

Article

Inhibition of Aminoglycoside 6'-N-Acetyltransferase Type Ib [AAC(6')-Ib]: Structure-Activity Relationship of Substituted Pyrrolidine Pentamine Derivatives as Inhibitors

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Abstract: The aminoglycoside 6'-N-acetyltransferase type Ib [AAC(6')-Ib] is a common cause of resistance to amikacin and other aminoglycosides in Gram-negatives. Utilization of mixture-based combinatorial libraries and application of the positional scanning strategy identified an inhibitor of AAC(6')-Ib. This inhibitor's chemical structure consists of a pyrrolidine pentamine scaffold substituted at four locations (R1, R3, R4, and R5). The substituents are two S-phenyl (R1 and R4), an S-hydroxymethyl (R3), and a 3-phenylbutyl (R5) groups. Another location, R2, does not have a substitution, but it is named because its stereochemistry was modified in some compounds utilized in this study. Structure-activity relationship (SAR) analysis using derivatives with different functionalities, modified stereochemistry, and truncations were carried out by assessing the effect of the addition of each compound at 8 μ M to 16 μ g/ml amikacin-containing media and performing checkerboard assays varying the concentrations of the inhibitor analogs and the antibiotic. The results showed that: 1) the aromatic functionalities at R1 and R4 are essential, but the stereochemistry is essential only at R4, 2) the stereochemical conformation at R2 is critical, 3) the hydroxyl moiety at R3 as well as stereoconformation are required for full inhibitory activity, 4) the phenyl functionality at R5 is not essential and can be replaced by aliphatic groups, 5) the location of the phenyl group on the butyl carbon chain at R5 is not essential, 6) the length of the aliphatic chain at R5 is not critical, 7) all truncations of the scaffold resulted in inactive compounds. Molecular docking revealed that all compounds preferentially bind to the kanamycin C binding cavity, and binding affinity correlates with the experimental data for most of the compounds evaluated. The SAR results in this study will serve as the basis for the design of new analogs in an effort to improve their ability to induce phenotypic conversion to susceptibility in amikacin-resistant pathogens.

Keywords: aminoglycoside resistance; structure-activity relationship; aminoglycoside modifying enzymes, acetyltransferase, *Acinetobacter*

1. Introduction

A growing number of Gram-negative pathogens are rapidly acquiring resistance to most and, in some cases, all antibiotics in use [1]. As a consequence, treatment of severe infections caused by multidrug-resistant (MDR) bacteria is becoming more complicated and prohibitively expensive [2]. The magnitude of the problem is illustrated by the inclusion of MDR *Acinetobacter baumannii* and other Gram negatives such as *Klebsiella pneumoniae* and *Pseudomonas aeruginosa* as "Priority 1:Critical" in the World Health Organization Priority Pathogens list for Research and Development of new antibiotics [3]. The urgency to develop new treatments against these pathogens requires not only the design of novel antibiotics but also the finding of adjuvants that, in combination with

existing drugs, circumvent the resistance [4]. This latter strategy extends the useful life of antibiotics already in use, but that are becoming ineffective due to the dissemination of resistance traits. This strategy has been successful for β -lactams, in which case several β -lactam/ β -lactamase inhibitor formulations are currently in use [5,6]. On the other hand, the identification or design of inhibitors of resistance to other classes of antibiotics has not progressed beyond the research laboratory.

Aminoglycosides are bactericidal antibiotics that interfere with translational fidelity producing proteins with incorrect primary sequences that lead to a myriad of toxic physiological effects and, ultimately, cell death [7,8]. These antibiotics have been instrumental in treating life-threatening infections caused by Gram-negative and, in combination with other antimicrobials, Gram-positive bacteria [7,9]. Although bacteria have developed various mechanisms to resist aminoglycosides, enzymatic inactivation is the most prevalent in the clinical setting [8,10,11]. There are numerous reports of compounds that interfere with the inactivation of the antibiotic molecule by different molecular mechanisms or enhance the cellular uptake [8,12-23]. However, despite their demonstrated activity, none of them could be turned into formulations for clinical use.

The aminoglycoside 6'-N-acetyltransferase type Ib [AAC(6')-Ib] causes resistance to amikacin and other aminoglycosides in Gram-negative bacteria [11]. Since this is the most common enzyme among AAC(6')-I-producing Gram-negative pathogens, it was selected as the target in the quest for inhibitors that, in combination with amikacin, could be used to treat resistant infections. In particular, it could help control those caused by strains resistant to carbapenems, which are antimicrobials of last resort for treatment of several MDR infections [6]. We have recently identified an inhibitor of AAC(6')-Ib using mixture-based combinatorial libraries and the positional scanning strategy [23]. The compound consists of a pyrrolidine pentamine scaffold with two S-phenyl, an S-hydroxymethyl, and a 3-phenylbutyl groups at the positions shown in Table 1. The structure-activity relationship (SAR) study described in this article was carried out to better understand this compound's properties as an inhibitor of AAC(6')-Ib and design related compounds with more robust activity.

2. Materials and Methods

2.1. Bacterial strains

A. baumannii A155, originally isolated from a urinary sample at a hospital in Buenos Aires, Argentina [24]. It belongs to the clonal complex 109, it is multiple drug resistant and naturally carries *aac(6')-Ib* [25,26].

2.2. General methods

Routine cultures were carried out in Lennox L broth (1% tryptone, 0.5% yeast extract, 0.5% NaCl), and 2% agar was added in the case of solid medium. For determination of levels of resistance to amikacin the culture medium used was Mueller-Hinton broth.

Checkerboard assays were performed in Mueller-Hinton broth with variable concentrations of the compound to be tested (0, 4, 8, 16, and 24 μ M) and amikacin (0, 8, 16, 32, and 64 μ g/ml) in microtiter plates using the BioTek Synergy 5 microplate reader (BioTek Synergy 5) as described before [23]. All compounds that did not show a significant reduction ($p < 0.01$, Two-Sample T-Test versus compound 2637.001) in the initial screening were chosen for checkerboard assay. Since there is a chance that the testing compounds have some residual antimicrobial activity, data was analyzed using an approach that quantifies exact levels of synergy [23,27]. The model considers that amikacin and the compounds to be tested have independent antimicrobial mechanisms of action. The percent activity of the mixture of the two chemicals is modeled as:

$$\%_{\text{amikacin \& compound}}(x_1, x_2) = \%_{\text{amikacin}}(x_1) + \%_{\text{compound}}(x_2) - \%_{\text{amikacin}}(x_1) \cdot \%_{\text{compound}}(x_2)$$

In this equation x_1 and x_2 are the concentrations of amikacin and tested compound, respectively. To calculate the effective percent activity of the antibiotic alone at a given concentration, after accounting for compound activity the previous equation can be rearranged as follows:

$$\text{Eff\%amikacin}(x_1) = (\%_{\text{amikacin \& compound}}(x_1, x_2) - \%_{\text{compound}}(x_2)) / (1 - \%_{\text{compound}}(x_2))$$

This methodology informs the actual change in amikacin resistance levels. Four checkerboard assays were performed for each compound, and the above methodology was applied to the median of the four values at each dose combination.

Once applied to the checkerboard data, a 95% confidence interval for the mean effective concentration of amikacin to achieve 50% inhibition (IC_{50}) at each dose of potentiating compound was determined using standard curve fitting of Hill's equation.

2.3. *Synthesis and purification of small molecule compounds*

All molecules screened were synthesized at Torrey Pines Institute for Molecular Studies (now the Center for Translational Science at Florida International University) using solid-phase chemistry as previously described [23].

2.4. *High-performance liquid chromatography (HPLC) purification*

All purifications were performed on a Shimadzu Prominence preparative HPLC system, consisting of LC-8A binary solvent pumps, an SCL-10A system controller, a SIL-10AP auto sampler, and an FRC-10A fraction collector. A Shimadzu SPD-20A UV detector set to 214 nm was used for detection. Chromatographic separations were obtained using a Phenomenex Gemini C18 preparative column (5 μm , 150 mm \times 21.5 mm i.d.) with a Phenomenex C18 column guard (5 μm , 15mm \times 21.2mm i.d.). Prominence prep software was used to set all detection and collection parameters. The mobile phases for HPLC purification were HPLC grade obtained from Sigma-Aldrich and Fisher Scientific. The mobile phase consisted of a mixture of acetonitrile/water (both with 50mM acetic acid). The initial setting for separation was 2% acetonitrile, which was held for 2 min, then the gradient was linearly increased to 6% acetonitrile over 4 min. The gradient was then linearly increased to 35% acetonitrile over 29 min. The HPLC system was set to automatically flush and re-equilibrate the column after each run for a total of four column volumes. The total flow rate was set to 15 mL/min, and the total injection volume was set to 2000 μL . The fractions corresponding to the desired product were then combined and lyophilized.

2.5. *Liquid chromatography–mass spectrometry (LCMS) analysis of purified material*

Purity and identity of compounds were verified using a Shimadzu 2010 LCMS system consisting of a LC-20AD binary solvent pump, a DGU-20A degasser unit, a CTO-20A column oven, and a SIL-20A HT auto sampler. A Shimadzu SPD-M20A diode array detector scanned the spectrum range of 190 – 400 nm during the analysis. Chromatographic separations were obtained using a Phenomenex Luna C18 analytical column (5 μm , 150 mm \times 4.6 mm i.d.) with a Phenomenex C18 column guard (5 μm , 4 \times 3.0 mm i.d.). All equipment was controlled and integrated by Shimadzu LCMS solutions software version 3. Mobile phase A for LCMS analysis was LCMS grade water, and mobile phase B was LCMS grade acetonitrile obtained from Sigma-Aldrich and Fisher Scientific (both with 0.1% formic acid for a pH of 2.7). The initial setting for analysis was 5% acetonitrile (v/v), and then linearly increased to 95% acetonitrile over 14 min. The gradient was then held at 95% acetonitrile for 2 min before being linearly decreased to 5% over 2 min and held until stop for an additional 2 min. The total run time was 20 min, and the total flow rate was 0.5 mL/min. The column oven and flow cell temperature for the diode array detector was 40 °C. The auto sampler was at room temperature, and a 5 μL aliquot was injected for analysis.

2.6. Modeling

The structures of the compounds were converted to 3D structures with added polar hydrogen bonds using Open Babel [28]. The structure of AAC(6')-Ib complexed with kanamycin C and AcetylCoA [29] was obtained from the protein data bank. The AAC(6')-Ib protein with kanamycin C removed, as well as the compounds were prepared in the pdqt format using AutoDockTools 4.2 [30]. A cavity in the kanamycin C binding region of the protein was selected as the target site for virtual screening. Vina from AutoDockTools 4.2 [30] was used to perform docking and screening. The docking scores were sorted and ranked based on their predicted binding energies. LigPlot+ [31] was used to generate a 2-D ligand-protein interaction map. PyMol 2.3 (Schrodinger) was used for visualization and rendering.

3. Results

3.1. Synthesis and preliminary analysis of analogs to compound 2637.001

The recent identification of an inhibitor of the AAC(6')-Ib opened new possibilities to formulate combinations with aminoglycosides to treat resistant infections. This compound's chemical structure consists of a pyrrolidine pentamine scaffold substituted with two S-phenyls, an S-hydroxymethyl, and a 3-phenylbutyl groups at the positions R1, R3, R4, and R5, respectively (compound **2637.001**, Table 1). (Note **2637.001** was referred to as compound **2155-206** in the previous publication [23]). A SAR set of experiments with a series of compound **2637.001** analogs was initiated to gain insights into the different chemical groups' contribution to the AAC(6')-Ib inhibitory effect. The primary goal of this preliminary SAR study was to assess the relative importance of each specific functionality and stereochemistry as well as determining the minimal pharmacophore needed. Therefore, compounds were designed with a single substitution at each of the R positions or truncation of a specific scaffold fragment (Table 1). The effect of the addition of each compound at 8 μ M concentration to 16 μ g/ml amikacin-containing medium on growth of the *aac(6')-Ib*-harboring *A. baumannii* A155 strain was tested. The concentration of amikacin was chosen based on previous studies showing that this strain grows in the breakpoint concentration 16 μ g/ml amikacin [32]. Bacterial growth was assessed measuring OD₆₀₀ after 20 h incubation, and the values were used to calculate the percentage of inhibition of resistance (Table 1). The different degrees of growth inhibition observed in these assays indicate that the structural changes of the analogs with respect to the compound **2637.001** must affect the AAC(6')-Ib inhibitory efficacy.

Table 1. Properties of 2637.001 analogs

Compound name	Chemical structure	Functionalities	%Inhibition (Average, n=10)	Standard Error	Delta G Kcal/ml (Average, n=3)
2637.001		R1: <i>S</i> -phenyl R2: <i>S</i> -pyrrolidine R3: <i>S</i> -hydroxymethyl R4: <i>S</i> -phenyl R5: 3-phenylbutyl	62	4	-9.5±0.1
R1 analogs					
2637.002		R1: <i>S</i> -methyl R2: <i>S</i> -pyrrolidine R3: <i>S</i> -hydroxymethyl R4: <i>S</i> -phenyl R5: 3-phenylbutyl	18*	2	-8.7±0.1
2637.003		R1: <i>S</i> -benzyl R2: <i>S</i> -pyrrolidine R3: <i>S</i> -hydroxymethyl R4: <i>S</i> -phenyl R5: 3-phenylbutyl	20*	3	-8.0±0.1
2637.020		R1: <i>R</i> -phenyl R2: <i>S</i> -pyrrolidine R3: <i>S</i> -hydroxymethyl R4: <i>S</i> -phenyl R5: 3-phenylbutyl	73	4	-8.5±0.3
R2 analogs					
2637.021		R1: <i>S</i> -phenyl R2: <i>R</i> -pyrrolidine R3: <i>S</i> -hydroxymethyl R4: <i>S</i> -phenyl R5: 3-phenylbutyl	28*	4	-9.5±0.3
R3 analogs					
2637.004		R1: <i>S</i> -phenyl R2: <i>S</i> -pyrrolidine R3: <i>S</i> -methyl R4: <i>S</i> -phenyl R5: 3-phenylbutyl	39	6	-9.2±0.1

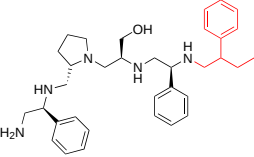
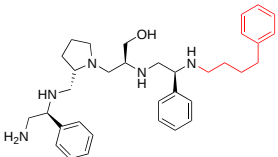
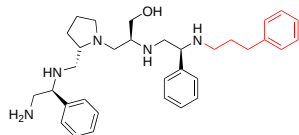
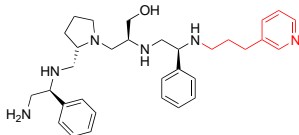
2637.005		R1: <i>S</i> -phenyl R2: <i>S</i> -pyrrolidine R3: <i>R</i> - hydroxymethyl R4: <i>S</i> -phenyl R5: 3-phenylbutyl	24*	2	-8.2±0.1
2637.019		R1: <i>S</i> -phenyl R2: <i>S</i> -pyrrolidine R3: <i>R</i> - methyl R4: <i>S</i> -phenyl R5: 3-phenylbutyl	74	9	-9.2±0.1

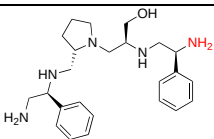
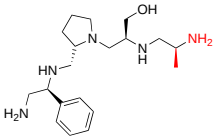
R4 analogs

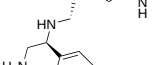
2637.006		R1: <i>S</i> -phenyl R2: <i>S</i> -pyrrolidine R3: <i>S</i> - hydroxymethyl R4: <i>S</i> - methyl R5: 3-phenylbutyl	23*	2	-7.9±0.2
2637.022		R1: <i>S</i> -phenyl R2: <i>S</i> -pyrrolidine R3: <i>S</i> - hydroxymethyl R4: <i>R</i> - phenyl R5: 3-phenylbutyl	28*	4	-9.6±0.1

R5 analogs

2637.007		R1: <i>S</i> -phenyl R2: <i>S</i> -pyrrolidine R3: <i>S</i> - hydroxymethyl R4: <i>S</i> -phenyl R5: ethyl	60	1	-9.5±0.1
2637.008		R1: <i>S</i> -phenyl R2: <i>S</i> -pyrrolidine R3: <i>S</i> - hydroxymethyl R4: <i>S</i> -phenyl R5: butyl	17*	2	-9.4±0.1
2637.010		R1: <i>S</i> -phenyl R2: <i>S</i> -pyrrolidine R3: <i>S</i> - hydroxymethyl R4: <i>S</i> -phenyl R5: pentyl	71	2	-9.1±0.2

2637.012		R1: <i>S</i> -phenyl R2: <i>S</i> -pyrrolidine R3: <i>S</i> -hydroxymethyl R4: <i>S</i> -phenyl R5: 2-phenylbutyl	40	6	-8.5±0.2
2637.011		R1: <i>S</i> -phenyl R2: <i>S</i> -pyrrolidine R3: <i>S</i> -hydroxymethyl R4: <i>S</i> -phenyl R5: phenylbutyl	66	6	-8.1±0.2
2637.013		R1: <i>S</i> -phenyl R2: <i>S</i> -pyrrolidine R3: <i>S</i> -hydroxymethyl R4: <i>S</i> -phenyl R5: phenylpropyl	62	3	-9.1±0.1
2637.014		R1: <i>S</i> -phenyl R2: <i>S</i> -pyrrolidine R3: <i>S</i> -hydroxymethyl R4: <i>S</i> -phenyl R5: (pyridiin-3-yl)propyl	46	3	-9.1±0.2

Truncation Analogs					
2637.015		R1: <i>S</i> -phenyl R2: <i>S</i> -pyrrolidine R3: <i>S</i> -hydroxymethyl R4: <i>S</i> -phenyl R5: hydrogen	26*	3	-8.7±0.1
2637.016		R1: <i>S</i> -phenyl R2: <i>S</i> -pyrrolidine R3: <i>S</i> -hydroxymethyl R4: <i>S</i>-methyl R5: hydrogen	20*	3	-8.2±0.1

2637.017	 R1: S-phenyl R2: S-pyrrolidine R3: S-hydroxymethyl R4: modified to ethyl R5: nothing	21*	3	-8.4±0.1
2637.018	 R1: S-phenyl R2: S-pyrrolidine R3: S-hydroxymethyl R4: nothing R5: nothing	17*	2	-8.2±0.1

*P-Value < 0.01, Two-Sample T-Test with Bonferroni-Holm Correction when compared to 2637.001. Delta G Kcal/ml values are the binding scores obtained from molecular docking and reported as averages of the top three docking scores. For comparison, Amikacin docking gave Delta G of -8.2±0.2.

The importance of the *S*-phenyl at the R1 position was assessed by modifying the chemical group or the stereochemistry (Table 1). In compound **2637.002**, the aromatic phenyl group was removed leaving an *S*-methyl functionality, and in compound **2637.003**, the phenyl moiety was separated from the backbone by the addition of a methylene group (Table 1). In both cases, the *S* conformation was maintained. In compound **2637.020**, the *S*-phenyl was replaced by an *R*-phenyl functionality changing only the stereochemistry. Table 1 shows that replacing the aromatic functionality with a methyl group significantly reduced the percentage of inhibition (hereafter referred to as inhibitory activity) with respect to that observed when compound **2637.001** is tested (62% vs. 18%). Interestingly a methylene group placed between the scaffold and the phenyl functionality (benzyl) (**2637.003**) also affected the inhibitory activity reducing it to 20%. This reduction in inhibitory activity could be due to the loss of the aromatic group's ability to interact or stabilize the interaction with the appropriate region of AAC(6')-Ib. Conversely, the absolute stereochemistry at this position does not appear to be critical as the *S* and *R* conformations produced similar inhibitory activities (**2637.001**, 62% vs. **2637.020**, 73%).

Position R2 was not originally considered a location for addition of functionalities. However, in this study the relative importance of the stereochemistry at this position was assessed (Table 1). An analog, **2637.021**, was synthesized, where the R2 stereocenter was modified from *S* to *R*. This change resulted in a compound with significantly reduced capability to inhibit resistance to amikacin (**2637.001**, 62% vs. **2637.021**, 28%). This result demonstrated that absolute stereochemistry plays a crucial role at this position.

At the R3 position, which has an *S*-hydroxymethyl in **2637.001**, analogs that modify the functionality or the stereochemistry were assessed (Table 1). Compound **2637.005** differs from **2637.001** in the stereoconfiguration, which was altered from *S* to *R*. This compound was used to determine the relative importance of the absolute conformation at this position. Table 1 shows that the modification significantly impacted the inhibitory activity (**2637.001**, 62% vs. **2637.005**, 24%), probably by impeding the appropriate interaction between the hydroxy moiety and the target.

The two other analogs with modifications at the R3 positions were compounds **2637.004** and **2637.019**, in which the hydroxy group was eliminated (Table 1). The conformation was maintained in the former and changed to *R* in the latter. It was interesting that when the stereochemistry of the parent compound was preserved, the inhibitory activity was slightly reduced (**2637.001**, 62% vs. **2637.004**, 39%), suggesting that the hydroxyl group is needed and likely is involved in hydrogen bonding between the parent compound **2637.001** and AAC(6')-Ib. However, when the stereochemistry was changed, the

compound showed a comparable capacity for inhibiting growth to the parent compound (**2637.019**, 74% vs. **2637.001**, 62%). This result suggests a different binding motif for this analog that does not require the hydroxyl group present in the parent compound.

The *S*-phenyl group at the R4 position in the compound **2637.001** was replaced by *S*-methyl (**2637.006**) or *R*-phenyl (**2637.022**) groups (Table 1). The results obtained when adding compound **2367.006** or **2637.022** to the culture medium showed the importance of the aromatic functionality and the *S* conformation at the R4 position respectively. The reduction in levels of inhibition of resistance indicates that both modifications had significant effects suggesting an important role of the phenyl moiety and its steric configuration.

Modifications at the R5 position resulted in three groups of analogs (Table 1). The first set was designed to examine the effect of removing the aromatic phenyl group and replacing it with aliphatic groups with various carbon chain lengths. In compound **2637.008**, the phenyl group has been removed from the 3-position of the butyl group. The phenyl group was removed in compounds **2637.007** and **2637.010**, and the aliphatic chains were modified to contain either two or five carbons, respectively. The compounds **2637.007** and **2637.010**, in which the phenyl group was removed, and the carbon chain length was reduced or lengthened with respect to compound **2637.001**, exhibited similar levels of inhibition of resistance to the parent compound (**2637.001** 62%, **2637.007** 60%, and **2637.010** 71%). These results suggest that the phenyl functionality is not essential. In the case that an aliphatic functionality is used at this position, there are potential options regarding sizing, branching, and additional substituents, thus allowing for lipophilic optimization as needed.

The second set of analogs is characterized by modifications in the location of the phenyl group on the butyl carbon chain (Table 1). In compounds **2637.011** and **2637.012**, the phenyl group is bound to the second or the fourth carbon, respectively. Analog **2637.011** produced a similar inhibitory effect compared to the parent compound (**2637.001**, 62% vs. **2637.011**, 66%). Having comparable activities, analog **2637.011** presents the added benefit that by placing the substituent at position 4, the undefined stereocenter on the parent compound is eliminated. The result obtained with analog **2637.012** had lower but still evident activity (40%), suggesting that the phenyl group located at the R5 position can be moved without a drastic loss of activity.

The last set includes compounds **2637.013** and **2637.014** (Table 1). Compound **2637.013** maintains the phenyl group on the terminal carbon, but it is bound to a shorter aliphatic carbon chain (propyl). This conformation examines the effects of eliminating the last carbon moiety at the R5 position of the parent compound and provides an analog without a stereocenter in this position. It was encouraging to note that this analog also maintained activity (62%) as it allows for another compound where the undefined stereochemistry is eliminated at the R5 position. Compound **2637.014** builds on **2637.013** by examining the effect of introducing an aromatic heterocyclic moiety at the R5 position. While this compound had lower activity (46%), it could suggest that a heterocyclic moiety can be introduced at this position.

The last set of compounds is composed of truncated analogs (Table 1). This set was utilized to assess the minimal pharmacophore needed to preserve inhibitory activity when scanning from the R1 to the R5 direction. In compounds **2637.015** and **2637.016**, the R5 functionality was eliminated, and compound **2637.016** was reduced further by removing the phenyl group of the R4 functionality. Compound **2637.017** was further reduced, eliminating the primary amine and *S*-methyl groups. Finally, compound **2637.018** was designed to lack both the R4 and R5 groups from the parent compound. None of the analogs from this set produced significant inhibitory activity in the primary assay suggesting that the entire scaffold is essential. However, future studies will be necessary to confirm the essentiality of other scaffold regions.

3.2. Molecular Docking

The data from the SAR study suggests that specific changes in absolute stereochemistry or elimination of key functional groups can affect the compound's ability to enhance the amikacin antibacterial response in the primary screening assay (Table 1). The binding poses of the twenty compounds in Table 1 as well as amikacin against AAC(6')-Ib were investigated to explore potential critical interactions responsible for these changes in inhibitory efficacy. A blind docking revealed that the compounds all preferentially bind to the kanamycin C binding cavity, and therefore, this site is considered as the target site for docking. To incorporate the flexibility of the sidechains around the target site, flexible docking was performed with W49, Y65, E73, V75, Q91, Y93, S98, D100, W103, D115, D152, and D179 as the flexible residues. The screening revealed that **2637.001** is one of the top compounds to bind AAC(6')-Ib effectively, and based on the Delta G value obtained from docking, it is predicted to bind more effectively than amikacin (Table 1). Figure 1a shows the AAC(6')-Ib-compound complex, showing the binding pose of **2637.001** in the kanamycin C binding site. This is the same binding site predicted for amikacin (Figure S1). A 2D map of the ligand in the binding site shows that Q91 and D179 make hydrogen-bond interactions, in addition to other residues involved in hydrophobic interactions (Figure 1b).

Some correlations comparing the predicted binding efficacies and poses to the screening data and SAR observations were found (Table 1, Delta G, and Figures 1-4). For example, looking in detail at the R3 position analogs (**2637.001**, **2637.004**, **2637.005**, **2637.019**), some trends from the docking study support the SAR observations noted previously. Compound **2637.004** eliminates the hydroxyl group from the parent compound, which is shown to hydrogen bond with the target protein (Figure 1). Compound **2637.004** is predicted to have a slightly lower binding efficiency than **2637.001** (Table 1), and the binding pose (Figure 2, a and b) shows that it now only interacts with the Asp179 residue thus providing a rationale for the reduction in inhibitory activity noted for **2637.004** (Table 1). In compound **2637.005** the absolute stereochemistry of the hydroxy group is changed. This modification significantly affected the inhibitory activity and predicted binding efficacy (Table 1). The 2D map (Figure 3, a and b) shows that the compound has a preferential reorientation in the binding pocket so that the hydroxy group no longer interacts with the target protein. In compound **2637.019** the hydroxy group at the R3 position was eliminated and the absolute stereochemistry at this position was modified. This compound maintained inhibitory activity (Table 1, % inhibition) and was predicted to have slightly less binding efficiency than the parent compound (Table 1, Delta G). Looking at the 2D map (Figure 4, a and b), it appears that **2637.019** potentially compensates for the loss of the hydroxyl group by maintaining a hydrogen bond interaction with Asp115 as well as increasing an intramolecular pi stacking between the R1 and R5 benzyl groups.

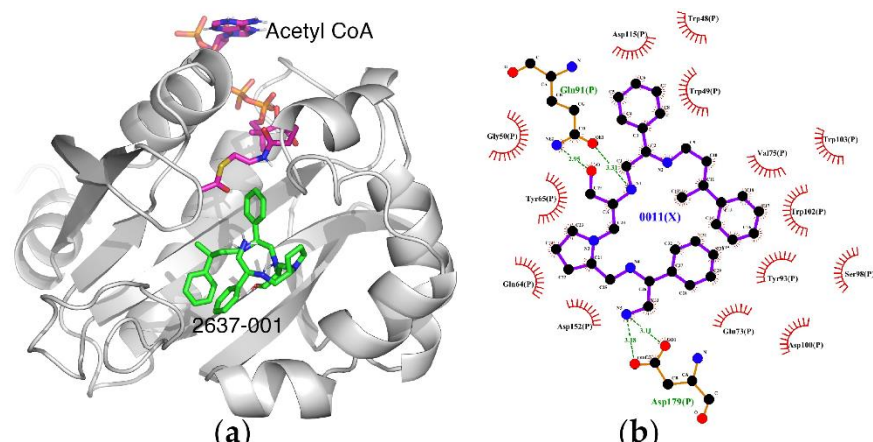


Figure 1. (a) The complex of **2637.001** and AAC(6')-Ib obtained from molecular docking. The bound acetyl CoA is also shown. (b) Interaction map of the ligand in its binding site of the AAC(6')-Ib receptor. The map shows the hydroxyl functionality interaction with Gln91 and the primary amine of **2637.001** hydrogen bonding with Asp179.

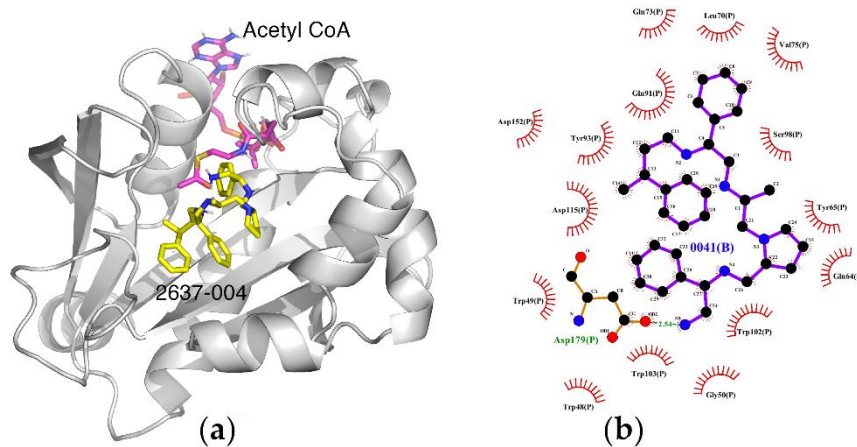


Figure 2. (a) The complex of **2637.004** and AAC(6')-Ib obtained from molecular docking. The bound acetyl CoA is also shown. (b) Interaction map of the ligand in its binding site of the AAC(6')-Ib receptor. The primary amine of **2637.004** maintains a hydrogen bond interaction with Asp179.

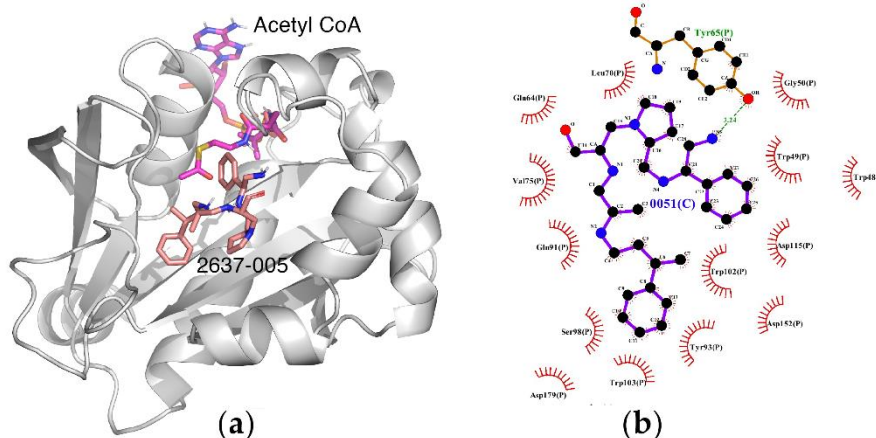


Figure 3. (a) The complex of **2637.005** and AAC(6')-Ib obtained from molecular docking. The bound acetyl CoA is also shown. (b) Interaction map of the ligand in its binding site of the AAC(6')-Ib receptor. Compound **2637.005** adopts a different orientation in the pocket and now the primary amine interacts with the phenol group of Tyr65 and the hydroxyl group is no longer close enough to hydrogen bond with Gln64.

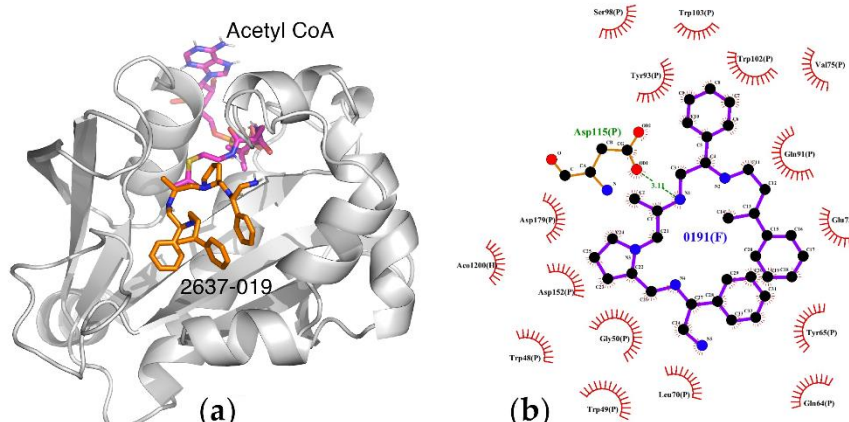


Figure 4. (a) The complex of **2637.019** and AAC(6')-Ib obtained from molecular docking. The bound acetyl CoA is also shown. The figure shows a potential for an intramolecular pi stacking between the R1 and R5 phenyl groups. (b) Interaction map of the ligand in its binding site of the AAC(6')-Ib receptor. Compound **2637.019** still maintains a hydrogen bond interaction with Asp115.

3.3. Potentiation

A more precise analysis of the ability of the analogs listed in Table 1 that did not show a significant deviation ($p<0.01$) in inhibitory activity from **2637.001** was carried out using checkerboard assays. The percent growth inhibition results at all doses are shown in Figure S2 and the analyzed results in Table 2. The checkerboard assays confirmed that compounds **2637.020**, **2637.007**, **2637.010**, **2637.011**, and **2637.013** do not significantly differ in potentiation behavior from **2637.001**. For example, amikacin has a potentiated IC_{50} of 9.5 μ M (95% C.I. (7.5, 11.4)) in the presence of 8 μ M of **2637.020**, versus equivalent values of 8.5 μ M (95% C.I. (5.5, 11.5)) for **2637.001**. Similar overlap in confidence intervals for the other compounds are shown in Table 2.

Three compounds, **2637.004**, **2637.012**, **2637.014**, showed slight reduction in the initial screening, and were confirmed to have significantly lower potentiating ability in the checkerboard assay. For example, amikacin has a potentiated IC_{50} of 20.6 μ M (95% C.I. (14.9, 26.2)) in the presence of 8 μ M of **2637.004**, versus equivalent values of 8.5 μ M (95% C.I. (5.5, 11.5)) for **2637.001**. Similar non-overlap in confidence intervals for the other compounds are shown in Table 2.

Compound **2637.019** showed antimicrobial activity on its own when tested in the checkerboard assay, with a median percent inhibition of 19.1% at 16 μ M with no amikacin. No other compound tested in the checkerboard assay exceeded 6.3% inhibition at this dose (See Figure S2). The adjusted valued after discounting the antimicrobial activity showed that the compound **2637.019** did not show any consistent difference in potentiation ability versus **2637.001** (as seen in overlap of confidence intervals in Table 2).

Table 2. Summary of checkerboard assays

Compound name	R group modified	Compound Dose (μM)									
		0	4	8	16	24	0	4	8	16	24
IC ₅₀ μM	95% CI		IC ₅₀ μM	95% CI		IC ₅₀ μM	95% CI		IC ₅₀ μM	95% CI	
2637.001	NA	24.9	18.2	31.5	12.9	11.1	14.7	8.5	5.5	11.5	0.8
2637.020	R1	23.6	21.2	26.0	12.6	10.3	14.8	9.5	7.5	11.4	0.3
2637.004	R3	33.9	28.5	39.3	25.0*	18.6	31.3	20.6*	14.9	26.2	10.1*
2637.019	R3	31.9	24.6	39.2	33.1*	25.6	40.5	11.5	9.3	13.7	0.3
2637.007	R5	29.9	25.2	34.6	14.8	10.3	19.3	8.8	6.9	10.7	4.0*
2637.010	R5	31.0	25.6	36.4	9.2	7.6	10.8	5.5	4.1	6.9	1.7
2637.012	R5	33.3	27.4	39.1	25.9*	21.8	29.9	20.6*	17.9	23.3	10.4*
2637.011	R5	22.7	18.3	27.0	16.0	14.0	18.0	11.0	9.9	12.2	0.6
2637.013	R5	29.7	24.2	35.2	15.3	12.6	18.1	8.0	5.9	10.0	0.7
2637.014	R5	33.6	29.1	38.1	24.9*	20.4	29.4	19.5*	14.8	24.2	9.1*

IC₅₀ with 95% Confidence intervals for compounds tested in the checkerboard assays. All values are based on curve-fitting of Hill's equation. * indicates a non-overlapping confidence intervals demonstrating a significant reduction in potentiation ability of amikacin at that dose. Yellow highlighting indicates IC₅₀ values less than half of the minimal checkerboard dose, thus being interpolate estimates. NA, not applicable.

4. Discussion

The quest to confront the antibiotic resistance crisis, one of the top threats to human health, requires multifactorial approaches to stop the selection and dissemination of resistant pathogens, design or discover new antimicrobials, and devise strategies to prolong drugs' useful life [4,33,34]. A very successful approach to achieve this latter objective in the case of β -lactams was developing inhibitors of β -lactamases that are administered in combination with the antibiotic to eliminate the pathogen's ability to hydrolyze the antibiotic[5,6]. Unfortunately, such a successful alternative has not yet been fully developed for aminoglycosides. Despite significant efforts, no formulations that combine an aminoglycoside and an inhibitor of the resistance have been approved for human use [4,35]. Although there are many mechanisms and variations by which bacteria resist aminoglycosides, the presence of AAC(6')-Ib in the majority of Gram-negative amikacin-resistant clinical strains implies that the search to find inhibitors that permit their use in a significant number of infections may not be as insurmountable as it seems [4,8,36]. In particular, effective inhibition of AAC(6')-Ib-mediated resistance would restore the efficiency of amikacin as treatment of the currently most dangerous MDR carbapenem-resistant infections [37,38].

Using mixture-based combinatorial libraries and the positional scanning strategy led to identification of an inhibitor of AAC(6')-Ib that, when supplied in combination with amikacin, overcame resistance in several bacteria[23]. The chemical structure of this compound consists of a pyrrolidine pentamine scaffold with two S-phenyl, an S-hydroxymethyl, and a 3-phenylbutyl groups at the positions shown in Table 1. In this study, a series of analogs were analyzed to gain insights into the role that parts of the scaffold, stereochemistry, and functional groups decorating the scaffold play in driving inhibitory activity and ultimately potentiation. For most of the positions (R2, R3, and R4) the absolute stereochemistry of the parent compound was critical for maintaining the inhibitory activity of the compound. The truncation studies showed that the complete pyrrolidine pentamine scaffold is necessary for maintaining inhibitory activity (though further studies remain with truncations in the opposite direction to conclusively determine if the entire scaffold is necessary). At most of the positions (R1, R3, and R5) there was at least a single point substitution analog that maintained the level of inhibitory activity and ultimately the ability to potentiate amikacin at levels comparable to the parent **2637.001** compound.

A molecular docking approach showed that the compounds compete for the same binding site as amikacin against AAC(6')-Ib, further validating the potential mechanism by which the compounds potentiate amikacin. Through this same molecular docking approach it was evident that some of the compounds with better inhibitory activity have more binding interactions with AAC(6')-Ib than those with weaker inhibitory activity.

Taken together, the results shown in this study, validate the concept that inhibiting AAC(6')-Ib is a potential venue to preserve the antimicrobial efficacy of amikacin against Gram-negative amikacin-resistant clinical strains. A medicinal chemistry approach that incorporates molecular modeling to explore additional analogs based on the pyrrolidine pentaamine scaffold holds promise of identifying clinical candidates.

Supplementary Materials: Supplementary information is available online at www.mdpi.com/xxx/s1, Figure S1: Complex between AAC(6')-Ib and amikacin, Figure S2: Check-erboard assays.

Author Contributions: Conceptualization, M.E.T., C.P., M.G., R.S., and P.C.; investigation, K.R., J.M., C.R., K.P., P.V., C.O-H, M.S.R., T.L.V., H.D., P.C., and S.K.; software, R.S. and P.C.; validation, formal analysis, M.E.T., C.P., M.G., R.S., K.R., J.M., and P.C.; resources, M.E.T., C.P., M.G., R.S., and P.C.; writing—original draft preparation, M.E.T., C.P., P.C., and M.G.; writing—review and editing, M.E.T., C.P., M.G., R.S., M.S.R., and P.C. supervision, M.E.T. and C.P.; funding acquisition, M.E.T. All authors have read and agreed to the published version of the manuscript.

Funding: This research was funded by the National Institute of Allergy and Infectious Diseases, National Institutes of Health, grant number R15AI047115. 60
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Conflicts of Interest: The authors declare no conflict of interest. 62

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