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Targeting Class A and C Serine β -Lactamases with a Broad-Spectrum Boronic Acid Derivative

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ABSTRACT: Production of β -lactamases (BLs) is the most widespread resistance mechanism adopted by bacteria to fight β -lactam antibiotics. The substrate spectrum of BLs has become increasingly broad, posing a serious health problem. Thus, there is an urgent need for novel BL inhibitors. Boronic acid transition-state analogues are able to reverse the resistance conferred by class A and C BLs. We describe a boronic acid analogue possessing interesting and potent broad-spectrum activity vs class A and C serine-based BLs. Starting from benzo(b)thiophene-2-boronic acid (BZBTH2B), a nanomolar non- β -lactam inhibitor of AmpC that can potentiate the activity of a third-generation cephalosporin against AmpC-producing resistant bacteria, we designed a novel broad-spectrum nanomolar inhibitor of class A and C BLs. Structure-based drug design (SBDD), synthesis, enzymology data, and X-ray crystallography results are discussed. We clarified the inhibitor binding geometry responsible for broad-spectrum activity vs serine-active BLs using double mutant thermodynamic cycle studies.



INTRODUCTION

Production of β -lactamases (BLs) (EC 3.5.2.6) is one of the major mechanisms by which bacteria develop resistance to β -lactam antibiotics. BLs are classified into serine-based BLs (classes A, C, and D) and metallo- β -lactamases (MBLs), which are class B BLs. These enzymes hydrolyze the amide bond in the β -lactam ring, which constitutes the key feature of this type of antibacterial agent. The substrate spectrum of BLs now includes carbapenems and the latest generation of cephalosporins.¹

The use of a BL mechanism-based inhibitor in combination with a β -lactam antibiotic is a well-established strategy to fight resistance.² However, the three commercially available β -lactam inhibitors (clavulanate, sulbactam, and tazobactam) are effective primarily against class A BLs (Figure 1B), while class C and D enzymes are only poorly inhibited by any of the three commercial inhibitors. Moreover, no clinically useful inhibitor of MBLs has been identified to this point.³

These BL inhibitors share a β -lactam ring, making them susceptible to resistance stemming from up-regulation of BL production, selection for new BLs, and other mechanisms that evolved over millions of years of chemical exposure between bacteria and β -lactam-producing microorganisms (Figure 1B).^{4,5}

Among class A plasmid-encoded BLs, TEM and CTX are the most commonly observed extended-spectrum BLs (ESBLs);

they are found mainly in *Escherichia coli* and *Klebsiella* pneumoniae and confer a high level of resistance to available broad-spectrum cephalosporins. The TEM strains responsible for nosocomial epidemics are usually multiresistant to antibiotics, and most of them produce ESBLs.^{1,6,7} CTX-M is a group of class A BLs that are particularly resistant to extended spectrum β -lactam antibiotics such as cefotaxime, which itself was developed to counter bacterial resistance to first-generation penicillins and cephalosporins (Figure 1C).⁸ Since its discovery in the late 1990s, CTX-M has become the most frequently observed ESBL in many regions of the world.

These families of BLs present a significant clinical threat, with CTX-M-14 and CTX-M-15 being the most prominent ESBLs worldwide and TEM BLs exhibiting the most variants.⁹

Regarding class C, resistance due to plasmid-mediated AmpC enzymes is produced by BL overexpression, conferring resistance to broad-spectrum cephalosporins (i.e., *Enterobacter aerogenes* and *Enterobacter cloacae* infections) and causing outer-membrane porin modifications (carbapenem resistance) and plasmid transmission (*E. coli, K. pneumoniae* and *Proteus mirabilis* infections).¹⁰

To treat antimicrobial multiresistant pathogens, a second-generation BL inhibitor era has already begun, which mainly

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Figure 1. β -Lactamases inhibitors and β -lactam antibiotics. (A) Starting lead compound 1, BZBTH2B, benzo(b)thiophene-2-boronic acid. (B) The BL inhibitor clavulanic acid. (C) Cefotaxime, III generation cephalosporin. (D) The last generation BL inhibitor avibactam. (E) Broad spectrum inhibitor compound 5, 3-(2-carboxyvinyl)benzo(b)thiophene-2-boronic acid. (F) Inhibitor SMS2 ((1R)-1-(2-thienylacetylamino)-1-(3-carboxyphenyl)methylboronic acid.

focuses on novel non- β -lactam inhibitors showing broadspectrum profile.^{2,3,11-18} Derivatives such as avibactam and its analogues have now reached in combination with ceftazidime clinical phase II, representing a promising weapon against bacterial resistance (Figure 1D).^{19–21} Conversely, an ideal MBL inhibitor remains to be found despite the large number of potential molecules already described.²²

Among novel non- β -lactam inhibitors, we introduced boronic acid transition-state analogues that bind to AmpC BL with nanomolar affinities: this novel chemistry was able to reverse the resistance conferred by these enzymes, in particular for those belonging to class C.^{16,18–20}

Starting from benzo(b)thiophene-2-boronic acid (BZBTH2B, compound 1, Figure 1A), a nanomolar non- β lactam inhibitor of AmpC able to potentate the activity of a third-generation cephalosporin against class C-producing resistant bacteria without inducing overexpression of AmpC, we designed a broad-spectrum nanomolar inhibitor. Serinedependent BLs have similar folding and contain a S-X-X-K motif, with serine acting as the catalytic residue. Starting from the available crystal structure of compound 1 binding to the AmpC active site and its visual analysis in comparison with class A TEM-1 and CTX-M-9 BLs, we identified one potential binding spot in the enzymes and a corresponding (or reciprocal) chemical point of derivatization in the skeleton of compound 1 that was exploited to produce stronger binding affinity and, most important, broad-spectrum activity.^{23–25} In particular, our comparative analysis suggested the possibility of introducing a polar, carboxylated lateral chain (i.e., 2-carboxyvinyl) in compound 1 to gain new favorable electrostatic and H-bond interactions with the C3(4') carboxylate consensus binding pocket, common to all serine-active BLs.

The newly designed inhibitor, compound **5** (Figure 1E) was then synthesized and tested, resulting in a potent and broadspectrum inhibitor. Its binding orientation in CTX-M-9 and its critical binding interactions with TEM-1 were subsequently clarified and confirmed by means of X-ray crystallography and free energy double-perturbation thermodynamic analysis.

Thermodynamic cycles can be used to confirm and/or quantify the interaction between a functional group of a ligand and an active site residue.²⁶⁻²⁹ In these experiments, wild-type and "mutant" versions of both the enzyme and ligand are used, with each "mutant" lacking the interacting groups of interest. The energetic costs of removing the ligand functional group both in the presence and absence of the residue with which interactions occurs are calculated. The difference between these energies provides the energy involved in the key interaction. Quantitative knowledge of the energy exploited in the new interactions that were critical for binding to TEM-1 and indicated fundamental directions for further optimization.

RESULTS AND DISCUSSION

3-D Comparative Analysis of Serine-Based BLs and Inhibitor Design. Starting from compound 1, we were interested in obtaining a validated lead compound with conserved or improved affinity vs class C and, most important, with extended activity vs class A. To localize the more appropriate point of derivatization and to rationally select the chemical groups suitable for derivatization, we referred to three X-ray structures. The complex of AmpC BL with compound 1 $(PDB \text{ code } 1C3B)^{23}$ and the two complexes of SMS2, a glycylboronic acid inhibitor ((1R)-1-(2-thienylacetylamino)-1-(3-carboxyphenyl)methylboronic acid) (Figure 1F) binding TEM-1 (PDB code 1NXY)²⁴ and CTX-M-9 (PDB code 1YM1),³⁰ were analyzed (2a,b). X-ray structures for comparative analysis were selected based on the broad activity spectrum of SMS2. In the derivatization of glycylboronic acids, the introduction of a carboxyphenyl group not only improved the affinity toward AmpC but, most important, also extended the spectrum of inhibition vs TEM-1 and CTX-M-9 via hydrogen-bond with Arg244 in TEM-1 and Arg276 in CTX-M-9. Both residues are located in the β -lactam carboxylate binding pocket.³¹ Our approach was guided by previous results: within boronic acid, a carboxy group designed to mimic the C3(4)'carboxylate of β -lactam substrates is a key recognition feature in class A and C $\beta\text{-lactamases.}^{15,24,25}$

The lead compound **1** in the AmpC binding site (PDB code 1C3B) forms quadrupole–quadrupole interactions with Tyr221 and quadrupole–dipole interactions with Asn152 (Figure 2a). Distances and angles are consistent with quadrupolar



Figure 2. (a–c). 3-D comparative analysis of class A and C β lactamases guided inhibitor design. (a) Binary complex of BZB2THB in AmpC BL. Key residues and interactions are highlighted. Carbon atoms are in green, nitrogen atoms are colored blue, oxygen atoms red, and sulfur atoms yellow. Red spheres represent water molecules. Picture was prepared using CCP4mg.⁵³ (b) Glycylboronic acid inhibitor SMS2 complexes with CTM-M-9 (carbon atoms in blue) and TEM-1 (carbon atoms in orange) are superimposed. The interactions with C3(4')carboxylate binding pocket of β -lactams are highlighted. Nitrogen atoms are colored blue, oxygen atoms red, and

Figure 2. continued

sulfur atoms yellow. Red spheres represent water molecules. Picture was prepared using CCP4mg.⁵³ (c) Molecular surface of the superimposed active site region of CTX-M-9, TEM-1, and AmpC. Glycylboronic acid inhibitor SMS2 complexes with CTM-M-9 (carbon atoms in blue) and TEM-1 (carbon atoms in orange) were superimposed with BZB2THB in complex with AmpC (carbon atoms in green). The C3(4')carboxylate binding pocket of β -lactams and the C3 position in BZB2THB selected for chemical derivatization are highlighted. The surface contributed by nitrogen atoms is colored blue, oxygen atoms are colored red, and carbon atoms are colored gray. Picture was prepared using CCP4mg.⁵³

interactions observed in other protein structures.^{32,33} An observed interaction is the hydrogen bond between the boronic acid O2 atom and Tyr150. Two well-ordered water molecules appear in the active site, making extensive interactions that involve the inhibitor and Tyr150. The first water molecule interacts through a hydrogen bond with the boronic acid O2 atom and with the active site residue Thr316. The second water molecule forms a hydrogen bond with the first water molecule. This second water molecule also interacts with catalytic residues Asn346 and Arg349. Additionally, the boronic acid O1 atom hydrogen binds with the backbone nitrogens of Ser64 and Ala318 and also with the carbonyl oxygen of Ala318.²³

Regarding the binary complex of TEM-1 binding SMS2 (PDB code 1NXY),²⁴ we can observe that the inhibitor adopts a deacylation transition-state analogue conformation in which the boronic acid O1 atom is still in the "oxyanion" while the configuration around the boron has inverted; therefore, the boronic acid O2 atom hydrogen binds with the catalytic base Glu166, displacing the ordered, catalytic water molecule (Figure 2b).³⁴ The phenyl ring of SMS2 stacks with the aromatic ring of Tyr105, making quadrupole interactions.^{32,33} The carboxylate group hydrogen binds with Arg244, Ser235, and an ordered and highly conserved water molecule. These interactions between Arg244, Ser235, and water are also seen with the C3(4)'carboxylate of β -lactams, consistent with our carboxylated inhibitor designed to mimic this ubiquitous substrate group (Figure 2b).³⁵

In the binary complex CTX-M-9–SMS2 (PDB code 1YM1), key interactions between the boronic acid group and CTX are analogous to those already described for TEM-1. The aromatic ring of the inhibitor makes van der Waals contacts with the aromatic ring of Tyr105 (distances from 3.3 and 4.6 Å). Moreover, it is 3.0 Å away from the Ser130 O γ atom, with the hydroxyl group perpendicular to the face of the ring, in a typical dipole–quadrupole interaction geometry.³² The carboxylic acid groups of SMS2 interact with O γ 1 of Thr235 and three well-ordered water molecules in the ubiquitous C3(4')carboxylate binding pocket, as previously observed in TEM-1 complexes. Arg-276, which is implicated in the catalytic process of CTX-M enzymes, in one monomer interacts with the *m*-carboxy group of SMS2 via an ordered water molecule.²⁴

When the three structures were superimposed for comparison, in addition to the main differences in the residues between class A and class C, position 3 of lead compound 1 always pointed toward key residues involved in the C3(4)'carboxylate interactions, namely Asn346 and Asn289 in AmpC (distance, 8.74 Å N δ 2 Asn346), Arg244 (8.14 Å NH1 Arg244), and Ser235 (8.92 Å O γ Ser235) in TEM-1 and Arg276 in CTX-M-9 (8.28 Å NH2 Arg276) (Figure 2c). The visual analysis identified position 3 in lead compound **1** as the most convenient derivatization point to introduce appropriate chemical groups able to reach the carboxylate binding site delimited in AmpC by Thr316 and Asn346, by Arg244 in TEM-1 and by Arg276 in CTX-M-9 (Figure 2c).³⁶

Having selected and confirmed position 3 for chemical derivatization, we decided to introduce a 2-carboxyvinyl chain able to mimic the distance, chemistry, and interactions of SMS2 C3(4)'phenylcarboxylate (Compound 5, Table 1). Compound

Table 1. Ki Values of BZB2THB Derivatives vs Serine β -Lactamases



4, the keto analogue of compound **5**, was additionally synthesized as a tool for free energy double-perturbation thermodynamic cycle, thus confirming our hypothesis of the importance of the carboxylate moiety in modulating affinity and potency vs BLs.

Synthesis. 2-(4,4,5,5-Tetramethyl-1,3,2-dioxaborolan-2-yl)benzo[b]thiophene-3-carbaldehyde (3) was designed and synthesized as a scaffold for synthetic elaboration. We first reported the formylation reaction of compound 2 as a key step in the derivatization at the 3 position of compound 1. In Scheme 1, we report the synthetic route for the synthesis of compound 4 and 5. The boronic acid group of 1 was protected using 2,3-dimethyl-2,3-butanediol, and the resulting compound 2 was treated with dichloromethyl methyl ether in the presence of Tin(IV) chloride to produce compound 3 with 75% yield. A

Scheme 1^a

Wittig reaction from 3 and 1-(triphenylphosphoranylidene)-2propanone in a 1:1 molar ratio was performed to produce compound 4 as pinacol-protected boronic acid. A Wittig reaction between *tert*-butoxycarbonylmethylene-triphenylphosphorane and compound 3, followed by hydrolysis, was performed, yielding compound 5 as pinacol-protected boronic acid.

BLs Inhibition Studies. To investigate the effectiveness of the new analogues, we employed CTX-M-9, TEM-1, and AmpC BLs and a UV absorbance-based biochemical assay to obtain binding affinities. As previously reported for other boronic acid inhibitors of BLs, tested compounds were competitive inhibitors and no incubation effect was detected. The inhibitory activity of analogues 4 and 5 carrying a lateral chain at position 3 of compound 1 was evaluated, with the expectation that compound 5 would form favorable electrostatic interactions with the carboxylate binding pocket of BLs, as previously described. As expected, the 3-carboxy derivatives 3-(2-carboxyvinyl)benzo(b)thiophene-2-boronic acid (compound 5) showed broad-spectrum activity vs classes A and C; compound 5 inhibits with low nanomolar potency class A, while its affinity vs AmpC was not diminished compared to the starting lead, compound 1 (K_i vs TEM-1 and CTX-M-9 40 nM, K_i vs AmpC 90 nM, Table 1). On the other hand, compound 4, bearing a keto group instead of a carboxy group, maintains nanomolar affinity vs AmpC (8 times less active) but loses potency vs class A, highlighting how critical the carboxylic group in class A is for molecular recognition (125 times less active vs class A with respect to class C). Moreover, compound 5 was also tested vs the AmpC mutant N289A, in which the wild-type asparagine is replaced by an alanine; its affinity was not reduced (K_i 50 nM vs N289A), thus supporting the hypothesis that in AmpC the carboxylate interacts with Asn346, as the β -lactam do and not with the adjacent Asn289.

X-ray Crystallographic Structure Determination. To investigate the structural basis for the relative affinities and to understand the details of recognition along the reaction coordinates, we determined the crystal structure of CTX-M-9 enzymes in complex with compound **5** to 1.50 Å resolution. Excluding proline and glycine, 91.5% of residues were in the favored region, 8.0% of residues were in the allowed region of the Ramachandran plot, and only 0.4%, were in the generously



"(a) 2,3-Dimethyl-2,3-butanediol, dry ethyl ether, RT; (b) tin chloride, dichloromethyl methyl ether, dichloromethane, -20 to 0 °C; (c) 1-(triphenylphosphoranylidene)-2-propanone, dry dichloromethane, RT; (d) *tert*-butoxycarbonylmethylene-triphenyl posphorane, dry dichloromethane, RT; (e) dichloromethane trifluoroacetic acid, 50%, RT.



Figure 3. (a) Electron density of compound **5** in complex with CTM-M-9. The $2F_o - F_c$ electron density map is represented by the green cage and is contoured at 1.0σ . The inhibitor is covalently attached to Ser70. Carbon atoms of CTM-M-9 and compound **5** are colored green, Nitrogen atoms are colored blue, oxygen atoms red, and sulfur atoms yellow. Picture was prepared using Pymol.⁵⁴ (b) Key interactions observed in the complex of **5** with CTM-M-9, site B. The inhibitor is covalently attached to Ser70. Carbon atoms of CTX are colored gray, carbon atoms for **5** are colored green. Nitrogen atoms are colored blue, oxygen atoms red, and sulfur atoms yellow. Red spheres represent water molecules. Residues Ser130 and Ser237 are in double conformation. Picture was prepared using CCP4mg.⁵³

allowed region.³⁷ Several double conformations were revealed, some of which were involved in active site geometry (chain A, 59, 98, 186, 201, 222, 262, 271, and 281; chain B, 75, 89, 113, 158, 186, 237, 249, 274, and 281). In the binding site, the position of inhibitor 5 was unambiguously identified in the initial $F_{o} - F_{c}$ electron density difference map contoured at 1σ (Figure 3a). Electron density connects the $O\gamma$ of the catalytic Ser70 to the boron atom of boronic acid. The geometry of the boronic acid group is tetrahedral (Figure 3a,b). In the CTX-M-9-compound 5 complex, the 021 boronic acid hydroxyl interacts via a hydrogen bond with the "oxyanion" hole N and the O backbone atom of residue 237 (distance 2.88 and 2.74 Å, respectively, in chain A, Figure 3b, Table 2) and N of Ser70 (distance 2.69 Å in chain A, Figure 3b, Table 2). The 022 boronic acid hydroxyl interacts with $O\delta 1Asn170$ (2.63 Å) and with $O\varepsilon 1$ Glu166 (2.66 Å), replacing the catalytic water molecule, which is not present in this complex. In this structure, the O22 atom represents the position of the catalytic water molecule in the deacylation transition state, consistent with similar interactions observed in a TEM-1 boronic acid structure.³⁸ Overall, the boronic acid complex of compound 5 adopts a conformation consistent with a deacylation transitionstate analogue.³⁹ The benzothiophene ring interacts with Tyr105 ($C15-C\beta$ Tyr105, 3.4 Å). The lateral chain of compound 5 makes contact with Ser130 (C β Ser130–C18 3.4 Å). The carboxylic acid groups of compound 5 interact with surrounding residues and a well-ordered water molecule in a network different from one chain to the other. In chain A, O23 interacts with the Oy1 of Thr235 (O23-Oy1Thr235 2.75 Å), with Gly236 (NG236- O23 3.10 Å) and via a well-ordered water molecule with Tyr105 (O23-Wat410 2.70 Å, Wat410-OHTyr105 3.6 Å). O23 also interacts with three well-ordered water molecules in an extended H-bond network that involves Arg-276, which is implicated in the catalytic process of CTX-M enzymes.40 This residue adopts a different conformation in chain B; the interaction between Arg276 and the carboxylate group is closer, and only one ordered water molecule mediates the H-bond (N&Arg276-Wat229 2.98 Å, Wat229-O23 3.2 Å). Moreover, in chain B as well as in chain A, several well-ordered water molecules participate in an extended H-bond network stabilizing the complex. The described interaction pattern is consistent with the predicted inhibitor carboxylate group role:

Table 2. Key Interactions in the Complex Structure

	23 0 24 0 0 0 8 0H 21	
	distance (Å)	
interactions	monomer A	monomer B
S70N-O21	2.76	2.69
S237N-O21	2.94	2.88
S237O-O21	2.78	2.74
\$1300γ-024	3.63	3.34
E166O <i>ε</i> 1–O22	2.59	2.66
E166O <i>e</i> 2-O22	3.33	3.32
N170Oδ1−O22	2.65	2.63
N170Nδ2−O22	3.19	3.26
G236N-O23	3.10	
Τ235Ογ1–Ο23	2.75	3.34
Τ235Ογ1-Ο24	3.42	2.46
Κ234Νζ-Ο24	2.94	3.45
Wat 61-O23	2.86	
Wat482-O23	3.19	
Wat 488-O23	2.82	
Wat229-O23		3.20
R276N ϵ -Wat229		2.98
Wat410-O23		2.70

the introduced chemistry mimics the ubiquitous C(3)4' substrate carboxylate, as previously observed in the TEM and CTX complexes.^{24,25}

Double Mutant Thermodynamic Cycle Analysis (DPA). To study the energetic effect of the 3-(2-carboxy-vinyl) group on compound 5 activity vs TEM-1, a double-perturbation analysis was performed. Although it was clear from the X-ray complex CTX-M-9-5 that the 3-(2-carboxy-vinyl) group interacts via hydrogen bond with Arg276, the energies of these interactions were unknown. In addition, lacking a 5– TEM-1 X-ray complex, we wanted to confirm this key interaction also for TEM-1. For instance, moving from 1 to its (2-carboxy-vinyl) derivative **5**, binding energy vs TEM-1 improves by 2.8 kcal/mol, but it was unclear how much of this energy is specifically derived from the hydrogen bond with Arg276. This binding energy could alternatively result from ionic interactions between the carboxylate and positively charged protein residues, or through hydrogen bonds with ordered water molecules. Quantitative knowledge of the energy of these interactions responsible for the high affinity of the lead compound should be preserved in any future iteration. Moreover, DPA could fill the gap of information derived from a missing compound 5-TEM-1 X-ray complex.

To quantify the energetic contributions of the hydrogen bond with Arg276 in CTX-M-9, to confirm the role of the carboxyl group in the H-bond interaction in TEM-1 and to definitively identify which residue in TEM-1 was involved in the aforementioned interaction, we prepared DPA. In the experiments, we used wild-type and "mutant" versions of both the enzyme and ligand, with each "mutant" lacking the interacting groups of interest (Scheme 2a). The energetic

Scheme 2. Double Perturbation Analysis: (a) Ligands, (b) Double Perturbation Analysis Cycle^{*a*}



^aThermodynamic cycle to quantify the hydrogen bond energy between Arg244 and 5. E = wild-type enzyme, E' = mutant enzyme. L = wild-type ligand, L' = "mutant" ligand. $\Delta\Delta\Delta G = (\Delta\Delta G_1 - \Delta\Delta G_4)$ or $(\Delta\Delta G_2 - \Delta\Delta G_3)$. E = Tem-1, E' = Tem-30 R244S. L = 5, L' = 4. $\Delta\Delta\Delta G_{1-4} = 2.73$ kcal/mol. $\Delta\Delta\Delta G_{2-3} = 2.75$ kcal/mol.

cost, $\Delta\Delta G$, of removing the ligand functional group both in the presence and the absence of the residue with which it interacts was calculated, and the difference between these energies, $\Delta\Delta\Delta G$, is the energy of the interaction (Scheme 2b).

Compound 5 loses its affinity vs TEM-1 when in TEM-30 Arg244 is replaced by a Ser residue (220-fold less active, K_i from 0.04 to 8.8 μ M). In terms of binding interaction, the energetic cost is 3.18 kcal/mol ($\Delta\Delta G_2$), suggesting a critical interaction with Arg244. On the other hand, varying the

inhibitor structure by masking the carboxylate functionality as with the keto derivatives 4, the inhibitor loses activity again vs TEM-1 wild-type (120-fold less active, K_i from 0.04 vs 4.8 μ M, $\Delta\Delta G_1$ 2.8 kcal/mol), supporting the hypothesis of an H-bond between the carboxylate and Arg244. The following doublemutation steps, when both the enzyme and inhibitor are "mutated," confirmed this finding. The binding energy of the keto derivative 4 vs TEM-30 with respect to compound 5 vs TEM-30 did not change (less than 1-fold, K_i 8.8 vs 9.9 μ M, respectively, $\Delta\Delta G_4$ 0.07 kcal/mol). Moreover, when 4 was tested against TEM-30, the affinity with respect to the binding vs TEM-1 drops by only 2-fold (K_i from 4.8 vs 9.9 μ M, $\Delta\Delta G_3$ 0.43 kcal/mol). When we calculated the $\Delta\Delta\Delta G$ for the four mutation steps, we found that the binding energy contribution of the carboxylate group vs Arg244 was in perfect agreement with the presence of an H-bond (Scheme 2b: DPA cycle, $\Delta\Delta G_1$) $-\Delta\Delta G_4 = 2.73$ kcal/mol, $\Delta\Delta G_2 - \Delta\Delta G_3 = 2.75$ kcal/mol). We can certainly affirm that in TEM-1, Arg244 picks up the Hbond interactions with the carboxylate group of compound 5, as crystallographically observed in the compound 5-CTX-M-9 complex, where in turn Arg276 is primarily involved in key interactions with compound 5 as well as several well-ordered water molecules. Superposition of the CTX-M-9-5 complex with TEM-1 structure supports and clarifies our finding: TEM-1 Arg276 corresponds in CTX-M-9 to Asn276; however, TEM-1 Arg244 is in the proximity of the compound 5 carboxylate group and can easily pick up H-bond interactions as Arg276 does in CTX-M-9 (Figure 4).

DISCUSSION AND CONCLUSION

The identification of compound **5** as a novel, non- β -lactambased broad-spectrum inhibitor of class A and class C β lactamases is a promising result in the bacterial resistance mediated by BLs.

The purpose of our studies was, beginning with a known, nonextended spectrum lead, to take advantage of key regions present along all BL serine-based active site responsible for binding recognition, e.g., the carboxylate binding domain, in order to introduce broad-spectrum activity.

Our designed ligand 5, through the introduction of a new functional group in the starting lead 1, exhibits low nanomolar affinity vs CTX-M-9, TEM-1, and AmpC BLs. The gain in affinity vs class A with respect to the starting lead 1 was 1250-fold vs CTX and vs TEM-1. Comparing compound 5 activity profile with that of the glycylboronic derivative SMS2, we found that its potency remains unchanged vs TEM-1 (0.040 and 0.064 μ M, respectively) while it is reduced toward AmpC (0.090 vs 0.001 μ M, respectively). Interestingly compound 5 gains affinity toward CTX-M-9 (0.040 vs 0.578 μ M).

The results highlight the importance in inhibitor binding recognition of Arg276 in CTX-M-9 as well as Arg244 in TEM-1; both residues, which are part of the C3(4')carboxylate consensus binding pocket and common to all serine-active BLs, have been observed to interact with β -lactam substrates in X-ray complexes with CTX-M-9 and TEM-1. Therefore, our studies suggested that these residues are hot spots useful for broad-spectrum BL inhibitor design.

Comparing the CTX-M-9–compound **5** (4LEN) complex and the glycil boronic derivative SMS2 binding CTX-M-9 (1YM1), we see that the carboxylate lateral chain is oriented as we predicted in the design, H-bonding to those residues, which are highly conserved in CTX-M-9 and present in TEM-1, that were intentionally targeted (Figure 2b, Figure 4).



Figure 4. Interpreting DPA results. Compound **5** (carbon atoms in green) in binary complex with CTX-M-9 (carbon atoms in gray) is superimposed TEM-1 (carbon atoms in orange). Key residue Arg276, which in CTM-M-9 interacts with compound **5** carboxylate group, is replaced by Asn276 in TEM-1. However, in TEM-1, as suggested from DPA, key interaction could be picked up from Arg244, oriented at H-bond distance from compound **5**. Picture was prepared using CCP4mg.⁵³

In addition, the lateral chain of compound **5** interacts with an opened part of the active site, in a different conformation in chain A with respect to chain B, highlighting a certain mobility and flexibility in the interactions between ligand and protein. In both binding sites, several well-ordered water molecules participate in an extended H-bond network that contribute to stabilize the complex. In compound **5**, the rigidity conferred to the lateral chain by the double bond probably contributes to the overall binding energy due to its restricted mobility, resulting in a tighter binding.

Despite its novelty and broad-spectrum enzymatic activity, compound 5 resulted in being unstable in microbiology assay conditions over time (data not shown). These finding directed the design and synthesis of more stable analogues of 5 now under development.

As a final comment, structurally guided optimization of novel extended BL inhibitors active against class A and C has confirmed binding hot spots that can be targeted to produce high-affinity inhibitors. These hot spots, i.e., the carboxylate binding domain, are shared by therapeutically important groups of BLs, suggesting the hot spots' potential for broad-spectrum inhibition of BLs. The nanomolar potency of **5** poses this compound as an interesting broad-spectrum inhibitor of BLs, which deserves further investigation.

EXPERIMENTAL METHODS

Synthesis. 5-Methylbenzo[b]thiophene and benzo[b]thien-2-ylboronic acid were purchased from Lancaster. All reagents were purchased from Sigma-Aldrich and Fluka and were of reagent grade. Reaction progress was monitored by TLC on precoated silica gel 60 F254 plates (Merck). Silica gel (60 M; 230–400 mesh, ASTM) was used for column chromatography. The purity of all synthesized compounds was determined by elemental analyses, performed on a PerkinElmer 240C instrument, and all values were within ±0.4% of the theoretical values (data not shown). Yields refer to purified products

and were not optimized. All compounds were characterized by ¹H NMR and ¹³C NMR on AC200 and Bruker MX400 WB instruments (CIGS, University of Modena e Reggio Emilia). Unless otherwise stated, spectra were recorded in DMSO- d_6 or CDCl₃. Chemicals shifts are reported in ppm from tetramethylsilane as an internal standard. LC-MSIT analysis was performed through HPLC Agilent 1200 Ion Trap LC/MS Agilent 6310 device. Source of ionization: Electrospray. Column: ZORBAX SB-C18 2.1 mm × 30 mm, particles of 3.5 μ M.

2-(4,4,5,5-Tetramethyl-1,3,2-dioxaborolan-2-yl)benzo[b]thiophene (2). To 0.30 g (1.68 mmol) of benzo[b]thien-2-ylboronic acid (1) in dry diethyl ether (40 mL), 2,3-dimethyl-2,3-butanediol (0.199 g, 1.68 mmol), and a catalytic amount of trifluoroacetic acid was added. The mixture was stirred under nitrogen for 15 min. The solvent was removed under vacuum and crude residue extracted with *n*-pentane, producing **2.** Yield: 0.381 g, 87%. ¹H NMR (DMSO): δ 1.31 (s, 12H), 7.42 (m, 2H), 7.91 (s, 1H), 7.95 (m, 1H), 8.02 (m, 1H). Anal. (C₁₄H₁₇BO₂S) C, H, N.

2-(4,4,5,5-Tetramethyl-1,3,2-dioxaborolan-2-yl)benzo[b]thiophene-3-carbaldehyde (3). Tin chloride (575 μ L, 4.9 mmol) was added dropwise to a stirred solution of **2** (0.425 g, 1.63 mmol) in dry dichloromethane (6 mL) at -20 °C under nitrogen. Dichloromethyl methyl ether (177 μ L, 1.96 mmol) was added dropwise and the mixture allowed to warm to 0 °C. After the mixture had been stirred for 7 h, it was poured into diluted hydrochloric acid and extracted with dichloromethane CH₂Cl₂. The extracts were washed with sodium chloride solution, dried with sodium sulfate, and evaporated. The residue was crystallized with dichloromethane/*n*-hexane, producing **3**. Yield: 0.415 g, 75%. ¹H NMR (CDCl3): δ 1.41 (s, 12H), 7.50 (m, 2H), 7.90 (m, 1H), 8.85 (m, 1H), 10.69 (s, 1H). Anal. (C₁₅H₁₇BO₃S) C, H, N.

4-[2-(4,4,5,5-Tetramethyl-[1,3,2]dioxaborolan-2-yl)-benzo-[b]thiophen-3-yl]-but-3-en-one (4). A mixture of 3 (0.1 g, 0.347 mmol) and 1-(triphenylphosphoranylidene)-2-propanone (0.11 g, 0.348 mmol) in dry dichloromethane (1.5 mL) was stirred at room temperature under nitrogen for 24 h. The solvent was removed under vacuum, and the crude residue was extracted several times with diethyl ether to remove triphenylphosphine oxide. The organic phase was dried on magnesium sulfate and the solvent removed under vacuum, producing 4 as a white crystalline solid. Yield: 0.055 g, 48%.

3-[2-(4,4,5,5-Tetramethyl-[1,3,2]dioxaborolan-2-yl)-benzo-[b]thiophen-3-yl]-acrylic Acid (5). A mixture of 3 (0.30 g, 1.04 mmol) and *tert*-butoxycarbonylmethylene-triphenylphosphorane (0.391 g, 1.04 mmol) in dry dichloromethane (4 mL) was stirred at room temperature under nitrogen for 24 h. The solvent was removed under vacuum, and the crude residue was extracted several times with diethyl ether to remove triphenylphosphine oxide. After solvent removal, the crude extract was treated with dichloromethane/ trifluoroacetic acid 50% and stirred at room temperature for 2 h. The solvent was removed under vacuum and residue washed with diethyl ether, producing 5 as a white crystalline solid. Yield: 0.120 g, 35%. ¹H NMR (DMSO): δ 1.34 (s, 12H), 6.59 (d, 1H), 7.52 (m, 2H), 8.07 (dd, 1H), 8.18 (dd, 1H), 8.25 (d, 1H), 12.41 (b all., 1H). Anal. (C₁₇H₁₉BO₄S) C, H, N.

Inhibition Assays. Versus AmpC. Boronic acids were dissolved in DMSO stock solutions at 50 mM; more dilute stocks were subsequently prepared as necessary by dissolving them in 50 mM phosphate buffer at pH 7. Compounds isolated as pinacol-protected boronic acid were tested without further ester cleavage reaction; compounds were hydrolyzed to the free acid form by dissolving them in 50 mM phosphate buffer at pH 7.4.^{18,41} AmpC from *E. coli* was expressed and purified to homogeneity as described.³⁶ Kinetic measurements were performed using nitrocefin as a substrate in 50 mM Tris buffer, pH 7.0, and monitored in an HP8453 UV-vis spectrophotometer. The K_m of nitrocefin for AmpC in this buffer was 127 μ M. The concentration of AmpC was determined spectrophotometrically in concentrated stock solutions made from lyophilized powder and subsequently diluted; this enzyme had been previously expressed and purified as described.²³ The concentration of enzyme was 1.75 nM. Inhibition K_i values were obtained from IC₅₀ plots assuming competitive inhibition, an assumption consistent with both previous inhibition patterns in this series and with experiments investigating the effect of increasing substrate concentrations (not shown).⁴²

Versus CTX. Boronic acids were dissolved in DMSO stock solutions at 50 mM; more dilute stocks were subsequently prepared as necessary by dissolving them in 50 mM phosphate buffer at pH 7. CTX-M-9 was produced from a modified pET-9a plasmid in *E. coli* BL21 (DE3). The protein was purified by ion exchange and gel filtration, as previously described.⁴³ Enzymes were diluted from stock solutions to a final concentration of 1.5 nM. The enzyme assay was carried out in 50 mM potassium phosphate (pH 7.0) at room temperature and monitored in an HP8453 UV–vis spectrophotometer. The reaction was monitored at 340 nm using 6- β -furylacryloylamido-penicillanic acid (100 μ M, FAP, Calbiochem) as substrate (the K_m values for CTX-M-9 and CTX-M-16 were 17 and 6.5 μ M, respectively). The progress curves were measured at least three times for each substrate.²⁵ K_i values were obtained by comparing the progress curves in the presence and absence of the inhibitor using the method described by Waley.⁴⁴

Versus TEM-1. Boronic acids were dissolved in DMSO stock solutions at 50 mM; more dilute stocks were subsequently prepared as necessary by dissolving them in 50 mM phosphate buffer at pH 7. TEM-1 enzyme was expressed and purified as described.⁴⁵ TEM-1 enzyme assays used 100 mM 6- β -furylacryloylamido-penicillanic acid (FAP, Calbiochem) as a substrate, monitoring absorbance changes at 340 nm on an HP8453 spectrophotometer. Reactions were initiated with addition of 0.3 nM enzyme, using the same buffer as in the AmpC assays. The K_i values for compounds were obtained by the comparison of the progress curves in the presence and absence of the inhibitor. This method correlates well with full K_i analysis through coupled substrate and inhibitor concentration variation.⁴² TEM-30 used for double perturbation thermodynamic cycle was prepared as reported.⁴⁶

Crystallization. Cocrystals of CTX-M-9 enzyme in complex with 5 were grown by vapor diffusion in hanging drops equilibrated over a well solution of 1.6 M potassium phosphate buffer (pH 8.8) using micro seeding techniques and microcrystals of CTX-M-9 previously grown under the same conditions. To a solution of 15 mg/mL protein in 5 mM Tris (pH 7.0) and 30 mM NaCl was added an equal volume

of 1 mM inhibitor in a solution of 5% DMSO, 1.25 M potassium phosphate (pH 8.8). Crystals appeared in 8–24 h after equilibration at 20 °C. Before data collection, crystals were immersed in a cryoprotectant solution of 30% sucrose, 1.8 M potassium phosphate (pH 8.8) for approximately 30 s and then were flash frozen in liquid nitrogen.²⁵

Data Collection and Structure Determination. Data were measured at 100 K using an ADSC–CCD detector on Beamline 8.3.1 of the Advanced Light Source at Lawrence Berkley National Laboratory. Reflections were indexed, integrated, and scaled using the HKL software package.⁴⁷ The space group was P2₁, with two molecules in the asymmetric unit. Phases were calculated by molecular replacement with the program EPMR22 using the apoenzyme structure of CTX-M-9, with the water molecules and ions removed (Table 3).³⁰ The complex structures of CTX-M-9 with compound **S** at

 Table 3. X-ray Data Collection and Refinement Statistic for the Complex Structure

data collection	CNS
space group	$P2_1$
cell dimensions	
a (Å)	45.116
b (Å)	106.595
c (Å)	47.680
a (deg)	90
β (deg)	102.034
γ (deg)	90
resolution (Å)	20-1.52
no. reflections	93 642
R _{merge} (%)	8.8
$I/\sigma I$	20.3
completeness (%)	95.9
redundancy	1.9
resolution range for refinement (Å)	19.39-1.50
$R_{\rm work}/R_{\rm free}$ (%)	15.94/18.17
no. of protein residues (including double conformations)	540
no. of water molecules	773
B factors (Å ²)	
protein atoms; molecule 1 and 2	10.2
protein atoms molecule 2	17.063
rmsd bond length (Å)	0.006
rmsd bond angles (deg)	1.313

1.50 Å resolution were refined using the maximum likelihood method in CNS, including simulated annealing with an initial temperature of 2000 K, positional minimization, and individual B-factor refinement, with a bulk solvent correction.⁴⁸ σ A-weighted electron density maps were calculated using CNS and used in the steps of manual model rebuilding with the program Coot.⁴⁹ The final cycle of refinement was performed using the software Refmac from the CCP4 suite.^{50,51} Crossvalidation was employed throughout, and 5% of the data were used for the $R_{\rm free}$ calculation. The stereochemical quality of the models was monitored periodically with the program Procheck.^{52,53}

ASSOCIATED CONTENT

Accession Codes

The coordinates and structure factors for the binary complex of CTX-M-9-compound **5** have been deposited in the Protein Data Bank with the accession code 4LEN.

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Notes

The authors declare no competing financial interest.

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ABBREVIATIONS USED

BZB2THB, Benzo[b]-thiophene-2-boronic acid; BL, β -lactamase; DPA, double-perturbation analysis; PDB, Protein Data Bank; THF, tetrahydrofuran; TLC, thin-layer chromatography

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