Complementary Steric Engineering at the Protein−Ligand Interface for Analogue-Sensitive TET Oxygenases

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Supporting Information

ABSTRACT: Ten-eleven translocation (TET) enzymes employ O₂, earth-abundant iron, and 2-ketoglutarate (2KG) to perform iterative C−H oxidation of 5-methylcytosine in DNA to control expression of the mammalian genome. Given that more than 60 such C−H oxygenases are present in humans, determining context-dependent functions of each of these enzymes is a pivotal challenge. In an effort to tackle the problem, we developed analogue-sensitive TET enzymes to perturb the activity of a specific member. We rationally engineered the TET2−2KG interface to develop TET2 variants with an expanded active site that can be specifically inhibited by the N-oxalylglycine (NOG) derivatives carrying a complementary steric “bump”. Herein, we describe the identification and engineering of a bulky gatekeeper residue for TET proteins, characterize the orthogonal mutant−inhibitor pairs, and show generality of the approach. Employing cell-permeable NOG analogues, we show that the TET2 mutant can be specifically inhibited to conditionally modulate cytosine methylation in chromosomal DNA in intact human cells. Finally, we demonstrate application of the orthogonal mutant−inhibitor pair to probe transcriptional activity of a specific TET member in cells. Our work provides a general platform for developing analogue-sensitive 2KG-dependent oxygenases to unravel their functions in diverse signaling processes.

INTRODUCTION

Organisational multicellularity has been mechanistically linked to a steady increase in oxygen pressure in the atmosphere. This is likely due to the emergence of enzymes catalyzing oxygen-dependent biochemical reactions essential to metabolic and transcriptional regulations. An early class of such enzymes employ simple co-substrates such as 2-ketoglutaric acid (2KG) and iron for oxidative processing of proteins and nucleic acids. In humans, more than 60 such oxygenases catalyze C−H hydroxylation in diverse nuclear and cytoplasmic substrates and contribute significantly to cellular processes ranging from DNA repair to hypoxic signaling. Within this group, ten-eleven translocation (TET) enzymes perform C−H oxidation of 5-methylcytosine (5mC) to 5-hydroxymethylcytosine (5hmC), 5-formylcytosine (5fC), and finally 5-carboxylcytosine (5caC) provides a biochemical basis for active DNA demethylation. 5caC could be recognized and excised by thymine DNA glycosylase (TDG) to effect base excision repair (BER)-mediated replacement of 5caC with deoxycytidine (dC). Direct decarboxylation of 5caC could be another potential mechanism of active genome demethylation, although more evidence is needed. Gene regulation by such oxidative demethylation is critical for cellular differentiation and embryogenesis. For example, TET-mediated demethyla-

Figure 1. Logic of analogue-sensitive (as) TET enzymes. Conserved bulky amino acids are replaced to introduce hole-modified silent mutations in the active site. The mutants, although acting like wild-type TETs, could be specifically inhibited by “bumped” N-oxalylglycine (NOG) analogues that do not perturb native 2KG-dependent enzymes.
tion on the paternal chromosome is essential to early development of a fertilized egg. Aberrant gene regulation by TET overexpression and somatic mutation are implicated in multiple human conditions. Interrogating developmentally essential and clinically relevant TET members with distinct as well as compensatory functions in fluctuating cellular environments constitutes a major challenge.

Intriguing chemical transformations (iterative C–H functionalization under ambient conditions) coupled with asymmetric demethylation (paternal over maternal pronuclei) and the causal role in human disease have generated widespread interest in studying how a specific TET protein reprograms our genome in diverse physiological contexts. Diagramming a TET-specific gene regulatory network with genetic perturbation is difficult owing to its slow acting nature and oftentimes loss of the entire protein. Pharmacological perturbation involving small molecules provides a powerful alternative to dissect protein functions with high temporal control. However, attempts to develop small-molecule probes for TET enzymes with cellular activity have not met with much success as of yet. Gaining specificity from such molecules would be challenging due to the similarity in active site fold, cofactors, and co-substrates among 60 2KG-dependent human oxygenases.

To address these challenges, we envisioned developing orthogonal mutant—inhibitor pairs of TET enzymes by engineering a steric component of the TET–2KG interface. In this approach, known as analogue-sensitive chemical genetics or the bump-and-hole tactic, the active site of TET enzymes is modified by replacing bulky residues with alanine or glycine (Figure 1). The catalytically competent TET variants with expanded active site are then screened for selective inhibition by 2KG-competitive molecules. The mutant-specific inhibitors are obtained from N-oxalylglycine (NOG), a nonspecific inhibitor of 2KG-utilizing enzymes, by appending complementary steric bulk at a suitable position (Figure 1). Some of these NOG analogues are expected to be refractory to the wild-type enzymes due to a potential steric clash with the larger amino acids in the active site, thus providing chemical probes to interrogate a specific TET member. We selected NOG as the inhibitor scaffold because of its ability to bind the TET enzymes in a 2KG-competitive manner due to structural similarity. The presence of an amido acid functionality in NOG (in place of the ketoacid in 2KG) prevents its catalytic processing by the enzymes, thus making it an effective inhibitor. In this study, we describe our efforts toward development of the first analogue-sensitive TET–NOG pairs and their application in probing transcriptional activity of a specific TET member in human cells.

## RESULTS AND DISCUSSION

### Development of the Orthogonal TET2–NOG Pair

We selected TET2 as a representative DNA demethylase here because of its pivotal role in global methyl erasure during embryogenesis, its established link to cancer, and the availability of critical structural information. In order to develop TET2 variants with an expanded active site, we analyzed the crystal structure of TET2 bound with NOG and 5mC-containing DNA and identified several bulky residues that line up to hold 2KG in the active site (Figure 2a). Given that the majority of these amino acids are hydrophobic in nature and do not participate in strong electrostatic interactions, either with 2KG or DNA, we performed alanine scanning on these sites, one at a time. Mutant proteins carrying the TET2 catalytic domain were expressed and subjected to a MALDI-based C–H hydroxylation assay using a 5mC-containing short DNA and the cofactor 2KG 1 (Figure 2b). The wild-type TET2 and most of the mutants were capable of accepting 1 as their cofactor to generate 5hmC DNA albeit at varying degrees (Figures 2b, S1). F1377A and Y1902A mutants were however completely inert, likely due to the combined effect of active site misfolding and compromised DNA binding. Loss of H-bonding between 5mC and Y1902 may contribute to the catalytic inability of the Y1902A mutant.

This screening data provided us with a platform suitable for developing an analogue-sensitive TET2 enzyme, reminiscent of the protein kinase mutants where engineering a bulky gatekeeper residue sensitized the mutants toward “bumped” inhibitors. To obtain such allele-specific inhibitors, we synthesized a series of NOG analogues, 3–8, bearing alkyl and aryl moieties with variegated steric bulk at C4 of NOG 2 (Figure 2c, Schemes S1–S3). Next, we examined the inhibitory activity of the NOG analogues toward TET2 and its mutants using the MALDI-based hydroxylation assay (Figures 2c, S2, S3). Wild-type TET2 and the majority of its mutants were inhibited by NOG 2, reflecting their ability to bind and process 2KG 1 as cofactor. However, with the exception of the V1900A and V1935A mutants, which were moderately inhibited by the ethyl and propyl NOGs 3 and 4, respectively, the majority of the mutants remained inert toward the bulky analogues (Figures 2c, S4).

Careful analysis of the NOG-bound TET2 structure revealed that the isopropyl side chains of V1935 and V1900 reside very close (within 4.8 Å) to the C4 methylene group carrying the bulky substituents in NOG analogues. This led us to generate the V1935G and V1900G mutants with a slightly larger active

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**Figure 2.** Development of analogue-sensitive (a) TET enzymes. (a) Structure of TET2 bound with 5mC-DNA and NOG (PDB: 4NM6) showing conserved bulky hydrophobic amino acids. The selected residues are targeted for mutagenesis. (b) Bar diagram showing activity of selected alanine mutants of TET2 (% 5hmC formation) as judged by MALDI MS analysis. (c) Heat-map representation of % inhibition of selected hole-modified TET2 mutants by a set of NOG analogues. The best mutant T1393S/V1395G is referred to as TET2.

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site with intact demethylase activity (Figure S5). In the demethylation assay, multiple NOG analogues inhibited the V1395G mutant albeit to varying degrees (Figures 2c, S6). The smaller NOGs such as 4 and 5 inhibited the mutant more effectively. The analogues did not inhibit the V1900G mutant to a significant degree (Figures 2c, S7). Given that removing the steric block at V1395 site sensitizes TET2 mutants toward multiple inhibitors, we reasoned that expanding the active site further might help in gaining stronger inhibition by the bulkier NOGs. We therefore introduced a second site mutation to V1395G to generate multiple double mutants. The majority of the mutants lost their catalytic activity toward 5mC DNA. Remarkably, the T1393S/V1395G mutant retained its DNA demethylase activity as demonstrated by the hydroxylation assay (Figure S5). Furthermore, the double mutant was strongly blocked by NOG analogues 4–6 (Figures 2c, S8).

Leucine NOG 5 appeared to be particularly effective against the mutant, demonstrating the importance of complementary steric engineering to develop analogue-sensitive (as) TET2.

**Characterization of the Orthogonal TET2–NOG Pair.** We next sought to assess the inhibitory efficacy of NOG analogues (4–6) by determining their IC50 values against the selected TET2 variants. The MALDI-based hydroxylation assay is suitable for providing quantitative information on TET enzymatic activity because of the similar ionization potentials of SmC- and SmhC-containing DNAs. The propyl NOG 4 showed a better IC50 against the double mutant T1393S/V1395G (hereafter as TET2) compared to V1395G (120 vs 473 μM), consolidating the importance of the expanded TET2 active site for inhibitor potency of NOG analogues (Figure S9–S11). From our dose–response experiments, we found leucine NOG 5 to be the most effective inhibitor for as TET2, with an IC50 of 18 μM (Figures 3a, S12, S13). This compound did not show any inhibition toward wild-type TET2 in a wide range of concentrations (Figures 3a, S14), thus making it suitable for allele-specific perturbation of TET2.

To rationalize the binding potency and specificity of 5 to as TET2, we performed docking experiments using AutoDock energy minimization software. For this purpose, we used the coordinates from the NOG-bound TET2 crystal structure (PDB 4NM6). In the most stable conformation, 5 occupied the site of NOG 2 and maintained interactions with the as mutant in a manner similar to that of 2 with wild-type TET2 (Figure S15). Most importantly, in the energy-minimized model, the isobutyl side chain of 5 occupied the hole created by V1395G mutation in the as variant and induced a steric clash at the active site of the wild-type protein, which explains the observed specificity (Figure 3b,c). We further noticed that the S1393 residue in the mutant makes an extra H-bonding contact with the amido acid group in 5, potentially contributing to the improved inhibitor efficacy against as TET2 (Figure 3d). Overall, the docking study suggests that V1395G and T1393S mutations in as TET2 may contribute differentially to the analogue sensitivity, with the former bringing more steric complementarity and the latter more electrostatic affinity.

To further ensure that the “bumped” NOGs do not inhibit other human 2KG-dependent enzymes, we examined a set of structurally and functionally distinct demethylases: TET3 (DNA demethylase), FTO and ALKBH5 (RNA demethylases), KDM4A, C and KDM6B (histone demethylases). These wild-type proteins efficiently acted on their respective substrates in the presence of cofactor I and were inhibited by canonical NOG 2 but not the “bumped” NOG analogue 5 (Figures S16–S22). The fact that all seven native demethylases, including TET2, examined thus far remained refractory toward 5 lends further support for the orthogonality of the inhibitor across human 2KG-dependent enzymes.

Next, we examined the activity of the mutant—inhibitor pair on complex substrates (Figure 4a). We collected genomic DNA from human embryonic kidney 293T (HEK293T) cells. In a dot-blot assay, using the appropriate antibody, the isolated genomic DNA showed a robust level of methylated cytidine (5mC) and an almost undetectable amount of ShmC, consistent with minimal expression of the TET enzymes in these cells (Figure 4b). Hydroquinone treatment, however, induced ShmC formation in the genomic DNA of cultured cells, likely via activation of the endogenous TET enzymes as reported earlier (Figure S23). Upon incubating the genomic DNA with wild-type and as TET2 along with the cofactor I, we observed significant formation of ShmC (Figure 4b,c,d) in both cases, suggesting that the hole-modified mutant is capable of acting on a complex substrate with efficiency comparable to that of the wild-type protein. More importantly, leucine NOG 5 completely abrogated the formation of ShmC by as TET2 but not the wild-type enzyme, which was inhibited only by NOG 2. Collectively, these results demonstrate that the mutant— inhibitor pair is truly orthogonal and functional on a physiologically relevant substrate.

**Development of Analogue-Sensitive TET1 and TET3 Variants.** After demonstrating that as variant can be successfully developed for TET2, we sought to expand the technology to other members of the family. In mammals, the TET family consists of three members, TET1–3, each with distinct functions. Sequence analyses of TET oxygenases revealed that the amino acids corresponding to T1393 and
acids prevent binding of the KDM4A and TET2, respectively, reside in close proximity to analogue sensitivity for the 2KG-dependent human enzymes. priciple toward engineering these gatekeeper sites to achieve residues. Our studies disclose a potentially general design analogues to the wild-type proteins by acting as gatekeeper followed by a Chemoenzymatic functionalization of 5hmC modi...powerful method to map the genomic distribution of cytosine-3, respectively, as their V1685G and T1088S/V1090G double mutants of TET1 and anologues as observed for TET2. We generated T1683S/V1395 in TET2 are highly conserved among TET1...Activity of Figure 4. Activity of as TET2—NOG 5 pairs on the genomic DNA. (a) Schematic showing isolation of genomic DNA from human cells followed by enzymatic hydroxylation of SmC by as TET2 and its inhibition by 5 and subsequent biochemical analysis. (b) Dot-blot assay showing activity of wild-type TET2 on the genomic DNA. The activity was inhibited by NOG 2 but not by “bumped” NOG 5. (c) Dot-blot assay showing activity of as TET2 on the genomic DNA, which was inhibited by the “bumped” NOG 5. (d) Quantitative representation of the dot-blot results obtained in (b) and (c). (e) Sequencing data showing T1393 and V1395 of TET2 are fully conserved among other TET members. (f, g) Dot-blot assays showing activity of as TET1 and TET3, respectively, on the genomic DNA and their inhibition by NOG 5.

V1395 in TET2 are highly conserved among TET1–3 (Figure 4e). These bulky amino acids are very likely acting as gatekeeper residues that preclude binding of bulky NOG analogues as observed for TET2. We generated T1683S/V1685G and T1088S/V1090G double mutants of TET1 and -3, respectively, as their as variants. When examined for activity, the TET1 and -3 mutants efficiently produced ShmC in genomic DNA as measured in the dot-blot assay (Figure 4fg), confirming their catalytic activity. We subsequently showed that the as variants were selectively inhibited by NOG analogue 5 (Figure 4fg). Collectively, our results provide strong evidence for the existence of a bulky gatekeeper residue (an equivalent of V1395 in TET2) within the TET subfamily. Mutations at these sites led to, for the first time, analogue-sensitive DNA demethylases, suitable to deconvolute their specific functions. We have shown earlier that hole-modifying F185G mutation of KDM4A, a 2KG-dependent histone demethylase, sensitizes the mutant toward a series of bulky 2KG cofactor analogues. F185 and V1395 residues in KDM4A and TET2, respectively, reside in close proximity to the cofactor 2KG in the respective active sites. These amino acids prevent binding of the “bumped” cofactor/inhibitor analogues to the wild-type proteins by acting as gatekeeper residues. Our studies disclose a potentially general design principle toward engineering these gatekeeper sites to achieve analogue sensitivity for the 2KG-dependent human enzymes.

Functionlization of TET-Specific Activity on Genomic DNA. Chemoenzymatic functionalization of ShmC followed by affinity enrichment and sequencing has become a powerful method to map the genomic distribution of cytosine modifications and hence TET activity. A particular approach includes β-glucosyltransferase (β-GT)-mediated glycosylation of ShmC present in genomic DNA with 6-azidoglucose-UDP followed by “click” ligation with the appropriate reporter alkyne. We sought to combine the as TET variants with the chemoenzymatic modification of ShmC that would potentially allow direct probing of genome-wide activity of a specific TET member (Figure 5a). ShmC generated on a short DNA substrate by the as TET2 mutant underwent smooth glycosylation to 6-azidoglucose-UDP using β-GT followed by Cu-catalyzed “click” ligation with biotin-alkyne 9, as confirmed by MALDI MS (Figures 5b, S24, Scheme S4). However, no such ligated product was obtained when the TET2 assay was performed in the presence of “bumped” NOG 5 (Figure 5b), thus correlating the formation of biotinylated DNA to the activity of as TET2.

To further assess the ability of the engineered pair to modulate the tandem hydroxymethylation–biotinylation process in a genome-wide manner, the entire genomic DNA isolated from HEK293T cells was treated with the as TET2 variant with or without 5. The modified DNA was subjected to β-GT-mediated azidoglucosylation followed by “click” chemistry with 9. The ligated product was directly analyzed by dot-blot assay using antibody (Figure 5c). The chemiluminescent signal from the noninhibited sample was significantly reduced signal was observed when the TET assay was performed in the presence of NOG 5. These results provide a blueprint for the potential application of as TET enzymes in probing genome-wide activity of a specific TET member when coupled to next-generation sequencing.

Allele-Specific Perturbation of TET Activity in Cultured Human Cells. The three human TET proteins, each with distinct structural features, have compensatory as well as nonoverlapping functions in cells. We sought to examine the effectiveness of the NOG-based inhibitors in modulating demethylase activity of the analogue-sensitive TET...
enzymes in intact human cells. Toward this end, we synthesized octyl ester derivatives 10 and 11 of NOG 2 and leucine NOG 5, respectively. It has been shown that esterification of 2KG and 2HG (2-hydroxyglutarate) improves their cell permeability (Figure 6a, Scheme S5).32−44

Upon induced expression of the wild-type (or hole-modified) human TET2 catalytic domain in HEK293T cells, as confirmed by Western blot (Figure S25), the genomic DNA was isolated and subjected to dot-blot assay. A robust level of ShmC was observed only in cells expressing the enzymes and not in nontransfected cells, clearly demonstrating that both the enzymes are catalytically active in cells (Figure 6b,c). We observed a reduction in ShmC level, albeit to a smaller extent, when cells expressing wild-type TET2 were treated with 100 and 250 μM of 10 (Figure 6b). This is consistent with the weaker inhibitory activity of NOG 2 (IC_{50} ≈ 150 μM) toward wild-type TET2.26 However, no change in ShmC content was noticed when these cells were cultured in a concentration of the ester produg 11 was enough to reduce the ShmC level to a significant degree (~75%) in these cells, consistent with the stronger inhibitory activity of 5 (IC_{50} ≈ 18 μM) toward as TET2. The lack of hydroxylase activity of as TET2 in these cells must arise from allele-specific inhibition by the NOG analogue 11, as full recovery of TET2 activity ensued upon withdrawal of the compound from cell culture (Figure S26). These results demonstrate that the demethylase activity of TET2 variants on chromosomal DNA can be specifically and conditionally manipulated using a cell-permeable and reversible “bumped” NOG analogue in intact cells.

We also examined off-target activity of the NOG analogue 11 on 2KG-dependent endogenous histone demethylases. We extracted nuclear histones from HEK293T cells treated with 100 μM 11 and analyzed for four key trimethylated lysine residues in histone H3 (H3K4Me3, H3K9Me3, H3K27Me3, and H3K36Me3) using corresponding antibodies. Upon comparison with untreated cells, we did not observe any significant changes in histone methylation levels at these sites (Figure S27). This result further confirms the in cellulo selectivity of the “bumped” NOG analogue, making it suitable to study only the analogue-sensitive variant in human cells.

Gene regulatory function of TET enzymes is directly related to their catalytic ability to generate 5hmC in a loci-specific manner. Overexpression of TET2 is commonly associated with gene activation through the formation of ShmC at promoters and enhancers. We sought to examine whether the as TET2 and the “bumped” inhibitor pair could modulate expression of certain genes in cells. We focused on the epithelial cellular adhesion molecule (EpCAM) and transducin beta like 1 X-linked (TBL1X) proteins, as it has been shown that shRNA-mediated knockdown of TET2 leads to the silencing of both EPCAM and TBL1X genes primarily via the loss of ShmC at the promoter.35,46 Consistently, overexpression of TET2 leads to reactivation of EPCAM and TBL1X proteins. Nuclear extracts from HEK293T cells expressing the as TET2 variant and treated with or without 100 μM 11 were collected and analyzed for the expression of both the proteins using appropriate antibodies. Overexpression of as TET2 led to an increase in EPCAM and TBL1X protein levels compared to the control cells lacking as TET2 (Figures 6d,e, S28). These results confirm that as TET2 is functionally competent in cells and required for expression of certain genes much like its wild-type counterpart. We observed, however, more than a 3-fold reduction in EpCAM and TBL1X proteins when as TET2-expressing cells grew in the presence of the “bumped” inhibitor 11 (Figures 6d,e, S28). Expression of housekeeping proteins such as GAPDH remained unaffected in all the samples. Collectively, our results show that bulky NOG inhibitor can modulate the gene regulatory functions of a specific TET member in human cells. Our study further demonstrates that selective inhibition of the catalytic activity of TET2 is sufficient to silence expression of certain genes.

■ CONCLUSION

The diversity-based mechanism of protein evolution, in which many protein products originate from relatively few genes, is central to the formation of closely related protein families. This has significantly hindered the development of isoform/memberspecific chemical probes. In this piece of work, we employed structure-guided protein−ligand interface engineering to overcome the barrier imposed by active site degeneracy. TET mutants with a “hole” at the active site while still maintaining wild-type catalytic activity are specifically inhibited by designed NOG analogues carrying a complementary...
“bump”. These compounds do not interfere with the native TETs and a range of other 2KG-dependent enzymes and hence can serve as chemical probes for functional analysis of TET enzymes. Such specificity likely stems from a steric clash between the hydrophobic “bump” of the inhibitors and the conserved bulky gatekeeper residue in TET enzymes, as we have identified in the current study. Engineering at these gatekeeper residues in TET proteins extended the approach to all three members of the subfamily. Following successful probe development, we demonstrated their ability to conditionally manipulate 5hmC formation on genomic DNA. Furthermore, manipulation of gatekeeper residues in TET proteins extended the approach to work is dedicated to Prof. Goverdhan Mehta.

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Synthetic procedure and characterization of all the relevant compounds, methods for protein expression and purification, and biochemical assay, supplementary figures and tables, and NMR spectra (PDF)

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**Notes**

The authors declare no competing financial interest.

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