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Complete List of Authors:	Misson, Laetitia; University of Basel, Department for Chemistry Burn, Reto; University of Basel, Department for Chemistry Vit, Allegra; Helmholtz Zentrum für Infektionsforschung, Struktur und Funktion der Proteine Hildesheim, Juila; University of Basel, Department for Chemistry Beliaeva, Mariia; University of Basel, Department for Chemistry Blankenfeldt, Wulf; Helmholtz Zentrum für Infektionsforschung, Struktur und Funktion der Proteine Seebeck, Florian; University of Basel, Department for Chemistry

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Inhibition and regulation of the ergothioneine biosynthetic methyltransferase EgtD

Laetitia Misson,^{1,&} Reto Burn,^{1,&} Allegra Vit,^{2,&} Julia Hildesheim,¹ Mariia A. Beliaeva,¹ Wulf Blankenfeldt^{2,3} and Florian P. Seebeck^{1*}

¹Department for Chemistry, University of Basel, BPR 1096, Mattenstrasse 24a, Basel, Switzerland ²Structure and Function of Proteins, Helmholtz Centre for Infection Research, Inhoffenstr. 7, 38124, Braunschweig, Germany ³Institute for Biochemistry, Biotechnology and Bioinformatics, Technische Universität Braunschweig, 38106 Braunschweig, Germany

*To whom correspondence should be addressed: florian.seebeck@unibas.ch &These authors contributed equally to this work

Abstract. Ergothioneine is an emerging factor in cellular redox homeostasis in bacteria, fungi, plants and animals. Reports that ergothioneine biosynthesis may be important for the pathogenicity of bacteria and fungi raise the question as to how this pathway is regulated and whether the corresponding enzymes may be therapeutic targets. The first step in ergothioneine biosynthesis is catalyzed by the methyltransferase EgtD that converts histidine into N- α -trimethylhistidine. This report examines the kinetic, thermodynamic and structural basis for substrate, product and inhibitor binding by EgtD from *Mycobacterium smegmatis*. This study reveals an unprecedented substrate binding mechanism and a fine-tuned affinity landscape as determinants for product specificity and product inhibition. Both properties are evolved features that optimize the function of EgtD in the context of cellular ergothioneine production. Based on these findings we developed a series of simple histidine derivatives that inhibit methyltransferase activity at low micromolar concentrations. Crystal structures of inhibited complexes validate this structure- and mechanism-based design strategy.

Keywords. Methyltransferase, inhibitor design, mechanism, oxidative stress **Introduction**

Ergothioneine (**EGT**, Figure 1), the betaine of 2-mercaptohistidine, is a ubiquitous metabolite. Many bacteria¹⁻⁵ and most fungi biosynthesize EGT.^{1, 6} Plants and animals absorb EGT from their environment through a dedicated EGT transporter protein.^{7, 8} Active procurement of EGT by such a

diverse array of organisms indicates the EGT may play a fundamental role in cellular life. This hypothesis is more than half a century old but is now being tested and debated with increasing effort.⁹⁻¹⁴ Despite this recent attention, precise mechanisms by which EGT protects prokaryotic and eukaryotic cells are still elusive.^{11, 13, 14} The unusual redox activity and metal binding properties of the mercaptoimidazole side chain¹⁵⁻¹⁸ could enable EGT to participate in a broad range of processes¹⁹ including protection against reactive oxygen species,²⁰ reduction of oxidized heme-proteins,²¹ or passivating redox-active transition metals.^{22, 23}

Cellular dependence on EGT has been demonstrated for several microbial organisms.²⁴ Deletion of EGT biosynthetic genes in *Mycobacterium smegmatis*, *Mycobacterium tuberculosis*, *Streptomyces coelicolor*, *Neurospora crassa* and *Aspergillus fumigatus* produced strains with reduced resistance against oxidative stress.²⁵⁻³¹ In *M. tuberculosis* these deletions increased susceptibility to antimycobacterial drugs, and decreased viability in macrophages and in mice.³⁰ These recent findings raise the possibility that EGT biosynthesis – a process that does not occur in human cells – may be a target for novel antiinfective therapeutics. The genetic studies also agree that mutating the gene for the S-adenosylmethionine (SAM) dependent methyltransferase EgtD induces complete EGT deficiency in bacteria and fungi. This dependence validates EgtD as a potential target for EGT biosynthesis inhibitors.

EgtD initiates EGT biosynthesis by methylating histidine (HIS) to produce N- α -trimethylhistidine (TMH) via the intermediates N- α -monomethyl- (MMH) and N- α -dimethylhistidine (DMH).^{2, 32, 33} TMH is consumed by the oxygen- and iron-dependent sulfoxide synthase EgtB. This enzyme attaches the sulfur atom of γ -glutamylcysteine (γ GC) to carbon 2 on the imidazole ring of TMH.^{34, 35} Subsequent steps catalyzed by the amidohydrolase EgtC and the β -lyase EgtE result in EGT (Figure 1).^{36, 37} Fungal homologs of EgtB utilize cysteine instead of γ GC as sulfur donor, but the chemistry of this reaction is likely similar to that of mycobacterial enzymes.^{27, 38-40} Some cyanobacterial species recruited a homologous iron-dependent enzyme from a different pathway to act as an EgtB surrogate in EGT production.⁴¹ An even more surprising variation of this pathway occurs in anaerobic green sulfur bacteria. These organisms utilize a rhodanese-like enzyme (EanB) to attach sulfur to TMH in an oxygen-independent reaction.⁵ All these pathway variations include an EgtD-type methyltransferase, making this enzyme the sole indispensable component of EGT production (Figure 1).

Figure 1. Four biosynthetic pathways for ergothioneine (EGT) production in mycobacteria,² fungi, ^{27, 38-40} cyanobacteria,⁴¹ and anaerobic green sulfur bacteria.⁵

The growing recognition of EGT as a relevant factor in microbial metabolism and the key role of EgtD in EGT biosynthesis motivated us to examine the kinetic, thermodynamic and structural basis for

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ligand recognition by EgtD from *M. smegmatis*. This analysis revealed i) that EgtD binds its substrate by a mechanism which is unprecedented among SAM-dependent methyltransferases, ii) that EgtD activity is subject to stringent feedback regulation and iii) and that the EgtD active site can adapt to methylate a primary, a secondary and a tertiary amine with increasing efficiency. These findings were used to develop and validate the first designs of specific EgtD inhibitors.

Results and Discussion

Substrate-binding order. EgtD consumes three SAM equivalents to methylate HIS to TMH in three consecutive two-substrate – two-product reactions with MMH and DMH as reaction intermediates.⁴² In principle it is possible that trimethylation is processive, meaning that HIS only leaves the enzyme after all three methyl groups are installed. This model is unlikely. Assuming diffusion-limited substrate binding, and a dissociation constant of 4 uM for the EgtD:DMH complex, we find that unproductive dissociation of this complex is at least 10³-fold faster than turnover to TMH.³³ Hence, EgtD-catalyzed trimethylation is a distributive process. The efficiency of EgtD-catalyzed consumption of SAM is two-fold and three-fold less efficient when the methyl acceptor is HIS instead of DMH or MMH,³³ showing that the first methyl transfer is the rate limiting step of TMH production. Therefore, we concluded that the steady-state behavior of EgtD is dominated by the first methylation step.

The order of substrate binding was elucidated by measuring the apparent Michaelis-Menten parameters k_{cat} and K_{M} for HIS and SAM as a function of both substrate concentrations (Figure S1).³³ EgtD-catalyzed consumption of SAM was monitored by an enzyme-coupled UV assay.⁴³ The recorded data revealed that the apparent $K_{M,SAM}$ depends on [HIS], but that the apparent $K_{M,HIS}$ is largely independent of [SAM] (Figure 2). This behavior is diagnostic for an ordered sequential substrate binding mechanism, with HIS as the leading substrate.⁴²

Figure 2. Lineweaver–Burk plots of kinetic data used to examine the substrate binding mechanism of EgtD. **Top:** Primary and secondary plots with SAM as the variable substrate in presence of different concentrations of HIS. **Bottom:** Primary and secondary plots with HIS as the variable substrate in presence of different concentrations of SAM.

An obligatory binding order is consistent with the structure of EgtD (Figure 3). The active site of this enzyme is located in a cleft between the SAM-binding Rossmann-fold domain and the HIS-binding domain. The first domain is conserved among class I methyltransferases,^{44, 45} but the second domain is exclusive to *Methyltransf_33* enzyme family members, such as the Trp-, Tyr- and dimethylallyl-tryptophan methyltransferases.^{33, 46}

The crystal structure of EgtD in complex with DMH and SAH shows that the enzyme completely sequesters the methyl acceptor from bulk solvent (Figure 3). The only direct non-protein contact to DMH is provided by the sulfur atom of SAH. The only path for HIS in and out of this pocket leads through the SAM/SAH-binding site. Unless substrate binding is accompanied by large scale unfolding and refolding of the HIS-binding domain, the methyl-acceptor can reach its binding pocket only in the absence of SAM/SAH. Therefore substrate binding and product release must follow an ordered sequence.

This methyl-acceptor first binding order distinguishes EgtD from all characterized natural product methyltransferases.⁴⁷ Methyltransferases which methylate small substrates⁴⁷ usually follow a SAM-first or a random binding mechanism. Some DNA-, RNA- or protein-methyltransferases may follow an apparent substrate-first binding mechanism.⁴⁸ However, these enzymes often bind their macromolecular substrates through interactions outside the active side, which makes the comparison to enzymes with small substrates difficult.

Figure 3: Structure of EgtD in complex with S-adenosyl homocysteine (SAH, green) and N- α -dimethylhistidine (DMH, orange)(PDB: 4PIO).⁵ The substrate-binding domain (blue) is formed by residues 1 – 60 and 196 – 286. The SAM-binding domain is conserved in most SAM-dependent methyltransferases.

Product inhibition by TMH. EgtD is characterized by significant HIS-competitive inhibition by the product TMH. This behavior is also unusual for a SAM-dependent methyltransferase. To examine this trait of EgtD we recorded methyl-transfer activities in the presence of several TMH concentrations with either HIS or SAM as the substrate with variable concentration, while keeping the second substrate concentration constant. Plotting this data in form of Lineweaver–Burk plots showed that TMH behaves as a competitive inhibitor with respect to both substrates (Figure 4). The K_i for HIS-competitive inhibition of EgtD by TMH was determined by measuring the apparent $K_{M,HIS}$ in the presence of 500 µM SAM and three different concentrations of TMH. From this data K_i was calculated using the equation $K_i = K_M [TMH]/(K_{M,app} - K_M)$ (Table 1, Figure S2).

Table 1. Inhibition constants (K_i) for EgtD inhibitors^[a]

EgtD ligands	<i>K</i> i (μM) L-derivative	<i>K</i> i (μM) racemic
TMH	39 ± 6	-
1	-	8.5 ± 2.1
2	21 ± 3	41 ± 6
3	-	93 ± 11
4	-	49 ± 14

-	5.4 ± 1.6
-	72 ± 17
-	25 ± 1
2.6 ± 0.4	6.2 ± 1.5
-	8.2 ± 2.4
800 ± 200	-
100 ± 40	-
800 ± 200	-
200 ± 80	-

^[a] HIS-competitive inhibition of EgtD was quantified by measuring the apparent $K_{M,His}$ at three different inhibitor concentrations in the presence of 500 μ M SAM. Inhibition constants were determined using the equation $K_i = K_M[TMH]/(K_{M,app} - K_M)$

Table 2. Binding constants (K_D) binary and ternary EgtD complexes ^[a]

EgtD ligands	KD	ΔG	ΔH	$T\Delta S$
	(µM)	(kcal/mol)	(kcal/mol)	(kcal/mol)
HIS	290 ± 14	-4.8	- 8.0	- 3.3
HIS:SAH ^[b]	37 ± 1	-6.1	-10	-3.9
MMH ^[b]	70 ± 30	-5.7	- 13	-7.5
MMH:SAH ^[b]	14 ± 7	-6.6	-11	-4.5
DMH ^[b]	4 ± 2	-7.4	-5.0	2.4
DMH:SAH ^[b]	2 ± 1	-7.8	-27	-19
TMH	26 ± 4	-6.3	-8.0	-1.6
TMH:SAH	0.11 ± 0.01	-9.7	-9.2	0.2

^[a] Dissociation constants [*K*_D] were determined by isothermal calorimetry titration at 25 °C. ^[b] Data from Ref.³³

The value of K_i for TMH corresponds well with the dissociation constant (K_d) of the EgtD:TMH complex as determined by isothermal titration calorimetry (Table 2, Figure S3). The affinity of EgtD for TMH increased by 240-fold in the presence of 7 mM SAH (Figure S3). However, because the reaction mixtures used for the kinetic measurements contained SAH nucleosidase and adenine deaminase, SAH cannot accumulate,⁴³ and does not contribute to inhibition. Similarly, the SAH concentrations in living cells is also kept in the in the low micromolar range, suggesting that EgtD inhibition by SAH may not be significant in vivo.⁴⁹ On the other hand, stress factors that lead to accumulation of SAH might indeed interfere with EGT production.

Figure 4. Lineweaver–Burk plots of the data used to examine EgtD inhibition by TMH and **8. Top:** Primary plots with HIS or SAM as the variable substrate in presence of different concentrations of TMH **Bottom:** Primary plots with HIS or SAM as the variable substrate in presence of different concentrations of **8**.

Comparison to related methyltransferases. Methyltransferases are commonly inhibited by the consumed methyl donor, S-adenosyl homocysteine (SAH), which acts as a SAM-competitive inhibitor.⁴⁹ Inhibition by the methylated product is far less common among methyltransferases.

Therefore we wondered whether product inhibition by TMH is a specifically evolved feature of EgtD or whether this behavior is a mere consequence of the unusual substrate binding order or of the permethylation reaction. To address this question, two close EgtD homologs were examined. The first enzyme is the tyrosine betaine synthase (Ybs) from *Aspergillus nidulans*. Although Ybs shares only 28 % sequence identity with EgtD,³³ this fungal enzyme contains an almost identical set of active site residues. The only apparent differences between EgtD and Ybs map to the side chain binding pocket for the substrate. Hence, Ybs and EgtD should share all catalytic properties that are inescapable consequences of the protein architecture or the catalyzed reaction.

Using the same kinetic assay as described above we determined that Ybs catalyzes tyrosine methylation with similar efficiency as EgtD catalyzed methylation of HIS (Table 3, Figure S4). In contrast to EgtD, Ybs is not inhibited by its final product N- α -trimethyltyrosine (TMY, $K_i > 1$ mM, Figure S5). The efficiencies at which the two enzymes catalyze the conversion of DMH or DMY to TMH or TMY, respectively, were also determined. EgtD-catalyzed methyl transfer is three-fold more efficient when the methyl acceptor is DMH instead of HIS.³³ In contrast, Ybs-catalyzed methyl transfer is four-fold less efficient when the methyl acceptor is DMY instead of Tyr. As a consequence, EgtD and Ybs give rise to different product distributions when SAM is the limiting substrate. EgtD produces predominantly TMH, while Ybs produces predominantly DMY (Table 3).

Similar observations were made with an engineered EgtD variant (EgtD_{E282A,M252V}) that methylates tryptophan instead of HIS.³³ The variant contains two mutations in the substrate-binding domain that accommodate an indole instead of an imidazole ring. The crystal structure of this enzyme in complex with tryptophan (Trp) revealed an otherwise unchanged active site geometry.³³ EgtD_{E282A,M252V} catalyzed methylation of Trp with an efficiency only six-fold lower than that of the wild type enzyme with HIS as substrate (Table 3, Figure S6). However, methylation of N- α -dimethyltryptophan (DMW) to N- α -trimethyltryptophan (TMW) is 20-fold less efficient than the corresponding transformation of DMH by EgtD. The reduced efficiency is due to a reduced k_{cat} , suggesting that suboptimal positioning of the non-native substrate in the mutated active site specifically affects methyl transfer to DMW.

Based on the comparison of these three methyltransferases we conclude that efficient trimethylation and product inhibition as observed by EgtD are not inescapable consequences of the active site architecture, the catalyzed reaction type, or the substrate binding mechanism. More likely, the two behaviors rely on structural optimization of the EgtD active site and must have emerged by positive selection to serve a function. As will be discussed below, cooperative trimethylation and product inhibition may play important roles in quality control and regulation of EGT biosynthesis in Mycobacteria.

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Table 3. Kinetic parameters of the aromatic amino acid betaine sy	nthases ^a
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substrates	enzyme	k _{cat} (s ⁻¹)	<i>К</i> м (μМ)	k _{cat} / К _М (M ⁻¹ s ⁻¹)	MMH/DMH/TMH ratio (%)
HIS ^[b]	EgtD	5.8 x 10 ⁻¹	110	5.3 x 10 ³	
DMH ^[b]	EgtD	4.3 x 10 ⁻¹	32	1.7 x 104	<1/17/83
Tyr ^[b]	Ybs	1.1 x 10 ⁻¹	21	5.2 x 10 ³	
DMY	Ybs	5.0 x 10 ⁻²	43	1.2 x 10 ³	<1/60/40
Trp ^[b]	EgtD _{E282A,M252V}	1.1 x 10 ⁻¹	20	5.5 x 10 ³	
DMW	EgtD _{E282A,M252V}	0.9 x 10 ⁻²	11	8.2 x 10 ²	<1/95/5

^[a] Reaction conditions: 25 °C, Tris HCl 50 mM, pH 8, NaCl 50 mM, MnCl 200 μM, 500 μM SAM, SAH nucleosidase 5 μM, adenine deaminase 10 μM

^[b] Data from Ref.³³

Structure of EgtD in complex with TMH. In order to mediate efficient trimethylation and to allow for product inhibition, EgtD must be able to bind to a primary, a secondary, a tertiary and a quaternary amine. How can this enzyme accommodate the changing hydrogen-bonding requirements of its ligand? To examine this question, we solved the crystal structure of EgtD in complex with TMH (Table S1 & S2). This structure superimposes with the EgtD:HIS ⁵⁰ and the EgtD:DMH ³³ complexes a mutual r.m.s.d. of 0.4 Å (entire chain). The three ligands HIS, DMH and TMH occupy essentially the same position and almost all atoms of the active site residues superimpose. The important exceptions are residues in direct contact with the α -amino moiety. These residues are Asn166 and Gly161.

In the TMH complex two N- α -methyl groups make close contact with the backbone carbonyl of Gly161 (3.1 Å) and with the side chain carbonyl of Asn166 (3.2 Å). The carbonyls approach the N- α -methyl groups in a (O-C-N α) angle of 167° (Gly161) and 177° (Asn166) This geometry is suggestive of attractive interaction between the positively polarized methyl groups and the carbonyl functions. The third N- α -methyl group points toward the SAM-binding site.

In the DMH complex, one N- α -methyl group makes the same interaction with the Gly161 carbonyl function (d = 3.0 Å) and the other N- α -methyl group points toward the SAM-binding site. Asn166 moved 1.3 Å closer to the substrate to form a 2.7 Å hydrogen bond to the α -amino group of the ligand. In the HIS complex the ligand forms two rather loose hydrogen bonds to Asn166 (3.1 Å) and Gly161 (3.4 Å). In order to establish a hydrogen bond, the Gly161 carbonyl oxygen moved by 1.8 Å towards the α -amino group of HIS. This rearrangement is made possible by conformational changes of the backbone including residues 159 - 162.

The three structures show that EgtD solvates the N- α -amino moiety of its substrates and product by a highly polar pocket with adaptable size. Remarkably, HIS is by far the weakest binder, despite forming two classical hydrogen bonds (Table 2).³³ In the DMH complex the hydrogen bond to Gly161 is lost, but the hydrogen bond to Asn166 becomes shorter and hence, stronger. As a consequence, DMH is a 100-fold stronger EgtD ligand ($K_D = 4 \mu M$) than HIS. Surprisingly, TMH is still a comparatively strong ligand ($K_i = 40 \mu M$) even though both hydrogen bonds are lost. It is possible that the close interaction between two N- α -methyl groups and the carbonyls from Gly161 and Asn166 are attractive and partially compensate for the lack of hydrogen bonding.

Figure 5. Crystal structures of EgtD in complex A) HIS (PDB entry 4UY7, ⁵⁰); B) DMH (PDB entry 4PIN,³³); C) TMH; D) **8**; E) **2**; F) **3**. Unbiased m|F_{obs}|-D|F_{calc}| electron density (σ-level = 2) of the compounds is shown in green.

Catalytic cycle. The structure of the EgtD:TMH complex also illustrates why TMH is necessarily a competitive inhibitor with respect to both HIS and SAM. A model of EgtD in complex with SAM and TMH indicates that the third N- α -methyl groups of TMH and the sulfonium methyl group of SAM would clash (Figure S7). Hence, binding to the two ligands is mutually exclusive. In the conformation of DMH observed in the EgtD:DMH complex the same steric clash would prevent binding of SAM. In order to form the ternary complex DMH (**a**, Figure 6) must first adopt an alternative conformation in which the two N- α -methyl groups point towards Asn166 and Gly161 (**b**). This conformer can accept SAM (**c**) to form the reactive complex that decays via S- to N-methyl transfer to form the product complex (**d**). Based on this mechanistic model we hypothesized that DMH analogs that make the same interactions in the active site but cannot undergo the same conformational change could be potent EgtD inhibitors.

Figure 6. Mechanism of EgtD-catalyzed methylation of DMH. Residues Asn166 and Gly161 are shown in gray.

Strategies for inhibitor design. To test this idea we synthesized histidine derivatives with cyclic tertiary amines in place of the N- α -dimethyl amino moiety of DMH (Figure 5, see supporting information). The syntheses and characterization of compounds depicted in Figure 7 are described in the supporting information. The inhibitory activities were measured using the same assay as described above. Consistent with the design strategy compounds **1**, **2**, and **3** were not methylated by EgtD, but instead inhibit EgtD-catalyzed methylation of HIS (Figure S8 & S9, Table 1). In the presence of 0.5 mM SAM, inhibition by **1**, **2**, and **3** is characterized by inhibition constants (K_i) of 9, 40 and 90 μ M respectively (Table 1). Compounds **1**, **2** and **3** were synthesized in racemic form. Given that EgtD only interacts with L-amino acids, it is fair to assume that only the L-isomers of the inhibitors would bind (Figure S10). This assumption is corroborated by the finding that the pure L-form of **2** inhibits EgtD with a two-fold lower K_i than measured for the racemic mixture (Table 1).

To examine the binding mode of these inhibitors, we determined the crystal structures of EgtD in complex with **2** and with **3** (Figure 5). Both structures show that the tertiary amines of **2** and **3** form the same hydrogen bond to Asn166 (2.9 and 3.0 Å) as seen in the EgtD:DMH complex (Figure 5). The electron density around ligands clearly shows that EgtD binds only the L-isomer of **2** and **3**. One of the N- α -methylene carbons of **2** and **3** stacks against the carbonyl group of Gly161 (3.2 and 3.1 Å) and the rest of the pyrrolidino- and morpholino-rings block the space where the methyl group of SAM would approach the methyl acceptor. Both rings push the side chain of Thr163 which is pushed by 0.5 Å away from its position in the EgtD:TMH and EgtD:DMH structures. This steric stress also provides an explanation for why the size of the cyclic substituents in **1**, **2** and **3** correlates inversely with their inhibitory activity (Table 1).

Figure 7. Structure of tested EgtD inhibitors.

Avoiding competition with SAM. As an alternative design strategy we examined inhibitors that only compete with HIS, but do not compete with SAM. Such compounds could form inhibited complexes with EgtD that are not destabilized by the generally high cellular concentration of SAM. To test this idea we synthesized racemic histidine derivatives with a proton (4), a methyl group (5), a hydroxyl group (6), fluoride (7), chloride (8) or bromide (9) in place of the α -amine group. In kinetic assays these compounds, except for 5, proved stronger HIS-competitive inhibitors than TMH (Table 1). The inhibition constant of the L-form of 8 was again two-fold lower than that of the racemic form, indicating that the active site selectively binds one isomer. Determination of the inhibition mechanism of chlorohistidine (8, Figure 4) revealed HIS-competitive and SAM-uncompetitive inhibition, suggesting that the EgtD:8 complex can still bind SAM.

The crystal structure of EgtD in complex with **8** confirms that this ligand occupies the same active site position as all other co-crystallized histidine derivatives (Figure 5). Unlike the α -amine substituents in HIS, DMH, TMH, **2** and **3** the chloride substituent makes no direct contact with any protein residue. Solvation of the carboxylate and the imidazole ring by the active site apparently provides enough attractive interaction to induce strong inhibitory activity of the methyl- (**5**), chloro-(**8**) and bromo-substituted derivatives (**9**). The lower affinities of compounds **4**, **6** and **7** are most likely due to stronger solvation of the free ligand by water. The relatively poor affinity of EgtD for **6** mirrors the low affinity for HIS and corroborates the notion that the residues Gly161 and Asn166 are not particularly well positioned to engage in hydrogen bonding.

Bisubstrate inhibitors. A more common strategy to design inhibitors for methyltransferases targets the SAM-binding pocket. One successful way to increase the specificity towards one particular type of methyltransferase is to integrate structural motifs from SAM with those of the specific methyl acceptor into a single bisubstrate inhibitor.⁵¹⁻⁵⁶ To explore this approach for the design of EgtD inhibitors we synthesized four histidine derivatives (**10 – 13**) that are N-substituted to mimic the methionyl moiety of SAM. All four compounds displayed weak inhibitory activity (Table 1, Figure S11). One explanation could be that that the amino acid substituents are not recognized by the methionyl-binding site in EgtD. Indeed, reinspection of EgtD in complex with DMH and SAH showed the possibility that the chosen methionyl mimics may be too short to bridge the histidine binding site and the methionyl-binding site. It is also possible that the compounds do bind as intended, but that recognition of the methionyl-moiety does not produce enough attractive interaction to outweigh competition with SAM and HIS. The observation that the additional N- α -methyl group on compounds **11** and **13** increase the affinity by four to eight-fold provides further evidence that tertiary amines bind more strongly to EgtD than secondary amines.

Feedback inhibition. EgtD catalyzes the first step in EGT biosynthesis (Figure 1). This reaction converts the primary metabolites HIS and SAM to TMH as a substrate for the subsequent enzyme EgtB. Methyltransferases are very common contributors to biosynthetic pathways in natural product biosynthesis.⁵⁷ However, a cursory inspection of all methyltransferase entries in the Braunschweig Enzyme Database (BRENDA) shows that SAM-dependent methyl transfers rarely occur as first biosynthetic steps.⁵⁸ For example, alkaloids or phenylpropanoids usually receive methyl groups at later biosynthetic stages.⁵⁹⁻⁶²

The unique role of EgtD as the gateway to EGT production raises the specific problem of regulation. Two studies on a gliotoxin deficient strain of *A. fumigatus*, and a mycothiol-deficient strain of *M. smegmatis* revealed that these deficiencies are compensated by EGT overproduction. ^{63, 64} The mechanism by which ETG productivity is coupled to seemingly unrelated biosynthetic activity is not known. However, the two studies provide first indications that EGT production may be regulated.

Because methyl transfer from SAM to His is essentially irreversible, and because both substrates are abundant metabolites, regulation of EgtD activity is essential, either by transcriptional control, by reversible inhibition or by destruction of the enzyme. One regulatory mechanism has been proposed based on the finding that EgtD from *Mycobacterium tuberculosis* may be a substrate of the protein kinase PknD.³¹ According to this model the kinase phosphorylates a key active site residue of EgtD (Thr213) to block activity.

The observation that EgtD is subject to significant product inhibition highlights an alternative mode of regulation. Most SAM-dependent methyltransferases are inhibited by the side product S-adenosyl homocysteine (SAH). Therefore, methyltransferase activities are often modulated by the cellular concentration of SAH or the ratio of SAH/SAM.⁴⁹ Inhibition by the methylated product is far less common.⁶⁵⁻⁷⁰ The caffeoyl-coenzyme A 3-O-methyltransferase from *Petroselinum crispum* (Parsley) provides a rare exception. This enzyme is inhibited by its product feruloyl-CoA ($K_{i, \text{ feruloyl-CoA}} = 11 \mu M$) which allows strict regulation of the steady-state product concentration.⁷⁰

By analogy, we hypothesize that product inhibition of EgtD may also have physiological relevance. Incidentally, the value of the associated inhibition constant ($K_{i,TMH} = 40 \mu M$, Table 1) lies in the same range as the K_M for TMH ($K_{M,TMH} = 43 \mu M$) of the next enzyme in the pathway, EgtB (Figure 1).³⁴ These parameters ensure that TMH cannot accumulate to high concentrations even if EgtB activity decreases, for example due to limiting supply of the co-substrates γ GC and O₂. Consequently, the cellular supply of TMH is adjusted to the rate of EGT production. In addition, the stabilizing effect of SAH on the EgtD:TMH complex raises the possibility that EGT biosynthesis is also regulated by the cellular concentration of SAH. The underlying prediction that product inhibition of EgtD is a specifically evolved trait is corroborated by the finding that the homologous tyrosine betaine synthase Ybs is not inhibited by product despite significant active site similarity to EgtD.

Proofreading. In addition to making the first intermediate in EGT biosynthesis, EgtD also serves as a quality control element of this pathway. Although EGT has been identified from a large range of sources,^{1, 4, 5, 23} there are no isolation reports of EGT derivatives that lack one, two or all N- α -methyl groups. For reasons that are not exactly clear the betaine moiety of EGT is important for physiological function. Subsequent enzymes in the EGT pathway are unable to proofread the methylation state of their substrates.^{2, 36, 37} For example, EgtB from *M. smeqmatis* turns over DMH and TMH with almost the same efficiency.² EgtC and EgtE are unlikely to prevent alternative products because the reaction catalyzed by EgtB is irreversible. Hence, the only mechanism to prevent the formation of unwanted EGT derivatives is to limit the cellular concentration of DMH. Limiting this concentration is an important role of EgtD. As shown in Table 2 EgtD binds HIS, MMH and DMH with increasing affinity. As a result, each added methyl group on the methyl acceptor increases the probability of further methylation. The three methyl groups are transferred in a cooperative process that avoids the accumulation of MMH or DMH.³³ By contrast, the catalytic efficiency of the homologous enzymes Ybs and EgtD_{E282A,M252V} drops significantly after the first two methyl transfers to Tyr ad Trp, respectively (Table 3). Comparisons of the EgtD structure with the homology model of Ybs, and the crystal structure of EgtD_{E282A,M252V} do not reveal clear structural explanations for these different activities. One possibility is that transfer of the last methyl group is particularly sensitive to precise positioning of the N- α -dimethylated amino acid in the active site. Therefore, it is possible to

conclude that the ability of EgtD to catalyze cooperative trimethylation is also an essential and specifically evolved feature.

Mechanistic implications. Finally, we would like to summarize what can be learned about the catalytic mechanism and about inhibitor design from the ligand binding preferences of EgtD. EgtD can form up to thirteen binary and ternary complexes with its six native ligands HIS, MMH, DMH, TMH, SAM and SAH (Figure 8). Among these, EgtD:DMH and EgtD:TMH:SAH are the strongest binary and ternary complexes (Table 2). The interaction between the amide side chain of Asn166 and the α amino function of DMH shows that the α -amino function of the methyl acceptor is protonated in the binary complex (Figure 5). Apparently, the active site stabilizes a cationic moiety in this position. Unreactive histidine derivatives with neutral substituents (5, 8 and 9) form strong complexes with EgtD and SAM, suggesting that in the ternary complex the sulfonium moiety of SAM fully satisfies the requirement for a cationic charge in the active site. Consequently, the methyl acceptors HIS, MMH and DMH must lose a proton before or concomitant to SAM binding. Also, in order to make room for the second substrate, deprotonation must be accompanied by inversion of the α -amine. For example, DMH must turn the two N- α -methyl groups towards Gly161 and Asn166 in order to juxtapose the nucleophilic lone pair with the sulfonium methyl group of SAM (Figure 6). Each methyl transfer from HIS to TMH makes the ligand larger. The structures of EgtD in complex with HIS, DMH and TMH show how the active site undergoes stepwise expansion by repositioning of Gly161 and Asn166 to accommodate the growing size of the ligand.

To support efficient trimethylation the energy landscape of this expansion must be adjusted to increase the affinity for the methyl acceptor with each additional N- α -methyl group. As the thermodynamic binding data shows, EgtD follows exactly this expected behavior. A plot of the complex stabilities ($\Delta G_{binding}$) of EgtD with SAH and HIS, DMH or TMH shows that each additional methyl group on the methyl acceptor increases the complex stability by 1.2 kcal/mol (Figure 8, Table 2). A similar trend is apparent in the absence of SAH. Notably, the EgtD:TMH complex deviates from this trend. It seems possible that the stability of the EgtD:TMH complex is purposefully decreased to avoid inhibition by sub-micromolar concentrations of TMH.

A glance at the enthalpic and entropic contributions to the stability of the six complexes cautions that a purely structural interpretation of the binding data may be misleading (Table 2). For example, formation of the EgtD:DMH:SAH complex liberates more heat than formation of the EgtD:TMH:SAH complex. However, because the latter suffers almost no entropic penalty, the TMH complex is 20-fold more stable. The enthalpic term indicates that formation of a hydrogen bond to Asn166 and stacking one N- α -methyl group towards Gly161 in the EgtD:DMH:SAH complex amounts to more attraction

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than the two N- α -methyl interactions with Gly161 and Asn166 in the EgtD:TMH:SAH complex. The basis for the large entropic difference is more difficult to localize. It is also interesting to note that conversion of the EgtD:HIS:SAH complex to the EgtD:TMH:SAH complex is accompanied by 300-fold stabilization, which is entirely due to entropic contributions. The same trend applies to the binary complexes in the absence of SAH. This result indicates that the two N- α -methyl interactions with Gly161 and Asn166 can at least partially compensate for the loss of the two hydrogen bonds in the EgtD:HIS:SAH complex. One interpretation of this result is that the two N- α -methyl interactions with protein carbonyl groups are at least weekly attractive.

Figure 8. Top: Complete reaction scheme of EgtD catalyzed trimethylation of HIS. EgtD can combine with its six native ligands to 13 binary and ternary complexes. **Bottom:** the stability of EgtD complexes as determined by ITC (Table 2).

Conclusion. This report describes the unusual substrate binding mechanism of the SAM-dependent methyltransferase EgtD. Unlike most methyltransferases, this enzyme follows an obligatory sequential binding mechanism with the methyl acceptor as the leading substrate. Secondly, this enzyme can regulate EGT production by way of product inhibition. Third, the enzyme ensures efficient permethylation of its substrate and suppresses the accumulation of mono- and dimethylated intermediates. Product inhibition and efficient permethylation are the result of specific evolutionary optimization. These findings were exploited to design three types of substrate competitive EgtD inhibitors. The most efficient inhibitors (**5** & **8**) are very simple histidine derivatives that provide promising leads for further development EGT biosynthesis inhibitors.

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Supporting Information Available: Detailed descriptions of all experiments, supporting figures (S1 – S11), schemes (S1 - S8) and tables (S1 – S2) are shown in the supporting information. This material is available free of charge via the internet at http://pubs.acs.org.

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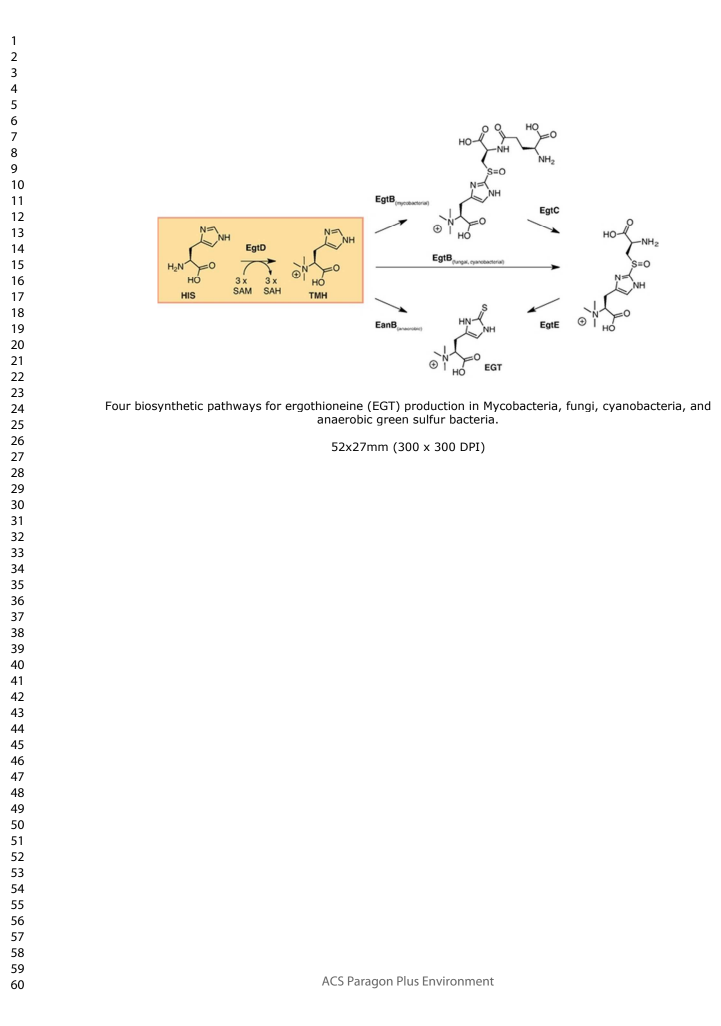
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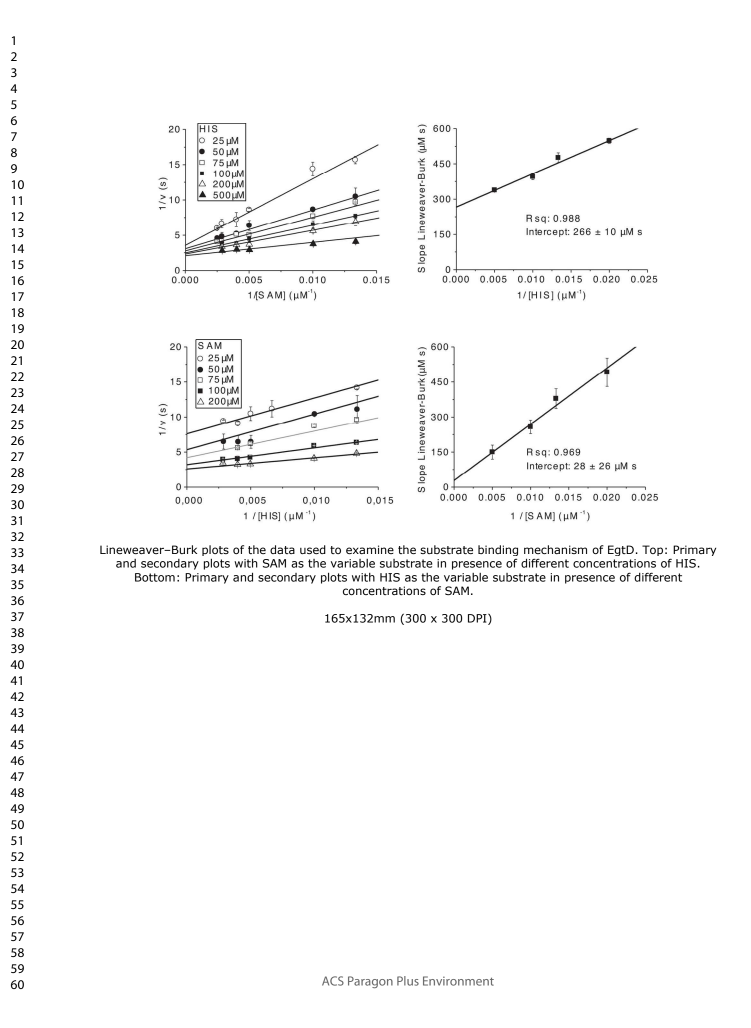
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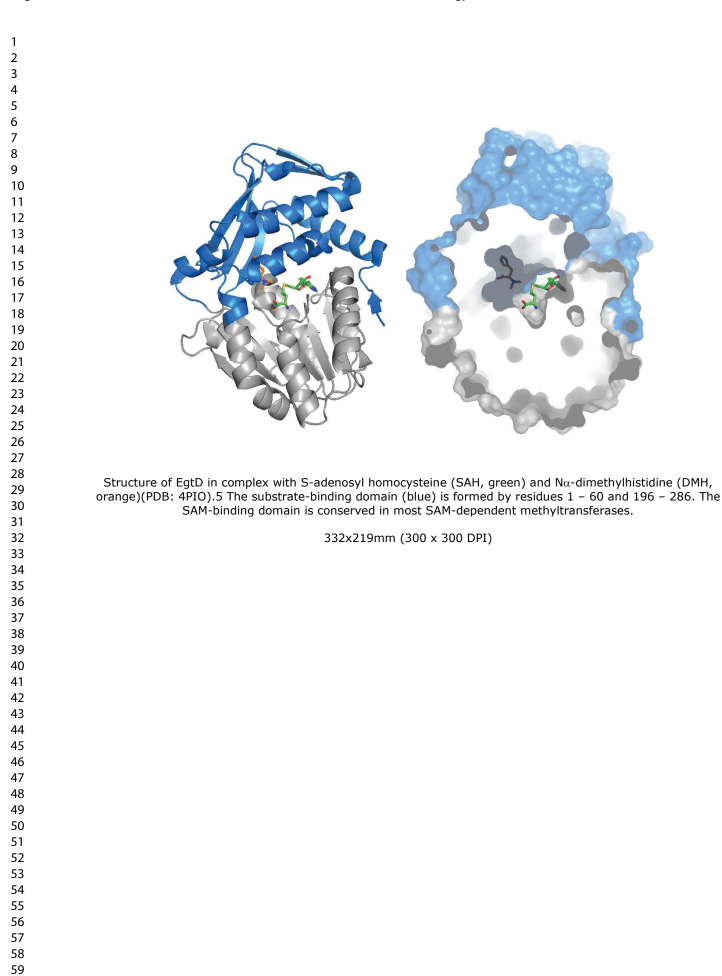
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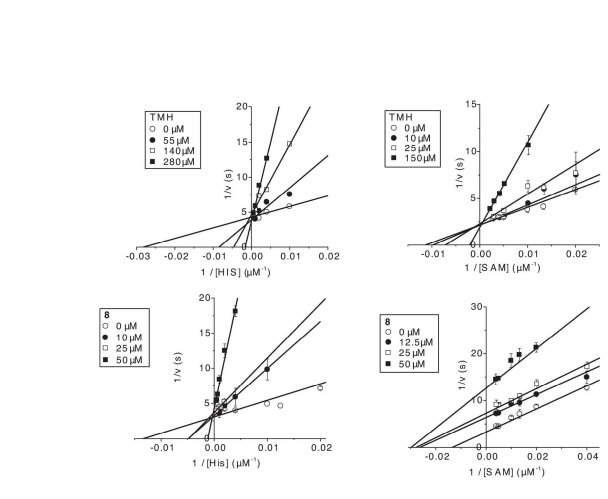
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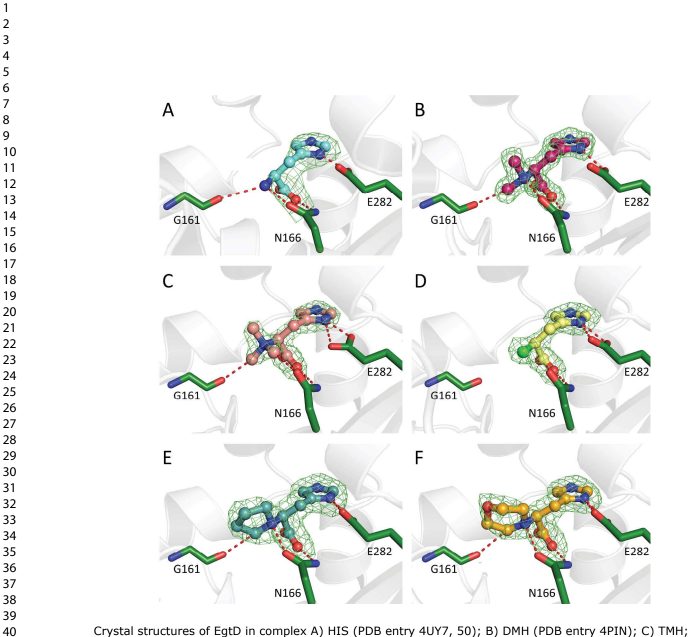






Lineweaver–Burk plots of the data used to examine EgtD inhibition by TMH and 8. Top: Primary plots with HIS or SAM as the variable substrate in presence of different concentrations of TMH Bottom: Primary plots with HIS or SAM as the variable substrate in presence of different concentrations of 8.

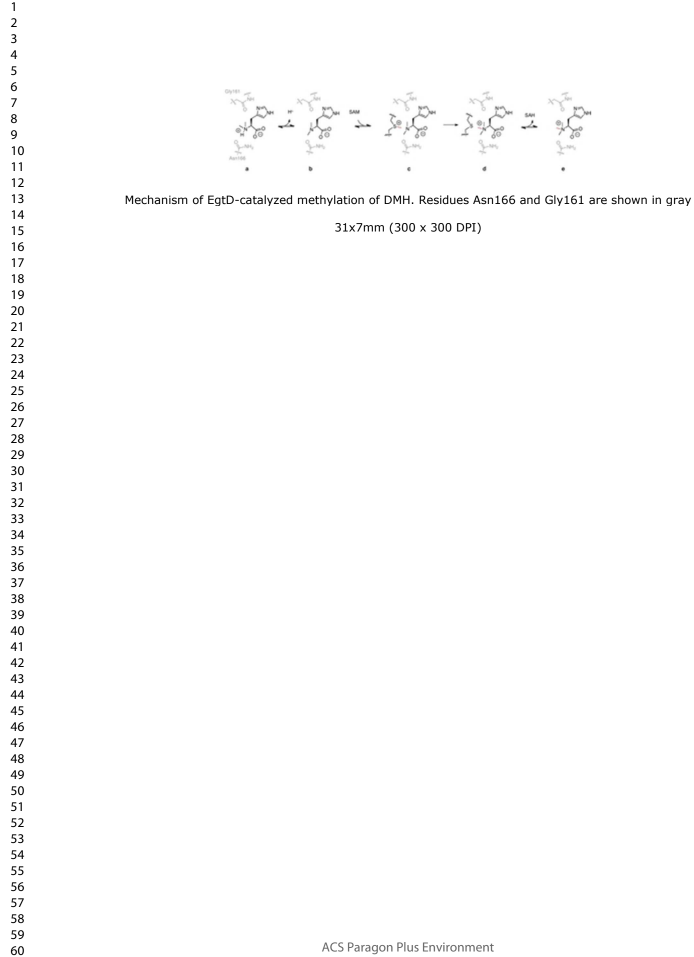
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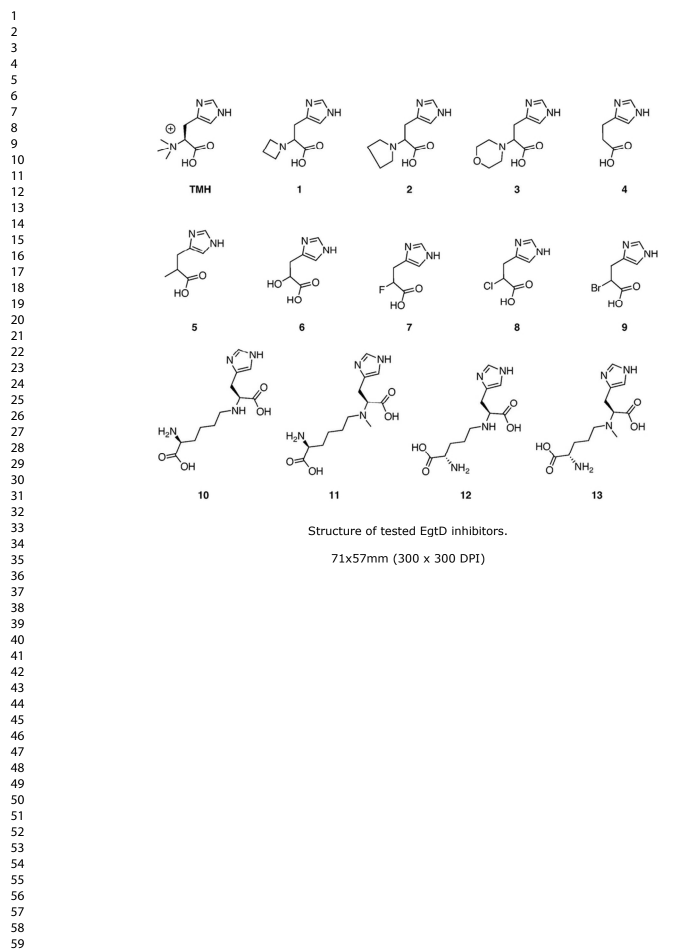


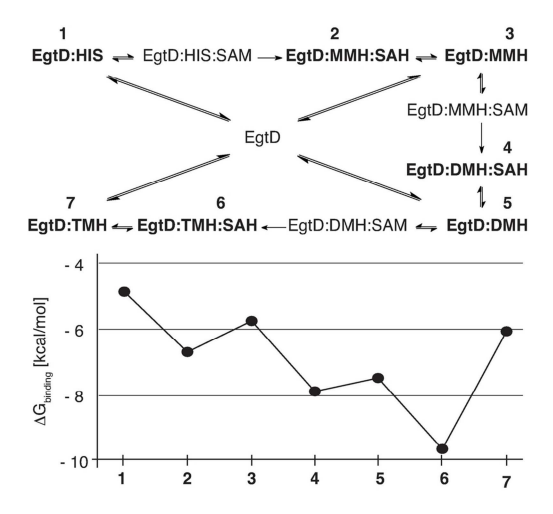
Crystal structures of EgtD in complex A) HIS (PDB entry 4UY7, 50); B) DMH (PDB entry 4PIN); C) TMH; D) 8; E) 2; F) 3 . Unbiased m|Fobs|-D|Fcalc| electron density (σ -level = 2) of the compounds is shown in green.

514x527mm (600 x 600 DPI)

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Top: Complete reaction scheme of EgtD catalyzed trimethylation of HIS. EgtD can combine with its six native ligands to 13 binary and ternary complexes. Bottom: the stability of EgtD complexes as determined by ITC (Table 2).

80x75mm (300 x 300 DPI)

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