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Articles

# Comparative Study of Adenosine Analogs as Inhibitors of Protein Arginine Methyltransferases and a *Clostridioides difficile*-Specific DNA Adenine Methyltransferase

Jujun Zhou,  $\bigtriangledown$  Youchao Deng,  $\bigtriangledown$  Iredia D. Iyamu, John R. Horton, Dan Yu, Taraneh Hajian, Masoud Vedadi, Dante Rotili, Antonello Mai, Robert M. Blumenthal, Xing Zhang, Rong Huang,\* and Xiaodong Cheng\*



**ABSTRACT:** S-Adenosyl-L-methionine (SAM) analogs are adaptable tools for studying and therapeutically inhibiting SAMdependent methyltransferases (MTases). Some MTases play significant roles in host-pathogen interactions, one of which is *Clostridioides difficile*-specific DNA adenine MTase (CamA). CamA is needed for efficient sporulation and alters persistence in the colon. To discover potent and selective CamA inhibitors, we explored modifications of the solvent-exposed edge of the SAM adenosine moiety. Starting from the two parental compounds (**6e** and 7), we designed an adenosine analog (**11a**) carrying a 3phenylpropyl moiety at the adenine N6-amino group, and a 3-(cyclohexylmethyl guanidine)-ethyl moiety at the sulfur atom off the ribose ring. Compound **11a** ( $IC_{50} = 0.15 \ \mu M$ ) is 10× and 5× more potent against CamA than **6e** and 7, respectively. The structure of the CamA–DNA–inhibitor complex revealed that **11a** adopts a U-shaped conformation, with the two branches folded toward each other, and the aliphatic and aromatic rings at the two ends interacting with one another. **11a** occupies the entire hydrophobic surface (apparently unique to CamA) next to the adenosine binding site. Our work presents a hybrid knowledge-based and fragment-based approach to generating CamA inhibitors that would be chemical agents to examine the mechanism(s) of action and therapeutic potentials of CamA in *C. difficile* infection.

# INTRODUCTION

S-Adenosyl-L-methionine (SAM, SAMe, or AdoMet)<sup>1</sup> is a metabolite that is biosynthesized naturally in almost all species (excepting only some obligate intracellular parasites that import it) and is the second most-used cofactor after ATP.<sup>2</sup> SAM is involved in a wide range of biochemical reactions including (but not limited to) transmethylation reactions,<sup>3,4</sup> radical SAM reactions,<sup>5–8</sup> polyamine biosynthesis,<sup>9</sup> and SAM-sensing riboswitches. <sup>10,11</sup> SAM-dependent methyltransferases (MTases) catalyze methyl transfers onto a wide variety of substrate molecules, including both macromolecules (DNA, RNA, polysaccharides, lipids, and proteins) and micromolecules (such as catechol, arsenic, histamine, nicotinamide, and thiopurine). Given the significance of epigenetic methylation of macromolecules in cancer, immunity, cardiovascular and many other diseases, as well as methylation of anticancer drugs

such as thiopurine,<sup>12,13</sup> SAM-dependent MTases are increasingly becoming therapeutic targets.<sup>14–19</sup>

SAM analogs are versatile tools for studying and therapeutically inhibiting SAM-dependent MTases.<sup>20–22</sup> Some of these compounds as inhibitors of human epigenetic enzymes,<sup>23,24</sup> as well as inhibitors of viral RNA MTases from Zika virus and SARS-CoV-2,<sup>25–27</sup> are in clinical trials. Interestingly, some SAM analogs designed as selective inhibitors of human protein arginine MTases (PRMTs)<sup>24,28,29</sup> and human DOT1L (a

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**Figure 1.** CamA inhibition by a pan-PRMT inhibitor (**6e**). (A) Chemical structures of five SAM-analogs. The  $IC_{50}$  and  $K_D$  values against CamA, as well as PDB codes for the structures of CamA–DNA–inhibitor complexes, are extracted from Zhou et al.<sup>31</sup> (B) Comparative inhibition of activity of CamA at a single concentration of 10  $\mu$ M. These assays included 50 nM CamA and 40  $\mu$ M SAM. (C) Chemical structures of compounds **6a–6n** carrying substituents at the R group, taken from Iyamu et al.<sup>48</sup> [**6j** and **6k** in ref 48 were no longer available at the time of testing.] (D, E)  $IC_{50}$  values of **6e** against CamA (panel D) or PRMT1 and PRMT3 (panel E).

histone H3 lysine 79 MTase)<sup>30</sup> also inhibited the *Clostridioides* difficile-specific DNA adenine MTase (CamA) in vitro at low micromolar concentrations<sup>31</sup> (Figure 1A). CamA exists in the genomes of all *C. difficile* sequenced to date, but closely related orthologs have not yet been found in any other bacteria.<sup>31,32</sup> CamA is needed for normal sporulation and for persistent *C. difficile* infection (CDI) in an animal model.<sup>32</sup> CDI is a leading cause of community-associated infection in children and of hospital-acquired diarrhea in older patients,<sup>33,34</sup> can have lethal consequences, and has been identified as an urgent threat to human health (https://www.cdc.gov/drugresistance/biggest-threats.html).

We showed that CamA<sup>35</sup> shares a conserved SAM binding site with PRMTs<sup>36–39</sup> and DOT1L<sup>40</sup> and many other Rossman-fold Class I MTases.<sup>41,42</sup> Given that similarity, CamA has an unexpectedly low binding affinity for the methyl donor SAM  $(K_m > 17 \ \mu\text{M})$ .<sup>35</sup> For comparison, most other MTases have much higher SAM affinity; for example, the *E. coli* DNA adenine MTase Dam has a SAM  $K_m$  of  $3-6 \ \mu\text{M}$ ,<sup>43–45</sup> for human PCIF1 it is  $0.7 \ \mu\text{M}$ , and for MettL5 it is  $1.0 \ \mu\text{M}$ .<sup>46</sup> The unusually low SAM affinity of CamA might be related to a coupled conformational rearrangement of the two N-terminal helices.<sup>35</sup> Whatever the explanation may prove to be, this unusual property triggered us to investigate if SAM analogs are specifically potent inhibitors of CamA activity in competition with the weakly bound SAM cosubstrate.

In the present study, we took advantage of recent advances in development of molecular tools and in the growing number of clinically used compounds capable of potently and selectively inhibiting PRMTs.<sup>47</sup> These PRMT inhibitors can work as competitors of SAM or substrate polypeptide or allosteric inhibitors.<sup>17</sup> We started from a small set of pan-PRMT inhibitors<sup>48</sup> together with our recently developed CamA-specific adenosine analogs.<sup>49</sup> The combination of unique moieties in these two chemotypes led us to design compound **11a** (YD905), which represents a more potent lead compound against CamA, having a half-maximal inhibitory concentration (IC<sub>50</sub>) of ~0.15  $\mu$ M.

# RESULTS

CamA Inhibition by a Pan-PRMT Inhibitor. We previously determined that a few known compounds showed in vitro inhibition of CamA at low micromolar concentrations.<sup>31</sup> These included the PRMT5 inhibitor [N]-64619178 (1),<sup>24</sup> the PRMT7 inhibitor SGC8158 (2),<sup>29</sup> and the DOT1L inhibitor SGC0946 (3).<sup>30</sup> These compounds are all SAM-competitive inhibitors, all contain an adenosyl moiety (Figure 1A), and all of them occupy the individual enzymes' SAM binding site. The three compounds demonstrated improved potency of CamA inhibition relative to sinefungin (4), a pan inhibitor of SAMdependent MTases, starting from IC<sub>50</sub> values of 24  $\mu$ M for sinefungin and progressively decreasing to 18  $\mu$ M (JNJ-64619178), 7.1 µM (SGC8158), and 4.7 µM (SGC0946). We also made use of the observation that LLY-283 (5), a SAM analog developed as a PRMT5 inhibitor,<sup>50</sup> has the shortest branch off its ribose moiety compared to compounds 1-3 and displayed much weaker inhibition of CamA<sup>31</sup> (Figure 1B),



**Figure 2.** Compound **11a** (YD905) is competitive with SAM. (A) Chemical structures of **6e** (II745), 7 (MC4741), **11a** (YD905), and **11b** (YD907). **11a** and **11b** differ in the length of the linker (2C vs 3C) off of the sulfur atom. (B) Structure of CamA–DNA in complex with compound 7 (magenta) (PDB 8CXZ). The 3-phenylpropyl moiety at the N6-amino of adenosine is bound to a largely hydrophobic surface (colored in gray). (C) Summary of inhibition (IC<sub>50</sub> values) of CamA, PRMT1, and PRMT3 by the four compounds shown in panel A (see Figure S1 in Supporting Information). (D) Structure of PRMT3 in complex with SAH (yellow), which is completely enclosed by PRMT3 residues (PDB 1F3L). (E) Michaelis–Menten saturation curves of CamA kinetics varying SAM concentrations (20, 30, 40, and 80  $\mu$ M) and inhibitor **11a** concentrations (0, 0.25, and 0.5  $\mu$ M). (F) Lineweaver–Burk double reciprocal plot suggests competitive inhibition by **11a** with respect to SAM.

suggesting a path to explore in further improving these inhibitors.

A CamA inhibition screen was carried out using a set of 12 compounds designed as bisubstrate analogs of pan-PRMT inhibitors. These compounds join a thioadenosine to various substituted guanidinium groups (which mimic the target arginine side chain of protein substrates), through a hydrophobic propyl linker (Figure 1C).<sup>48</sup> The 12 bisubstrate compounds vary at the guanidium end, having either an aliphatic or an aromatic group (Figure 1C). Among them, compound 6e (II745), which contains an aliphatic ring, most potently inhibited ~85% activity of CamA at 10  $\mu$ M concentration and in the presence of 40  $\mu$ M SAM (Figure 1B). We made the following two observations: (i) eliminating the one-carbon linker between the aliphatic ring and guanidine group (from 6e to 6d) decreased CamA inhibition (from ~85% to  $\sim$ 60%); and (ii) substituting the aliphatic ring in **6e** with the aromatic analog in 6g nearly eliminated inhibitory activity ( $\sim$ 10% inhibition). These two features, the one-carbon linker and aliphatic ring, might contribute to the flexibility of compound 6e.

We compared the inhibitory activity of compound **6e** for CamA with PRMT1 and PRMT3. With an IC<sub>50</sub> value of 1.5  $\mu$ M, **6e** has 12.5× weaker inhibition of CamA than of PRMT1 (0.12  $\mu$ M), but >4× stronger inhibition than of PRMT3 (6.5  $\mu$ M)

(Figure 1D-1E). Nevertheless, **6e** demonstrated >3× improved potency of CamA inhibition relative to SGC0946 (4.7  $\mu$ M).

Design and Synthesis of Compounds 11a and 11b. To further develop more potent inhibitors of CamA, we introduced a substituent onto the amino group (N6) of adenosine (Figure 2A). The rationale of this design stemmed from our recent structural characterization of CamA-DNA complexes in the presence and absence of methyl donor cofactor (SAM or its end product, S-adenosyl-L-homocysteine, SAH), which revealed a unique solvent-exposed edge of the SAM/SAH adenosine moiety (Figure 2B).<sup>31,35</sup> The N6-modified compounds bind in the pocket normally occupied by the SAM adenosine, with the N6-addition directed outward and resting against a mostly hydrophobic surface (see Discussion), making a unique mixture of interactions with CamA involving van der Waals, aromatic, and/or polar contacts.<sup>49</sup> Among them, compound 7 (MC4741) contains a 3-phenylpropyl moiety at the N6-amino group of adenosine and demonstrated an IC<sub>50</sub> value of 0.7  $\mu$ M against CamA<sup>49</sup> [Note compound MC4741 was numbered as 14 in ref 49]. Importantly, for potential therapeutic use in humans, compound 7 does not inhibit either PRMT1 or PRMT3 (Figures 2C and S1). Significantly, in PRMT3 there is no equivalent solvent-exposed surface next to the SAM/SAH adenosine moiety (Figure 2D; see Discussion on PRMT1). We reasoned that combining the unique features of 6e and 7 could lead to synergistic CamA inhibition.



**Figure 3.** Structure of CamA–DNA–**11a** complex. (A) Omit  $F_o - F_c$  electron densities of **11a** and **11b** contoured at  $3\sigma$  above the mean, from PDB 8FS1 and PDB 8FS2, respectively. (B) Structure of CamA–DNA with bound compound **11a** (in stick) within a dashed green box (PDB 8FS1). Surface representation is shown with carbon atoms (light gray), oxygen atoms (red), and nitrogen atoms (blue). The bound DNA substrate is shown in a cartoon model. (C) Enlarged solvent-exposed surface where **11a** (in green) binds. The surface is formed by aromatic residues Y30 (left), Y178 (right), and I115 (front), hydrophobic residues P167, I169, L174, L196, and F200 in the back (see panel H), and acidic residues (D149 and E174). In this view, the **11a** adenosine is partially obscured by I115. (D) Superimposition of CamA-bound compound **11a** (green; PDB 8FS1), SAH (PDB 7LT5), or sinefungin (yellow; PDB 7RFK), highlighting the U-shape of **11a**. (E) The aromatic side chain of Y30 conformationally changes between the SAH/sinefungin-bound state (yellow) and the **11a**-bound state (green). (F) The N-terminal helices ( $\alpha$ A and  $\alpha$ B) are disordered in the **11a**-bound state. (G, H) Compound **11a** contacts Y30 (left), E175, Y178, and D149 (right), and hydrophobic residues in the back (panel H). (I) The hydrophobic side chain of **116**9 packs against the aliphatic ring and the aromatic ring located, respectively, at the two ends of compound **11a** (green; PDB 8FS1) and compound **7** (magenta; PDB 8CXZ). (L, M) Two views of superimposition of three CamA-bound inhibitors, **11a** (green; PDB 8FS1), GSK0946 (cyan; PDB 7RFL), and sinefungin (yellow; PDB 7RFK).

We thus designed and synthesized two new compounds, **11a** (YD905) and **11b** (YD907), respectively, having a 2-carbon ethyl or 3-carbon propyl linker between the guanidinium group and the sulfur atom. Compound **11a** exhibited enhanced inhibition against CamA relative to the two parental compounds, with either 10× improvement from **6e** (IC<sub>50</sub> of 0.15  $\mu$ M vs. 1.5  $\mu$ M) or 5× improvement from 7 (IC<sub>50</sub> of 0.15  $\mu$ M vs. 0.7  $\mu$ M) (Figure 2C). However, increasing the linker from 2-carbon in **11a** to 3-carbon in **11b** resulted in 1.8× reduced inhibition of CamA relative to **11a** (i.e., the IC<sub>50</sub> value increased from 0.15  $\mu$ M to 0.27  $\mu$ M).

As expected, both **11a** and **11b** exhibit reduced inhibition of PRMT1 and PRMT3, as one of the parental compounds (7) does not significantly inhibit either of the two PRMTs. **11a** has a remarkable reduction (~116×) in inhibition of PRMT1 from that of **6e** (IC<sub>50</sub> of 14  $\mu$ M vs. 0.12  $\mu$ M) (Figure 2C). The same magnitude of negative effect on PRMT3 activity made the inhibition unmeasurable (IC<sub>50</sub> > 100  $\mu$ M). However, compound

**11b** has a much milder effect on PRMT1 inhibition than **6e**, resulting in a 10× reduction of PRMT1 inhibition (IC<sub>50</sub> increased to 1.2  $\mu$ M from 0.12  $\mu$ M), instead of the >100× reduction by **11a**. The lesser reduction of PRMT1 inhibition by **11b** is probably owing to the 3-carbon linker, given the reported optimal length between the thioadenosine and guanidinium group for PRMT inhibition by bisubstrate analogs.<sup>51</sup>

To further understand the mode of CamA inhibition by compound **11a**, we measured the inhibition constant ( $K_i = 90$  nM under our experimental conditions) by varying SAM concentration in the absence and presence of inhibitor at two concentrations (Figure 2E). A Lineweaver–Burk plot suggests that compound **11a** is a competitive inhibitor of SAM, as expected (Figure 2F). In agreement with our previous measurements ( $K_m > 17 \ \mu M$ ),<sup>35</sup> CamA has a weak binding affinity (as reflected by the  $K_m$  value) of 25  $\mu M$  for SAM (Figure 2E).



**Figure 4.** Selectivity of compounds **6e**, **7**, **11a**, and **11b** against activities of (A) CamA, (B) CcrM, (C) Dam, (D) MettL3–MettL14, (E) MettL5– Trm112, (F) MettL16, and (G) PCIF1. The oligonucleotide substrates and reaction conditions are listed in Table S2. Compound 7 used in the selectivity experiments was from ref 49. (H) Hydrophobic patch in CamA. The sequence logo shows that the eight residues forming this surface patch are very highly conserved among 58 nonidentical CamA orthologs, but they are not conserved among other DNA/RNA adenine MTases. The catalytic motif IV (D/N)PP(Y/F/W) is highlighted in yellow, the relevant P within the motif in green, and the other patch residues in cyan. In the non-CamA MTases, aligned different (but still hydrophobic) residues are shown in gray, and charged residues are shown in red circles.

**Compound 11a Binds to a Solvent-Exposed, Largely Hydrophobic Surface.** To define the basis for the enhanced binding and inhibition of CamA by compounds **11a** (YD905) and **11b** (YD907) relative to the parent compounds, we cocrystallized the CamA–DNA complex with each of the two compounds. The two complexes both crystallized in space group  $P2_12_12_1$ , and very similar structures (with root-mean-square deviation of 0.7 Å) were determined to resolutions of 2.74 Å (**11a**) and 2.59 Å (**11b**) (Table S1). While **11a** was fully observed in the electron density, the portion of **11b** where the 3carbon linker located was partially disordered (Figure 3A). In the following section, we describe CamA interactions with **11a** (Figure 3B), with a focus on the interactions involving the two branches off the guanidino group and the amino group of adenosine.

The adenosyl moiety (adenine and ribose rings) of compound 11a occupies the exact same binding site as the corresponding moieties of SAH and sinefungin (Figure 3C,D). However, unlike the homocysteine moiety of SAH and sinefungin, which exhibits an extended conformation, 11a adopts a U-shaped conformation, with the two branches folded toward each other, yielding a 3.5 Å separation between the aliphatic ring off the guanidinium group and the aromatic ring off the adenosine (Figure 3D). This separation of 3.5 Å reflects the optimum distance between the two ring carbon atoms with a van der Waals radius of 1.7 Å. Compound 11a is bordered by two tyrosine residues, Y30 adjoining the guanidinium branch and Y178 next to the adenosine branch (Figure 3D). Y30, one of the two residues that sandwich the target DNA adenine, rotates its aromatic ring  $\sim 90^{\circ}$  toward the guanidinium group (Figure 3E). The binding of compound 11a also resulted in disorder of the two N-terminal helices (residues 1-28), owing to the collision between the guanidinium branch and the helix  $\alpha B$  (Figure 3F).

The two branches of **11a** form extensive interactions with CamA and the bound substrate DNA target adenine (Ade), via a mixture of interactions involving van der Waals, aromatic, and

polar contacts (Figure 3G,H). The branch off the guanidinium group contacts Y30, the target Ade, P167, as well as I169 (Figure 3H). The hydrophobic side chains of I169 and L174 pack against the aliphatic ring and the aromatic ring located at the ends of the two branches of 11a (Figure 3H,I). The sulfur atom interacts with the DNA target Ade N6 amino group (Figure 3J). The 3-phenylpropyl moiety attached to the Ade N6-amino of 11a has the same conformation as that of compound 7 (Figure 3K). In sum, compound 11a has a  $\sim$ 520 A<sup>2</sup> interface with CamA  $(>100 \text{ A}^2 \text{ larger than that of compound 7})$ , occupying nearly the entire hydrophobic surface as shown in Figure 3C. As oriented in Figure 3C, the hydrophobic surface is formed by Y30 (left), Y178 (right), I115 (front), and P167, I169, L174, L196, and F200 in the back (Figure 3H). Finally, superimposition of three CamA-bound inhibitors, sinefungin (4), SGC0946 (3), and 11a, indicates that the extension off the ribose ring points in three different directions and engages in three different sets of interactions (Figure 3L,M). The 11a-mediated interactions involve the predominantly hydrophobic surface and help to explain its most potent inhibition by 11a.

Selectivity. Next, we examined four compounds, two parental compounds (6e and 7) and two derived compounds (11a and 11b), against a collection of related SAM-dependent MTases acting on nucleic acids (Figure 4), containing Escherichia coli Dam, Caulobactor vibrioides CcrM, and four human RNA (or dual RNA/DNA) adenine MTases (MettL3-MettL14, MettL5-Trm112, MettL16, and PCIF1) that have been enzymatically characterized in our laboratory.<sup>46,52</sup> While these four compounds showed robust ( $\geq$ 80%) inhibition of CamA activity (Figure 4A) at 10  $\mu$ M concentration in the presence of 40  $\mu$ M SAM, none of them showed >50% inhibition of the other tested MTases (Figure 4B–G). Compound 6e came close to the 50% threshold with complexes of MettL3-MettL14 and MettL5-Trm112 (Figure 4D,E), and 11b was close for MettL5–Trm112 (Figure 4E), but the difference from CamA inhibition was dramatic. As noted above, compound 6e was



**Figure 5.** Models of compounds **6e** or **11b** in binding MettL3, PRMT1, and MettL5. (A) Structures of MettL3–MettL14 bound with two inhibitors (PDB 6TTW and PDB 6TTX). (B) A model of **6e** binding MettL3 by superimposing the adenosine moiety. (C) Structure of PRMT1–SAH (PDB 1ORI) where the SAH binding site is exposed while the N-terminal residues are disordered in the absence of bound arginine substrate (circled). (D) Structure of PRMT1–SAH–GSK3368715 (PDB 6NT2) where SAH is largely buried in the presence of substrate-competitor inhibitor. The arrow indicates a narrow opening near the adenine N6-amino group. (E) The distance is 3.9 Å between the sulfur atom of SAH and the GSK inhibitor which occupied the substrate arginine binding site. (F) A model of compound **11b** binding PRMT1 in the open conformation. (G, H) A model of compound **11b** binding PRMT1 in the N6-addition. (I) Structure of MettL5–Trm112 in complex with SAM (PDB 6H2U). A model of **11e** binding MettL5 by superimposing the adenosine moiety, and the aliphatic and aromatic rings (circled) are visible from the surface.

developed as a pan-inhibitor of PRMT enzymes,<sup>48</sup> and it does inhibit CamA activity with an IC<sub>50</sub> of 1.5  $\mu$ M (this study). Thus, it is not too surprising that **6e** also modestly inhibits the activity of MettL3–MettL14 (with an estimated IC<sub>50</sub> > 10  $\mu$ M), as analogs of adenosine with *N*-substituted amide of ribofuranuronic acid off the ribose ring have been developed as inhibitors of MettL3–MettL14.<sup>59</sup>

As noted earlier, and discussed below, CamA differs from the other MTases in having a hydrophobic surface patch adjacent to the adenosine-binding pocket, and the N6-linked substituent in compounds 7, 11a, and 11b lies against this patch. The patch is conserved among CamA orthologs but is not seen in the other MTases (Figure 4H). For comparison, MettL16 has hydrophobic residues in five of the six aligned positions forming the CamA patch (Figure 4H), yet 11a shows just ~25% inhibition (Figure 4F), possibly because a negative charged glutamate is in the place of L174. 11a has about the same effect on CcrM (Figure 4B), which has hydrophobic residues in just two of the

six positions and also has a positively charged arginine in the place of Y178. *E. coli* Dam has no charged residues at the CamA hydrophobic patch positions, but has polar threonine residues at two of those positions, and shows about the same 25% inhibition as MettL16 and CcrM (Figure 4C).

# DISCUSSION

The majority (if not all) of SAM-dependent methyltransfer reactions involve a direct transfer of the methyl group from the charged methylsulfonium center to substrate, with an  $S_N$ 2-like mechanism.<sup>60</sup> The close juxtaposition between the sulfur atom of SAM (the methyl donor) and substrate (the methyl acceptor) allowed design and synthesis of bivalent (or bisubstrate) inhibitors<sup>61</sup> that have an effective linker between the two fragments, thus mimicking the transition state of the methyl transfer reaction. Compound **6e** is one such example, and it inhibits PRMT1 (IC<sub>50</sub> = 0.12  $\mu$ M), CamA (IC<sub>50</sub> = 1.5  $\mu$ M), PRMT3 (IC<sub>50</sub> = 6.5  $\mu$ M) and MettL3–MettL14 (IC<sub>50</sub> > 10



"Reagents and conditions of analog synthesis: (a) *n*-butanol, 3-phenylpropylamine, reflux, 12 h, 83%; (b) thioacetic acid, triphenylphosphine, diisopropyl azodicarboxylate, tetrahydrofuran, 0 °C to room temperature, overnight, 71%; (c) sodium methoxide, MeOH, room temperature, overnight; (d) HCOOH, H<sub>2</sub>O, 0 °C to room temperature, 2 d, 75–77% in two steps; (e) 1-ethyl-3-(3-dimethylaminopropyl)carbodiimide, *N*,*N*-diisopropylethylamine, dichloromethane, room temperature, overnight; (f) piperidine, dichloromethane, room temperature, 30 min, 53–57% in two steps.

 $\mu$ M). The common feature of the substrates of these enzymes is the amino group (NH<sub>2</sub>) of protein arginine (N $\epsilon$ ) or DNA/RNA adenine (N6). Variation in these bivalent inhibitors usually involves the extensions off the ribose ring, because of where the methylsulfonium atom is located. We modeled compound **6e** onto the existing structures of MettL3–MettL14 complexed with known adenosine analogs having varied IC<sub>50</sub> values<sup>59</sup> (~9  $\mu$ M to >250  $\mu$ M) (Figure 5A). It is evident that **6e** could be easily accommodated in the surface groove extended from the adenosine binding site (Figure 5B).

On the other hand, Ade N6-substituted SAH/SAM analogs have also been explored, initially targeted toward a mutant of the *S. cerevisiae* PRMT called RMT1.<sup>62</sup> Recently we characterized 42 adenosine analogs as potential inhibitors of CamA, which incorporated substituents at the adenosine C6-amino group (N6), including arylalkyl groups with a growing carbon-number alkyl chain, or variously substituted aminoalkyl groups.<sup>49</sup> Among them, compound 7 contains a 3-phenylpropyl moiety with the three-carbon aliphatic chain connecting the Ade N6 atom with the phenyl ring, and which inhibits CamA with an IC<sub>50</sub> of 0.7  $\mu$ M. However, compound 7 inhibits neither PRMT1 nor PRMT3 (Figure S1), nor does it significantly reduce the activities of a few other tested DNA/RNA adenine MTases (Figure 4).

Combining features of 6e and 7 gave rise to compounds 11a and 11b, with extensions projecting in two directions from ribose and adenine, respectively. We show here that 11a and 11b have synergistic effects on specificity, increasing inhibition of CamA and at the same time decreasing that of PRMT1 (Figure 2C). In addition to the direction of the effects differing between 11a and 11b on CamA and PRMT1, the strength of these respective effects also differs, which might be attributed to the length of hydrophobic linker joining the thioadenosine and guanidinium group. For CamA, compound 11a has 1.8× increased inhibition of CamA relative to 11b; the 3-carbon length in 11b might disturb the optimal distance between the aliphatic ring and the aromatic ring seen in 11a, and resulted in a partially disordered aliphatic ring (Figure 3A). For PRMT1, both compounds showed reduced inhibition from that of the parental compound **6e**; but compound **11b**, with an  $IC_{50}$  value of 1.2  $\mu$ M under the conditions tested (Figure 2C), has >11× stronger inhibition than 11a, in agreement with a previous study showing that the 3-carbon linker is preferred for the PRMT bisubstrate inhibitors.<sup>51</sup>

PRMT1–SAH structures have been determined to be in an open conformation in the absence of bound substrate<sup>37</sup> (Figure

5C) and in a closed conformation in the presence of a substratecompetitive inhibitor<sup>63</sup> (Figure 5D,E). While **6e** and **11b** could be easily modeled onto the open conformation (Figure 5F), the 3-carbon optimum length in **11b** for PRMT1<sup>51</sup> might force PRMT1 to undergo conformational change to accommodate the substituent at the N6-amino group (i.e., to open up a narrow opening near the N6-amino group indicated by an arrow in Figure 5D,G). Compound **11b** links a portion of SAH to a part of the substrate-competitive inhibitor (Figure 5H). In addition, compound **11b**, which at a concentration of 10  $\mu$ M showed close to 50% inhibition of MettL5–Trm112 (Figure 4E), could be modeled with two branches occupying the SAM and target adenine binding sites, respectively (Figure 5I).

Here, we present compound **11a** [1-(cyclohexylmethyl)-3-(2-((((2S,3S,4R,5R)-3,4-dihydroxy-5-(6-((3-phenylpropyl)amino)-9H-purin-9-yl)tetrahydrofuran-2-yl)methyl)thio)ethyl)guanidine] as a selective and potent CamA inhibitor. Like SAM/SAH, the inhibitor occupies the adenosine binding site with the N6-addition and the S-linked substitution directed outward and resting against a mainly hydrophobic surface, engaging in a mixture of interactions involving van der Waals, aromatic, and polar contacts (Figure 3C). The compound fully occupied the entire pocket-adjacent hydrophobic surface that appears to be unique to CamA among amino MTases examined in this study. We suggest that this hydrophobic surface patch contributes to CamA specific targetability.

To further improve our lead compound's ability, we will turn our attention to the space occupied by the homocysteine moiety of SAH/sinefungin, which points in a different direction from that of the S-linked substitute in **11a** (Figure 3D). A branched extension at or near the sulfur atom could be designed to merge **11a** and sinefungin or **11a** and SGC0946 (Figure 3L,M). For example, derivatives of AzaSAM, in which the nitrogen that replaces the SAM sulfur atom is coupled to various alkylamino groups, were developed as inhibitors of Set7/9 (a protein lysine MTase).<sup>64</sup> In sum, our work on discovering CamA inhibitors would specifically add to continuing exploration for other antimicrobial and/or antivirulence agents targeting *C. difficile* and could yield synergistic therapeutic approaches in clinical settings.

#### METHODS

**Chemistry.** Starting materials, reagents, and solvents were purchased commercially (Sigma Chemical Co. and Ambeed Inc.). Analytic and preparative high-pressure liquid chromatography (RP-HPLC) were performed on an Agilent 1260 Series system, which was run with a 5–95% acetonitrile/water gradient with 0.1% trifluoroacetic

acid (TFA). Flash chromatography was performed on a Teledyne CombiFlash Rf 100 system, run with a 0–5% methanol/dichloromethane gradient. The reaction yields were estimated based on the weight of the desired products. All <sup>1</sup>H and <sup>13</sup>C NMR spectra were recorded on a Bruker 500 MHz spectrometer. Based on HPLC analysis, all compounds used for IC<sub>50</sub> determination possessed a purity of >95%. High-resolution mass spectrometry (HRMS) spectra were logged on an Agilent high resolution 6550 quadrupole time-of-flight (Q-TOF) LC-MS instrument.

The desired compounds **11a** and **11b** were synthesized following the reported methods<sup>48</sup> (Scheme 1). Briefly, the commercially available 6-chloropurine ribonucleoside reacted with 3-phenylpropylamine to yield N6-phenylpropyladenosine (7), which was then subjected to the Mitsunobu reaction to produce compound **8**. Then the acetyl group of **8** was removed, followed by its substitution with Boc protected ethyl or propyl amine. Next, the Boc group was removed in the presence of 50% formic acid in water. Under vacuum, the volatiles were removed, and the residues were lyophilized to produce compounds **9a** and **9b** as a pale-yellow solid. Compound **10** was made as reported in quantitative yields,<sup>48</sup> and was then subjected to the reaction with **9a**–**9b**, followed by final Fmoc deprotection to generate **11a**–**11b**.

Synthesis of Compound 7. 6-Chloropurine ribonucleoside (2.86 g, 10.0 mmol) and 3-phenylpropylamine (6.76 g, 50.0 mmol), in 200 mL of anhydrous *n*-butanol, were stirred under reflux for 12 h. After cooling down to RT and under reduced pressure, the volatiles were removed, and the residue was purified by flash chromatography (DCM/MeOH, 95/5) to give 7 (3.20 g, yield 83%).

Synthesis of Compound 8. In a chilly solution of triphenylphosphine (2.62 g,10.0 mmol) in dry THF (100 mL), diethyl azodicarboxylate (1.6 mL, 10 mmol) was added drop by drop over 5 min and stirred for 30 min. Then 7 (1.92 g, 5.0 mmol) was added and continued to stir for 2 h. To the resulting yellow suspension was slowly added a solution of thioacetic acid (0.7 mL, 10.0 mmol) in dry THF (5 mL); the mixture was stirred overnight while the temperature increased slowly to RT. The crude oil was purified by flash chromatography (DCM/MeOH, 97/3) to yield 8 (1.57 g, yield 71%), after the removal of volatiles under reduced pressure.

Synthesis of Compounds **9a** and **9b**. Under an argon atmosphere, to a stirring solution of compound **8** (445 mg, 1.0 mmol) and *tert*-butyl (2-bromoethyl)carbamate or *tert*-butyl (3-bromopropyl)carbamate (1.5 mmol) in dry methanol (20 mL) was added sodium methoxide (30% in methanol, 0.6 mL, 3.0 mmol). The volatiles were removed under reduced pressure, after stirring overnight at RT, and the residue was purified by flash chromatography (DCM/MeOH, 97/3). The fractions were collected and dried under vacuum. The generated oil was dissolved in water (5.0 mL) and formic acid (5.0 mL) at 0 °C and stirred at RT for 2 days. Under reduced pressure, the volatiles were removed and the residues were lyophilized to afford the desired product **9a** (333 mg, yield 75%) and **9b** (352 mg, yield 77%), each as a pale-yellow solid.

Synthesis of Thiourea (10). To a stirring solution of Fmocisothiocyanate (281 mg, 1.0 mmol) in 25 mL at 0 °C, DCM was slowly added cyclohexylmethanamine (113 mg, 1.0 mmol). The reaction mixture was stirred for 1 h at RT. The volatiles were removed under reduced pressure to yield crude 10, which was directly used for the reaction.

Synthesis of Compounds 11a and 11b. To a stirring solution of amine 9a or 9b (0.1 mmol) in 10 mL DCM and thiourea 10 (59 mg, 0.15 mmol) were added 1-ethyl-3-(3-(dimethylamino)propyl)-carbodiimide (23 mg, 0.15 mmol) and *N*,*N*-diisopropylethylamine (19 mg, 0.15 mmol). The reaction mixture was stirred at RT overnight and was then diluted with 50 mL water and extracted with dichloromethane ( $3 \times 20$  mL). The combined organic layers were washed with water and brine and dried with anhydrous sodium sulfate. The volatiles were removed under reduced pressure to afford a crude oil, which was subsequently treated with 20% piperidine in 10 mL dichloromethane for 30 min. Then, the reaction was concentrated and separated with preparative HPLC (MeCN/H<sub>2</sub>O, 10–40%) to yield 11a (31 mg, yield 53%) and 11b (34 mg, yield 57%).

**11a.** HRMS m/z calcd for  $C_{29}H_{42}N_8O_3S$  [M + H]<sup>+</sup>: 583.3173. Found: 583.3185. <sup>1</sup>H NMR (500 MHz, CD<sub>3</sub>OD)  $\delta$  8.53–8.32 (m, 1H), 8.30 (s, 1H), 7.26–7.15 (m, 4H), 7.15–7.09 (m, 1H), 6.02 (d, *J* = 4.9 Hz, 1H), 4.75 (s, 1H), 4.33 (t, *J* = 5.1 Hz, 1H), 4.25–4.17 (m, 1H), 3.58 (s, 2H), 3.38 (t, *J* = 6.6 Hz, 2H), 3.11–2.91 (m, 4H), 2.75 (ddd, *J* = 15.4, 10.5, 6.5 Hz, 4H), 2.03 (s, 2H), 1.81–1.59 (m, 5H), 1.52 (ddd, *J* = 11.4, 7.9, 4.0 Hz, 1H), 1.31–1.11 (m, 3H), 0.93 (tt, *J* = 12.1, 6.3 Hz, 2H). <sup>13</sup>C NMR (126 MHz, CD<sub>3</sub>OD)  $\delta$  156.20, 152.55, 149.45, 147.88, 141.23, 140.52, 128.06, 128.03, 125.61, 119.57, 88.96, 84.46, 73.64, 72.62, 40.76, 37.21, 33.87, 32.61, 31.43, 30.20, 25.98, 25.41.

**11b.** HRMS m/z calcd for  $C_{30}H_{44}N_8O_3S$  [M + H]<sup>+</sup>: 597.3330. Found: 597.3342. <sup>1</sup>H NMR (500 MHz, CD<sub>3</sub>OD)  $\delta$  8.32 (s, 1H), 8.28 (s, 1H), 7.30–7.17 (m, 4H), 7.16–7.10 (m, 1H), 6.01 (d, J = 4.9 Hz, 1H), 4.75 (s, 1H), 4.32 (t, J = 5.0 Hz, 1H), 4.25–4.16 (m, 1H), 3.58 (s, 1H), 3.25 (t, J = 6.9 Hz, 2H), 3.09–2.87 (m, 4H), 2.74 (dd, J = 8.6, 6.8 Hz, 2H), 2.63 (t, J = 7.1 Hz, 2H), 2.03 (t, J = 7.6 Hz, 2H), 1.84 (q, J = 6.9 Hz, 2H), 1.80–1.59 (m, 6H), 1.52 (dt, J = 7.8, 4.0 Hz, 1H), 1.35–1.10 (m, 4H), 0.94 (tt, J = 12.0, 6.3 Hz, 2H). <sup>13</sup>C NMR (126 MHz, CD<sub>3</sub>OD)  $\delta$  156.20, 153.23, 150.38, 141.33, 140.12, 128.04, 125.58, 119.50, 88.84, 84.30, 73.63, 72.65, 39.88, 37.28, 33.90, 32.65, 30.21, 29.21, 28.34, 25.99, 25.42.

**MTases Used in the Study.** The MTases used in the current study were characterized previously and prepared in our laboratories: *Clostridioides difficile* CamA (pXC2184),<sup>35</sup> *Caulobacter crescentus* [now called *C. vibrioides*] CcrM (pXC2121),<sup>65</sup> *Escherichia coli* Dam (pXC1612),<sup>52,53</sup> human PCIF1 (pXC2055),<sup>46,58</sup> human MettL16 (pXC2210),<sup>46</sup> human MettL5–Trm112 (pXC2062–pXC2076),<sup>46</sup> and human MettL3–MettL14.<sup>55,56</sup> Human PRMT1 (residues 10–352)<sup>66</sup> and human PRMT3 (residues 211–531)<sup>67</sup> were purified according to published procedures.

**Inhibition Assay on PRMT1 and PRMT3.** To measure the inhibitory effect of the compounds on MTase activities of PRMT1 and PRMT3, a fluorescence-based SAHH-coupled assay<sup>68,69</sup> was utilized. A BMG CLARIOstar microplate reader measured the fluorescence signal for 18 min with excitation at 400 nm and emission at 465 nm. All experiments were performed in duplicate, and GraphPad Prism software 8.0 was used to process data.

For PRMT1, the assay was performed in 100  $\mu$ L including 0.1  $\mu$ M PRMT1, 10  $\mu$ M SAM, 5  $\mu$ M SAHH, and 10  $\mu$ M ThioGlo4 in 2.5 mM HEPES (pH 7.0), 25 mM NaCl, 25  $\mu$ M EDTA, 50  $\mu$ M tris(2-carboxyethyl)phosphine (TCEP), 0.01% Triton X-100. For PRMT3, the assay was performed in 100  $\mu$ L including 0.25  $\mu$ M PRMT3, 30  $\mu$ M SAM, 5  $\mu$ M SAHH, and 15  $\mu$ M ThioGlo4 in 20 mM Tris (pH, 7.5), 0.01% Triton X-100. After incubation with the inhibitors for 10 min at 37 °C (PRMT1) or 30 °C (PRMT3), the addition of 5  $\mu$ M histone H4 peptide (residues 1–21) initiated reactions.

Inhibition Assay of CamA-Mediated Methylation. The CamA methylation assay was performed in 100 mM NaCl, 50 mM Tris-HCl, pH 7.5, 1 mM dithiothreitol (DTT), 0.25% DMSO, and 0.1 mg mL<sup>-1</sup> BSA. CamA (50 nM final concentration) was preincubated with varied concentration of compound at RT (~22 °C) for 5 min followed by addition of 2.5  $\mu$ M double-stranded DNA (final concentration) (Table S2) and 40  $\mu$ M SAM (final concentration) (Figures 1B,D and S1). To determine the  $K_i$  of compound 11a against CamA, the methylation reaction was carried out with varying concentrations of SAM (20, 30, 40, and 80  $\mu$ M) and inhibitor 11a (0, 0.25, and 0.5  $\mu$ M) (Figure 2E). Reactions were incubated for 3 min at ~22 °C (RT), and quenched by adding TFA to 0.1%. The final reaction mixture (5  $\mu$ L) was transferred to a low-volume 384-well plate, and the SAH concentration was measured by Promega luminescence assay (MTase-Glo).<sup>70</sup> A Synergy 4 Multi-Mode Microplate Reader (BioTek) was used to detect the luminescence signal.

**Compound Selectivity Assays.** Table S2 lists the MT ases used in the selectivity assays (Figure 4), their respective substrates, and the reaction conditions under 10  $\mu$ M inhibitor concentration. Integrated DNA Technologies (IDT) synthesized oligonucleotides, and the mRNA cap analog (catalog number N-7113) was purchased from TriLink BioTechnologies. Inhibition activities at RT (~22 °C) were measured by Promega luminescence assay (MT ase-Glo).

X-ray Crystallography. An Art Robbins Gryphon Crystallization Robot set up the sitting drop crystals screening by vapor diffusion method. Crystals of CamA–DNA–11a (or 11b) grew under similar conditions after 3–4 days at ~19 °C (RT): 0.28 M potassium citrate, 0.1 M Tris-HCl, pH 7.0–7.5, and 21–24% (w/v) polyethylene glycol 3350. Crystals were quickly soaked in reservoir solution supplemented with 20% (v/v) ethylene glycol and flash frozen by liquid nitrogen.

At Argonne National Laboratory, X-ray diffraction data were collected at the SER-CAT beamline 22ID of the Advanced Photon Source. Crystallographic data sets were indexed, integrated, and scaled with HKL2000.<sup>71</sup> The difference Fourier method was used to solve the structures of the ternary CamA–DNA–11a (or 11b) complex, using the binary CamA-DNA structure as a template (PDB 7LNJ). Crystals were isomorphous essentially. In the first refinement cycle, rigid body refinement was used for positioning the new structures in the unit cell, and difference electron density maps  $(2F_o - F_c \text{ and } F_o - F_c)$  were used for locating bound inhibitor molecules. The grade web server (http:// grade.globalphasing.org) was used to generate CIF file restraints for 11a and 11b and for model building and refinement. COOT<sup>72</sup> and PHENIX REFINE<sup>73</sup> were used respectively and iteratively for model building and refinement of the structure. Table S1 summarizes the statistics of diffraction data collection and refinement. The PDB validation server<sup>74</sup> was used for validation of the final structural models. PyMol version 2.5.2 (Schrödinger, LLC) was used to prepare structure images.

**Computer Modeling of Compounds 6e and 11b Binding in MettL3, MettL5, and PRMT1.** COOT<sup>75</sup> was utilized to superimpose the structure of 11b-bound CamA (PDB 8FS2) with structures of other MTases (MettL3,<sup>59</sup> MettL5,<sup>76</sup> PRMT1<sup>37,63</sup>) having bound SAM/SAH or compounds also containing an adenosine moiety. The secondarystructure matching (SSM)<sup>77</sup> and least-squares (LSQ) options were primarily used for the superimposition. SSM aligns proteins based on their topology which works fairly well with the seven-stranded Class I MTases,<sup>41</sup> and LSQ allowed additional fine-tuning of the superimposition. Small rotations and translations of **11b** were made such that the adenosine moiety of **11b** exactly overlaid with that moiety in the other MTase structures. Finally, the edit chi angle function in COOT allowed rotation of permissible torsional angles to place the extensions of **11b** so as to eliminate clashes with the MTase and permit potential interactions. The **6e**-bound MettL3 was generated by deleting the 3phenylpropyl moiety from **11b**.

Sequence Alignment of Residues Forming a Hydrophobic Patch in CamA. CamA is the only known DNA adenine MTase with a hydrophobic surface patch adjacent to the SAM-binding pocket. A WebLogo analysis<sup>78</sup> of 58 nonidentical CamA orthologs<sup>32</sup> shows that the eight hydrophobic residues forming this surface patch are very highly conserved among CamA orthologs (the upper panel of Figure 4H). However, these residues are not conserved among other DNA/ RNA adenine MTases, CcrM from Caulobacter vibrioides (formerly crescentus) (YP 002515757.1), Dam from E. coli (NP 417846.1), and four human RNA MTases, MettL3 (XP\_011535270.1) (the catalytic subunit of MettL3-MettL14), MettL5 (NP\_054887.2) (the catalytic subunit of MettL5-Trm112), MettL16 (NP 076991.3), and PCIF1 (NP 071387.1). Only the more carboxyl-proximal six of the eight patch-forming residues are shown in the lower panel of Figure 4H, as the sequences in that region can be unambiguously aligned using the conserved (N/D)PP(Y/F/W) motif  $IV^{79}$  as an anchor point.

# ASSOCIATED CONTENT

#### **Supporting Information**

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acschembio.3c00035.

Summary of X-ray data collection and refinement statistics (Table S1); Inhibition reaction conditions (Table S2); Raw data of  $IC_{50}$  plots (Figure S1); NMR spectra of compounds **11a** and **11b**; HR-MS spectra of compounds **11a** and **11b**; Purity of compounds **11a** and **11b** (PDF)

Molecular formula strings (SMILES) and associated data (XLSX)

# Accession Codes

The experimental data that support the findings of this study are contained within the article. The X-ray structure (coordinates) and the source data (structure factor file) of CamA–DNA with bound inhibitors have been deposited to the PDB, and these will be released upon article publication under accession numbers PDB 8FS1 (11a) and PDB 8FS2 (11b).

### AUTHOR INFORMATION

#### **Corresponding Authors**

- Rong Huang Department of Medicinal Chemistry and Molecular Pharmacology, Institute for Drug Discovery, Center for Cancer Research, Purdue University, West Lafayette, Indiana 47907, United States; o orcid.org/0000-0002-1477-3165; Email: huang-r@purdue.edu
- Xiaodong Cheng Department of Epigenetics and Molecular Carcinogenesis, University of Texas MD Anderson Cancer Center, Houston, Texas 77030, United States; orcid.org/ 0000-0002-6967-6362; Email: XCheng5@mdanderson.org

#### Authors

- Jujun Zhou Department of Epigenetics and Molecular Carcinogenesis, University of Texas MD Anderson Cancer Center, Houston, Texas 77030, United States
- Youchao Deng Department of Medicinal Chemistry and Molecular Pharmacology, Institute for Drug Discovery, Center for Cancer Research, Purdue University, West Lafayette, Indiana 47907, United States
- Iredia D. Iyamu Department of Medicinal Chemistry and Molecular Pharmacology, Institute for Drug Discovery, Center for Cancer Research, Purdue University, West Lafayette, Indiana 47907, United States
- John R. Horton Department of Epigenetics and Molecular Carcinogenesis, University of Texas MD Anderson Cancer Center, Houston, Texas 77030, United States
- **Dan Yu** Department of Epigenetics and Molecular Carcinogenesis, University of Texas MD Anderson Cancer Center, Houston, Texas 77030, United States
- **Taraneh Hajian** Drug Discovery Program, Ontario Institute for Cancer Research, Toronto, ON M5G 0A3, Canada
- Masoud Vedadi Department of Pharmacology and Toxicology, University of Toronto, Toronto, ON MSS 1A8, Canada; Drug Discovery Program, Ontario Institute for Cancer Research, Toronto, ON MSG 0A3, Canada; orcid.org/0000-0002-0574-0169
- Dante Rotili Department of Drug Chemistry and Technologies, Sapienza University of Rome, 00185 Rome, Italy; orcid.org/0000-0002-8428-8763
- Antonello Mai Department of Drug Chemistry and Technologies and Pasteur Institute, Cenci-Bolognetti Foundation, Sapienza University of Rome, 00185 Rome, Italy; orcid.org/0000-0001-9176-2382
- **Robert M. Blumenthal** Department of Medical Microbiology and Immunology and Program in Bioinformatics, The University of Toledo College of Medicine and Life Sciences, Toledo, Ohio 43614, United States
- Xing Zhang Department of Epigenetics and Molecular Carcinogenesis, University of Texas MD Anderson Cancer Center, Houston, Texas 77030, United States

Complete contact information is available at:

https://pubs.acs.org/10.1021/acschembio.3c00035

# Author Contributions

<sup>∨</sup>J.Z. and Y.D. contributed equally to this work. J.Z. performed CamA and PCIF1 purification, crystallization, and inhibition assays; Y.D. performed compound synthesis (7–10) and PRMT inhibition; I.D.I. provided compounds of 6 series; J.R.H. helped with X-ray data collection, provided Dam enzyme, and performed compound modeling; D.Y. purified MettL16 and MettL5; T.H. and M.V. purified MettL3–MettL14 complex; D.R. and A.M. provided compound 7 used in the selectivity assays; R.M.B. performed bioinformatics, assisted in preparing the manuscript, and participated in discussion; X.Z. provided supervision, conceptualization, and project administration; R.H. and X.C. designed and organized the study, performed writing, reviewing, and editing of the manuscript, and provided conceptualization and funding acquisition.

#### Notes

The authors declare no competing financial interest.

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# ABBREVIATIONS

CamA, Clostridioides difficile-specific DNA adenine methyltransferase; CDI, Clostridioides difficile infections;  $IC_{50}$ , half maximal inhibitory concentration;  $K_D$ , dissociation constant; MTase, methyltransferase; PRMT, protein arginine methyltransferase; SAH, S-adenosyl-L-homocysteine; SAM, S-adenosyl-L-methionine; TFA, trifluoroacetic acid

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