 4b and 4c).

In conclusion, we report an initial structure-activity relationship for amiloride analogues that reveals that the 5-amino and guanidino groups of amiloride individually modulate potency of CVB3 3Dpol

Figure 4: Inhibitor docking to 3Dpol.
All active compounds interact in the same general orientation in the nucleotide binding site. The residues involved in significant interactions with inhibitors are shown (grey sticks) with polar interactions indicated as yellow dashed lines. The schematic 2D representations shown in the right column were created by LigPlot* v1.4.2 [13] with hydrophobic interactions shown as red semi-circles with lines radiating out in the direction of the interaction. (a) Amiloride (magenta carbons); (b) DMA (orange carbons); (c) EIPA (cyan carbons); (d) HMA (green carbons); (e) Benzamid (yellow carbons); (f) DCB (peach carbons); and (g) HMB (purple carbons).
inhibition and in combination can have an additive effect. We have investigated several approaches to molecular docking and have shown that the choice of the initial protein model plays a significant role in the ability of two alternate docking programs to find consistent solutions. The optimized docking model described here provides insight into the interactions required for inhibitor binding and allows the distinguishing of active versus inactive compounds. Two interactions have been identified that appear to be critical for inhibitory activity: a hydrogen bond between a nitrogen atom of the guanidino group of the active inhibitor and the backbone of Ile58, and a hydrogen bond between the 3-amino group of the inhibitor and the backbone of Leu175. The optimized molecular model can now be used to design more potent amiloride analogues to explore the structure-activity relationships of this compound class and facilitate the development of drugs for the treatment of CVB3 infections.

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