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Real-Time Monitoring of Human Guanine Deaminase Activity by an Emissive Guanine Analog

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provide a visible spectral window for a real-time study of GDA and its inhibition.

INTRODUCTION

Nucleobase and nucleoside deaminases play key roles in cellular pathways. Being responsible for the cellular levels of purine- and pyrimidine-based heterocycles, their over (or under) expression is frequently related to pathological conditions.^{1–4} Additionally, differences between human and bacterial enzymes provide intriguing opportunities for new therapeutic approaches, exploiting their distinct fidelity and inherent activities.¹

Guanine deaminase (GDA), a hydrolytic zinc-based enzyme, converts guanine to xanthine.^{5,6} Recent observations have shown it impacts neuronal morphology^{7,8} and implicated GDA in traumatic brain injury,⁹ memory dysfunction, and psychiatric diseases.¹⁰ The study of human GDA, from the aminohydrolase superfamily, has remained somewhat sparse compared to other deaminases, including adenosine deaminase (ADA)^{11,12} and cytidine deaminase (CDA),¹³ which convert adenosine to inosine and cytidine to uridine, respectively. As a result, real-time activity and inhibitory assays for GDA, especially the human isoform, have yet to be advanced, and its substrate scope remains somewhat elusive.¹⁴

Diverse methods have been utilized for assessing the enzymatic activity and inhibition of deaminases. Most protocols capable of directly measuring GDA activity rely on the distinct chromatographic behavior or the UV absorption signature of the substrate/product pair.¹⁵ The former techniques are accurate but can be time-consuming, while the latter tools frequently suffer from spectral interference as many inhibitors share similar chromophoric properties with the substrate or product. To facilitate the biochemical study of deaminases, define their substrate scope, and identify

inhibitors, innovative methods for monitoring their activity are needed. Although fluorescence spectroscopy might *a priori* appear most attractive due to its high sensitivity, it cannot be applied in such contexts due to the lack of any useful emissive features of the native purine nucleobases.

Over the past two decades our laboratory has designed, synthesized, and implemented minimally perturbing and responsive fluorescent nucleoside analogs. $^{15-29}$ The guiding criterion for their design has been the diminution of structural and functional perturbations, which are inevitable consequences of modifying any native building block. Nucleosides that fulfill such critical constraints are thus coined isomorphic. If they function identically to their native counterparts, we define them as being isofunctional. We have advanced two emissive RNA alphabets, based on thieno[3,4-d]pyrimidine (1st generation; ${}^{\text{th}}\mathbf{N})^{30}$ and isothiazole[4,3-d]pyrimidine (2nd generation; ${}^{tz}N$)³¹ heterocyclic cores (Figure 1a). The emissive adenosine analogs have been successfully used to advance realtime activity and inhibition assays of adenosine deaminase, exploiting the distinct emission profiles of the substrates (thA and ^{tz}A), and their corresponding deamination products (thI and ^{tz}I, respectively), with the latter (^{tz}A) being essentially isofunctional (Figure 1b).^{32,33} The red-shifted absorption and

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Figure 1. a) Heterocycle evolution pertaining to this study starting from the native purine core to thN to ^{tz}N. b) Adenosine deaminase conversion of A to I and isofunctional nucleoside analog ^{tz}A to ^{tz}I. c) Guanine deaminase conversion of G to X, isomorphic nucleobase analog thG_N to thX_N, and isofunctional nucleobase analog ^{tz}G_N to ^{tz}X_N.

visible-range emission of such fluorescent substrate surrogates opens a spectral window not accessible by other means, which facilitates the real-time monitoring of these reactions even in the presence of potentially interfering chromophores.³⁴

Applying such approaches to nucleobase-processing enzymes such as human GDA represents, however, a minimally explored territory, as highly emissive nucleobase analogs have not been broadly tested and scrutinized as substrates.^{6,} ¹³ The lack of the rather large and contact forming D-ribose residue presents the challenge of altering guanine without disturbing contacts at the active site of such a potentially fastidious nucleobase-processing enzyme.35 We therefore set out to investigate two emissive nucleobase analogs, thieno[3,4*d*]pyrimidine (thG_N) and isothiazole[4,3-*d*]pyrimidine (^{tz}G_N) (Figure 1c), and demonstrate that while the former is not a viable substrate, ${}^{tz}G_N$ does undergo facile GDA-mediated deamination to yield the fluorescently distinct xanthine analog, $t^{tz}X_{N}$. The spectral differences displayed by the substrate and product are exploited for a real-time monitoring of the enzymatic reaction. Moreover, the potential of ${}^{tz}G_N$ as a tool for GDA screenings is showcased by measuring GDA inhibition by a previously reported inhibitor and identifying two new inhibitors. Taken together, these findings highlight the necessity of the N7 for substrate recognition by GDA and provide information regarding its substrate and inhibitor scope.

RESULTS AND DISCUSSION

The four nucleobases, containing either a thienopyrimidinecore, ${}^{th}G_N$ and ${}^{th}X_{N}$, or an isothiazolepyrimidine-core, ${}^{tz}G_N$ and ${}^{tz}X_{N}$, were prepared according to established procedures (Figure 2a, Schemes S1–S3).^{30,31,35,36} Briefly, treatment of commercially available methyl 4-aminothiophene 3-carboxylate hydrochloride with chloroformamidine hydrochloride in DMSO₂ at 125 °C yields ${}^{th}G_N$ in one step. The corresponding xanthine analog, ${}^{th}X_N$, was synthesized in two steps by treating the same starting material with KOCN to yield the corresponding urea, which was then cyclized under basic conditions with sodium methoxide in methanol (Scheme S1).



Figure 2. a) Structures of ${}^{th}G_N$, ${}^{th}X_N$, ${}^{tz}G_N$, and ${}^{tz}X_N$ nucleobases. b) Absorption (dashed lines) and emission (solid lines) of ${}^{th}G_N$ (green), ${}^{th}X_N$ (red), ${}^{tz}G_N$ (blue), and ${}^{tz}X_N$ (orange) in water. c) Stokes shift correlations versus solvent polarity $[E_T(30)$ of water/dioxane mixtures] for ${}^{th}G_N$ (green), ${}^{th}X_N$ (red), ${}^{tz}G_N$ (blue), and ${}^{tz}X_N$ (orange). d) Change in optical density versus pH for ${}^{th}G_N$ (green), ${}^{th}X_N$ (red), ${}^{tz}G_N$ (blue), and ${}^{tz}X_N$ (orange).

For the isothiazole-based nucleobases ${}^{tz}G_N$ and ${}^{tz}X_{N}$, methyl 4aminothiazole 3-carboxylate hydrochloride, a key precursor, was prepared from methyl thioglycolate and N-tosylated Oxyma, followed by decarboxylation and esterification (Scheme S2).³⁷ Conversion of the isothiazole-based precursor to ${}^{tz}G_N$ and ${}^{tz}X_N$ followed the same cyclization procedures described above for the thiophene derivatives (Scheme S3). Crystal structures have been determined for all nucleobases (Figure 3a-d, Tables S1–S4).



Figure 3. X-ray crystal structures of nucleobases. a) thG_N b) thX_N c) ^{tr}G_N and d) ^{tz}X_N. MOE docking of e) xanthine as reference ($\Delta\Delta G 0$), f) ^{tz}X_N ($\Delta\Delta G 0.28$), and g) thX_N ($\Delta\Delta G 1.89$) in GDA active site (PDB ID 2UZ9, see Methods).

Table 1. Photophysical Data for Nucleobase Analogs

	$\lambda_{\mathrm{abs,max}}^{a}$	ε^{b}	$\lambda_{\rm em,max}^{a}$	ϕ
${}^{\mathrm{th}}\mathbf{G}_N$	315	3.0	439	0.40
${}^{\mathrm{th}}\mathbf{X}_N$	308	3.2	420	0.46
$^{tz}G_N$	320	5.4	446	0.07
$^{tz}X_N$	315	6.1	394	0.02
		1		

 ${}^a\lambda_{\rm abs,max}$ and $\lambda_{\rm em,max}$ are in nm. ${}^b\varepsilon$ is in 10³ M^{-1} cm⁻¹. Measured in triplicate in water.

To assess the likelihood of spectroscopically distinguishing between the substrates and their corresponding products, the fundamental spectroscopic properties of the nucleobases were determined (Tables 1 and S5). Absorption spectra in water displayed red-shifted maxima of the nucleobase analogs compared to their native counterparts, showing maxima at 315, 308, 320, and 315 nm for thG_N, thX_N, ^{tz}G_N, and ^{tz}X_N, respectively. Excitation at their absorption maxima gave rise to visible emission, which peaked at 439, 420, 446, and 394 nm for thG_N, thX_N, ^{tz}G_N, and ^{tz}X_N, respectively (Figure 2b). As expected, the emission quantum yields of the thiophene analogs, thG_N and thX_N (0.40 and 0.46, respectively) were higher than their isothiazole counterparts, ^{tz}G_N and ^{tz}X_N (0.07 and 0.02, respectively.)

Additionally, the sensitivity of photophysical parameters to environmental polarity was evaluated, and spectroscopically derived pK_{a} values were determined (Table S5). The absorption and emission parameters of each nucleobase were measured in dioxane, water, and mixtures thereof. By linearly correlating the calculated Stokes shift against the solvent polarity of each sample, the chromophore's responsiveness was defined. The four nucleobases showed differing levels of sensitivity to polarity, with the thiophene derivatives showing substantial impact on Stokes shift compared to the isothiazole derivatives (Figure 2c, Figure S1). The nucleobases also showed responsiveness to pH changes. pK_a values were extrapolated by plotting the change in optical density versus pH (Figure 2d, Figure S2). The guanine derivatives, ${}^{th}G_N$ and $^{tz}G_N$, show two p K_a values (p $K_a = 4.41$, 10.19 and p $K_a = 3.28$, 8.96, respectively), while the xanthine derivatives, ${}^{th}X_N$ and ^{tz}**X**_N, show one pK_a value ($pK_a = 9.92$ and $pK_a = 8.8$, respectively), as expected.^{31,38,39}

To compare the susceptibility of ${}^{th}G_N$ and ${}^{tz}G_N$ to GDAmediated deamination, enzymatic reactions were first analyzed by HPLC and then monitored in real-time by absorption and emission spectroscopy. HPLC analysis confirmed the complete transformation of native guanine to xanthine by GDA within 500 s (Figure 4a and S3a). The reaction of ${}^{tz}G_N$ with GDA showed conversion to the corresponding product, ${}^{tz}X_N$ (Figures 4a and S3b). However, the reaction of ${}^{th}G_N$ showed no conversion of substrate across all time points studied (Figure S3c). By plotting the area under the curve at different reaction times (Figure 4b), comparable reaction half-life values for guanine deamination (G $t_{1/2}$ = 27 s) and ^{tz}G_N deamination $({}^{tz}\mathbf{G}_{N} t_{1/2} = 21 \text{ s})$ were calculated assuming pseudo-first-order reaction conditions (Table 2). Kinetic parameters were also calculated by fitting HPLC-monitored deamination reactions to a set of ordinary differential equations consistent with Michaelis-Menten kinetics (see Methods). Similar values were obtained for guanine ($K_{\rm M} = 12 \pm 9 \ \mu \text{M}$; $k_2 = 36 \pm 27 \ \text{s}^{-1}$; $k_2/K_{\rm M} = 3.0 \pm 0.3 \ \mu \text{M}^{-1} \ \text{s}^{-1}$) and its isothiazolo analog ^{tz}**G**_N ($K_{\rm M} = 8 \pm 3 \ \mu \text{M}$; $k_2 = 32 \pm 7 \ \text{s}^{-1}$; $k_2/K_{\rm M} = 4.2 \pm 1.0 \ \mu \text{M}^{-1} \ \text{s}^{-1}$; see Table 2). We note that the $K_{\rm M}$ value obtained for the



Figure 4. a) Conversion of guanine to xanthine and conversion of isofunctional ^{tz}G_N to ^{tz}X_N by GDA. b) HPLC relative peak area versus time for consumption of guanine to xanthine at 260 nm (blue) and consumption of ^{tz}G_N to ^{tz}X_N at 320 nm (red). Inset: HPLC relative peak area versus time for the consumption of guanine (blue) and formation of xanthine (green) at 260 nm and the consumption of t^{tz}G_N (red) and formation of t^{tz}X_N (orange) at 320 nm from 0 to 100 s. c) Absorption changes upon enzymatic deamination of guanine to xanthine at 270 nm (blue) and t^{tz}G_N to t^{tz}X_N monitored by absorbance (at 355 nm, red), fluorescence (450 nm, excitation at 328 nm, green), and HPLC relative peak area over time (at 320 nm, blue).

deamination of guanine is comparable to previously published figures for rabbit liver GDA ($K_{\rm M}$ of 12.5 μ M),⁴⁰ and the $k_2/K_{\rm M}$ is comparable to $k_1/[{\rm GDA}]$ (G $k_1/[{\rm GDA}] = 2.6 \ \mu$ M⁻¹ s⁻¹) from the pseudo-first-order kinetic curves (Table 2).

Rewardingly, spectroscopic analyses agree with the chromatographic analysis (Figure 4c). When monitoring the enzymatic deamination of ${}^{tz}G_N$ spectroscopically, an increase in absorbance at 355 nm and an emission decrease at 450 nm when exciting at 328 nm, the isosbestic point, were observed. The reaction half-life for ${}^{tz}G_N$ was calculated using the same assumptions mentioned previously (${}^{tz}G_N$ abs $t_{1/2} = 27$ s; $t_{1/2}$ em = 29 s) and was found to be comparable to the native substrate (G abs $t_{1/2} = 31$ s), as shown in Table 2. The transformation of both native guanine to xanthine and ${}^{tz}G_N$ to ${}^{tz}X_N$ by GDA is also illustrated by the absorption maxima shift (Figure S4). Thienopyrimidine ${}^{th}G_N$ showed no change in absorption or emission in the presence of GDA over 500 s.

Reactions monitored by HPLC as well as real-time absorption and emission spectroscopy suggest that ${}^{tz}G_N$, the isothiazolepyrimidine substrate, is equally susceptible to GDAmediated deamination as native guanine. Intriguingly, ${}^{th}G_N$, the thienopyrimidine guanine analog, is unreactive and does not undergo GDA-mediated deamination. These findings showcase the functionality and increased isomorphicity of the isothiazolepyrimidine-based nucleobase alphabet, which reinstated the N7 moiety, compared to the thienopyrimidine-based nucleobases, even without the large and contact forming Dribose residue the nucleoside counterparts possess. Further support is obtained by Molecular Operating Environment

Table 2. GDA Reaction Rates

	HPLC				absorbance			emission				
	$\lambda_{\rm mntr}^{a}$	k_1^{b}	$t_{1/2}^{c}$	$K_{\rm M}^{d}$	k_2^d	k_2/K_M^d	$\lambda_{\rm mntr}^{a}$	k_1^{b}	$t_{1/2}^{c}$	$\lambda_{\rm mntr}^{a}$	k_1^{b}	$t_{1/2}^{c}$
G to X	260	26 ± 2	27	12 ± 9	36 ± 27	3.0 ± 0.3	270	22 ± 1	31			
${}^{\mathrm{th}}\mathbf{G}_N$ to ${}^{\mathrm{th}}\mathbf{X}_N$	320	NR	NR	NR	NR	NR	355	NR	NR	450 ^e	NR	NR
${}^{\mathrm{tz}}\mathrm{G}_{N}$ to ${}^{\mathrm{tz}}\mathrm{X}_{N}$	260	32 ± 1	21	8 ± 3	32 ± 7	4.2 ± 1.0	355	25 ± 1	27	450 ^f	24 ± 1	29

 ${}^{a}\lambda_{mntr}$ reported in nm and represents wavelength monitored. b Pseudo-first-order reaction kinetics slope of the exponential approximation in 10^{-3} s⁻¹. ${}^{c}t_{1/2}$ is reaction half-life calculated assuming pseudo-first-order kinetics. ${}^{d}K_{M}$, k_{2} , and k_{2}/K_{M} are reported in μ M, s⁻¹, and 10^{6} M⁻¹ s⁻¹, respectively. ${}^{e}\lambda_{exc}$ at 360 nm for ${}^{th}G_{N}$ reactions. ${}^{f}\lambda_{exc}$ at 328 nm for ${}^{tx}G_{N}$ reactions. Experiments done in triplicate.

(MOE) molecular docking experiments (Figures 3e–g, Figures S5 and S6) using the published crystallographic data for xanthine-bound human GDA (PDB ID 2UZ9). The enzyme recognition preferences for ^{tz}X_N over thX_N are illustrated by the markedly higher positive $\Delta\Delta G$ values obtained for the latter vs the former ($\Delta\Delta G$ 1.89 and 0.28, respectively). A similar trend was observed for the substrates (Figure S6). Docking also shows a plausible structure deformation induced by Arg235, which is projected toward the N7 position at the native substrate. This basic site, present in native G/X and the isothiazolo analogs, ${}^{tz}G_N/{}^{tz}X_N$, likely hydrogen bonds to Arg235. Clashing of this arginine side chain with the CH group of ${}^{th}G_N$ at the same position likely renders the thiophenopyrimidine-based substrate unrecognizable by GDA.

As stated, the presence of the nitrogen at a position equivalent to N7 in the purine skeleton endows $^{tz}G_N$, the isothiazole purine surrogate, with substrate recognition features that appear lacking in $^{th}G_N$. While comprehensive investigations of the GDA substrate scope have not been pursued, previous substrate and inhibitor analyses of amino-hydrolase isoforms suggest that the O6, N3, and N7 points of contact are important for substrate recognition. 6,17,40,41 Intriguingly, while $^{tz}G_N$ retains these contact points, it is lacking the NH group found in the purine's 9 position. Previously reported inhibitors, such as valciclovir and derivatives of azepinomycin, containing substituents at the N9 position suggest GDA can tolerate diverse groups at that position, which may explain the high tolerance and the native deamination rate displayed by $^{tz}G_N$.

The change in emission signal intensity upon GDAmediated deamination of ${}^{tz}G_N$ to ${}^{tz}X_N$ facilitates real-time monitoring of the enzymatic reaction and its inhibition. To demonstrate its potential, three inhibitors were tested, 5aminoimidazole-4-carboxamide (AICA), 4-aminoisothiazole-3carboxamide (ATCA), and 4-imidazolecarboxylic acid (ICA, Figure 5a). AICA is an established competitive inhibitor of GDA.⁴⁰ We synthesized ATCA, the isothiazole counterpart (Scheme S4). Reaction conditions used for previous experiments were sustained with the exception of inhibitor addition at various concentrations. Percent inhibition plots were fitted with sigmoidal Hill curves to obtain IC₅₀ values of the tested inhibitors. AICA (IC₅₀ = 100 μ M) showed comparable inhibition to ATCA (IC₅₀ = 80 μ M), while ICA is comparatively a less potent inhibitor (IC₅₀ = 2 mM, Figure 5b).

CONCLUSION

This study set out to assess the utility of two emissive nucleobase analogs, ${}^{th}G_N$ and ${}^{tz}G_N$, as substrates for the enzymatic reaction of GDA. We demonstrate that isothiazole-[4,3-d]pyrimidine, ${}^{tz}G_N$, behaves as an isomorphic and isofunctional emissive surrogate for guanine. The interconver-



Figure 5. a) GDA inhibitor structures of AICA, ATCA, and ICA. b) Overlay of % inhibition at 100 s and sigmoidal Hill plot for AICA (black), ATCA (red), and ICA (blue).

sion of ${}^{tz}G_N$ to ${}^{tz}X_N$ can successfully be tracked in real-time as the reaction substrate and product display distinct photophysical properties. The strategic replacement of the GDA substrate with emissive counterparts from both previously published emissive alphabets demonstrate the significance of N7 for GDA substrate recognition. Moreover, this discovery provides a biochemical tool to study the activity of human GDA and offers a spectral window for fabricating real-time high throughput discovery assays of GDA. We illustrate this through a small pilot study of inhibitor effects on GDA activity, which includes one previously reported inhibitor and two new inhibitors.

METHODS

Expression and Purification of Human GDA. Wild-type human GDA gene, cloned into pET-28B vector (Genescript), was expressed with an N-terminal hexahistidine tag. The construct was sequence-verified (GENEWIZ). The plasmid (4 μ g) was dissolved in water to a concentration of 28.8 ng/ μ L. *E. coli* BL21 (DE3) cells were transformed with the GDA containing plasmid. Overnight starter cultures were grown in Lubria Broth (LB) medium at 37 °C with 50 μ g/mL kanamycin. The overnight culture was used to inoculate 1 L of LB medium with 50 μ g/mL kanamycin. Cells were grown while shaking at 37 °C until an OD₆₀₀ of 0.6 was reached, and the culture was left to shake at 18 °C overnight. Cells were harvested after overnight growth by centrifugation at 3500 rpm for 35 min at 4 °C.

Cells were resuspended in lysis buffer [20 mM HEPES, 500 mM NaCl, 10% glycerol, 0.5 mM TCEP (pH 7.5), supplemented with one tablet of Complete EDTA-free protease inhibitor (Roche Applied Science)] and lysed by sonication. Cell lysate was centrifuged at 10,000 rpm for 1 h at 4 °C before the supernatant was decanted and filtered through a 0.45 μ m syringe filter. GDA was purified by immobilized metal affinity chromatography. An IMAC Ni-charged column [2 mL HisPur Ni-NTA Resin (Thermo Scientific)] was equilibrated with lysis buffer to which the lysate was then added. The column was washed with 50 mL of wash buffer supplemented with 20

mM imidazole. The bound protein was eluted from the column with elution buffer (wash buffer supplemented with 200 mM imidazole). Fractions were analyzed by SDS-PAGE, and those containing the target protein were pooled, subsequently concentrated, and buffer exchanged into storage buffer [20 mM Tris-HCl buffer (pH 8.0), 10% glycerol, 1 mM DTT] using a centrifugal filter device with a 10 kDa molecular weight cutoff. After protein expression and purification, the final protein concentration was 5.3 mg mL⁻¹ (0.1 mM) in a total volume of 1.7 mL. Protein aliquots were snap-frozen and stored at -80 °C until further use. Before being used in enzymatic reactions, the GDA protein stock was diluted 1:100 in sodium phosphate buffer [50 mM (pH 7.4)].

Synthetic Procedures. All synthetic procedures and characterizations of compounds are reported in the Supporting Information.

Photophysical Properties: General Methods. Absorption spectra were measured on a Shimadzu UV-2450 spectrometer with 0.5 mm resolution and setting the slit at 1 nm. Emission spectra were measured on a Horiba Fluoromax-4 equipped with a cuvette holder and stirring system. Emission measurements were taken with resolution at 1 nm and setting both the excitation and emission slits at 3 nm.

All measurements were carried out in a 10 mm four-sided quartz cuvette purchased from Helma. All spectra were corrected for the blank. Both instruments were equipped with a thermostat-controlled ethylene glycol-water bath, and all measurements were taken at 37 $^{\circ}$ C. Measurements were recorded after a 3 min temperature equilibration period.

Concentrated stock solutions for xanthine, ${}^{th}G_{N}$, ${}^{th}X_{N}$, ${}^{tz}G_{N}$, and ${}^{tz}X_{N}$ were prepared in DMSO, and a stock solution of guanine was prepared in water basified to pH 12 with sodium hydroxide. Samples for experiments were prepared with stock nucleobase diluted to a total sample volume of 3 mL in deionized water, mixed with a pipet for 10 s, and placed in the cuvette holder. All samples contain 0.3 v/v % DMSO, except guanine samples.

Quantum Yield Measurements. All sample concentrations were adjusted to optical density lower than 0.1 at the excitation wavelength (λ_{ex}) . The fluorescence quantum yield (ϕ) of each nucleobase was evaluated based on 2-aminopurine (0.68 in water, λ_{ex} 320 nm) as an external standard by using the following equation

$$\phi = \phi_{\text{STD}} \frac{I}{I_{\text{STD}}} \frac{\text{OD}_{\text{STD}}}{\text{OD}} \frac{n^2}{n_{\text{STD}}^2}$$
(1)

where ϕ_{STD} is the fluorescence quantum yield of the standard, *I* and I_{STD} are the integrated area of the emission band of the sample and the standard, respectively, OD and OD_{STD} are the optical density at the excitation wavelength for the sample and standard, respectively, and *n* and n_{STD} are the solvent refractive index of the sample and the standard solutions, respectively.

Sensitivity to pH. Sodium phosphate buffers with a final concentration of 50 mM were prepared and adjusted to the desired experimental pH values using HCl or NaOH prior to spectral measurements. Changes in optical density, at 310 and 350 nm, were plotted versus pH. The pK_a values were determined by interpolation of the fitting curves.

Sensitivity to Polarity. Experiments evaluating the effect of polarity were performed in water, dioxane, and mixtures of 20, 40, 60, and 80 v/v % water in dioxane. The sample $E_{\rm T}(30)$ values were determined by dissolving Reichardt's dye in an aliquot of the same solvent used to dilute the nucleobase DMSO sample. The observed wavelength absorption maximum $(\lambda_{\rm abs}^{\rm max})$ was then converted to the $E_{\rm T}(30)$ values (Table S6) using the following equation

$$E_{\rm T}(30) = \frac{28591}{\lambda_{\rm abs}^{\rm max}} \tag{2}$$

Enzymatic Deamination: General Methods. Reaction conditions were the same for all GDA reactions monitored by spectroscopy or chromatography. Concentrated stock solutions for xanthine, ${}^{th}G_N$, ${}^{th}X_N$, ${}^{tx}G_N$, and ${}^{tx}X_N$ were prepared in DMSO, and the stock solution of guanine was prepared in water basified to pH 12

with sodium hydroxide. Samples were prepared in a 10.00 mm foursided quartz cuvette from Helma. Reactions had a total reaction volume of 3 mL with nucleobase and enzyme concentrations of 3 μ M and 10 nM in sodium phosphate buffer [50 mM (pH 7.4)]. All measurements were taken at 37 °C, and GDA was introduced after a 3 min temperature equilibration period.

Real-Time Monitoring of GDA Reactions via Absorption and Emission. GDA-mediated conversion of guanine and its analogs (${}^{th}G_N$ and ${}^{tz}G_N$) was monitored by absorption and emission spectroscopy. Absorbance measurements of the enzymatic conversion of guanine (and its analogs) were performed on a Shimadzu UV-2459 spectrometer taking a point every 10 s for 600 s after the addition of GDA with a slit setting of 1 nm. The conversion was monitored at 270 nm for guanine to xanthine and at 355 nm for the analogs (${}^{th}G_N$ and ${}^{tz}G_N$ to ${}^{tz}X_N$). Emission measurements of the conversion of guanine (and analogs) were performed on a Horiba Fluoromax-4 with a cuvette holder and a built-in stirring system with excitation and emission slits set to 3 nm and taking points every 10 s for 600 s after the addition of GDA.

Emission was monitored at 450 nm with excitation at 360 and 328 nm, respectively, for ${}^{\rm th}G_N$ and ${}^{\rm tz}G_{N^*}$. The ${}^{\rm th}G_N$ reaction showed no change in absorption or emission intensity in the presence of GDA over 600 s. Each experiment was done in triplicate. There is a 6 s lag time after GDA is added to the cuvette after the time 0 measurement.

Steady State Absorption Measurements in the Presence of GDA. Steady state absorption spectra over time were performed on a Shimadzu UV-2450 spectrophotometer setting the slit at 5 nm, using a resolution of 0.5 nm, taking a measurement every 20 s. All spectra were corrected for the blank. Smoothing of data was done on spectra for plotting.

HPLC Analysis of Enzymatic Conversion of Native G to X and of ${}^{tz}G_N$ to ${}^{tz}X_N$. GDA-mediated conversion was monitored by chromatography. HPLC was carried out with an Agilent 1200 system with a Polaris 5 C18-A 250 × 4.9 mm column. Solutions of 0.1% formic acid (Honeywell Fluka) were prepared by dissolving 1 mL of formic acid in 1 L total volume of acetonitrile (J. T. Baker) or water. Solutions were filtered using Millipore type GNWP 0.2 μ m filters before use. Each injection (75 μ L) for guanine or ${}^{tz}G_N$ experiments was subjected to a linear gradient of 0.5% to 10% acetonitrile in water with 0.1% formic acid for 20 min, followed by a flush and equilibration for 10 min. ${}^{th}G_N$ injections were subjected to a linear gradient of 0.5% to 10% acetonitrile in water with 0.1% formic acid for 30 min, followed by a flush and equilibration for 10 min. Each run was monitored at 260, 280, and 320 nm with a calibrated reference at 650 nm and slit set at 1 nm.

A concentrated stock solution of guanine, ${}^{\text{th}}\mathbf{G}_N$, and ${}^{\text{tz}}\mathbf{G}_N$ was diluted in phosphate buffer. The solution was warmed to 37 °C for 3 min with stirring before addition of a GDA stock solution. After the addition of GDA, the enzymatic conversion was quenched in aliquots (after 20, 40, 60, 80, 100, 200, and 500 s) by adding formic acid (0.55 M) and placing the aliquots on ice. Each 100 μ L aliquot was filtered and analyzed by HPLC.

HPLC traces were corrected for the blank, and the relative areas were plotted as a function of time. Trend lines represent a loss of substrate and product apparition over time for a pseudo-first-order kinetic reaction.

A set of ordinary differential equations (ODEs) consistent with Michaelis–Menten kinetics (eqs 3–6) was solved using the Runge–Kutta method with a method with a variable time step in MatLab (function ode45). Initial concentrations used for enzyme and substrate were 10 nM and 3 μ M, respectively. The product and enzyme substrate complex were assumed to have initial concentrations of 0 μ M. The resulting fitted curves for each species were optimized by iteratively testing k values that maximized R.² This yielded k_1 , k_{-1} , and k_2 values from which $K_{\rm M}$ and $k_2/K_{\rm M}$ values were derived.

$$\frac{d[E]}{dt} = -k_1[E][S] + k_{-1}[ES] + k_2[ES]$$
(3)

$$\frac{d[S]}{dt} = -k_1[E][S] + k_{-1}[ES]$$
(4)

$$\frac{d[ES]}{dt} = k_1[E][S] - k_{-1}[ES] - k_2[ES]$$
(5)

$$\frac{\mathrm{d}[P]}{\mathrm{d}t} = k_2[ES] \tag{6}$$

Inhibition Studies. Fluorescence real-time monitoring of the enzymatic conversion of ${}^{tz}G_N$ to ${}^{tz}X_N$ with GDA in the presence of inhibitors was followed by emission spectroscopy by monitoring the intensity signal at 450 nm with excitation at 328 nm. 5-Aminoimidazole-4-carboxamide (AICA) was tested as an inhibitor in the enzymatic reaction for ${}^{tz}G_N$ to ${}^{tz}X_N$ using the same reaction conditions but supplemented with inhibitor concentrations of 0.1, 1, 10, 100, and 1000 μ M. 4-Aminoisothiazole-3-carboxamide (ATCA) was also tested as an inhibitor in the reaction under the same conditions but supplemented with inhibitor concentrations of 0.1, 1, 10, 100, 250, and 500 µM. 4-Imidazolecarboxylic acid (ICA) was also tested as a potential inhibitor under the same reaction conditions with 10, 100, 1000, 2000, and 3000 μ M concentrations. IC₅₀ values were determined from plots of percent inhibition against respective inhibitor concentration on a logarithmic axis, fitted with sigmoidal Hill curves. All measurements were done in triplicate.

Docking of Guanine, Xanthine, and Analogs in GDA. Docking simulations were performed with the Molecular Operating Environment (MOE) 2020 software suite. The crystal structure of human guanine deaminase (guaD) in complex with zinc and its product xanthine was obtained from the Protein Data Bank (PDB ID 2UZ9). The crystal structure was chosen because it contains the native product xanthine. The chosen ligands were placed by the Triangle Matcher method and ranked with the London dG values scoring function. A total of 30 poses were refined using the rigid receptor method. Then five poses were rescored using a GBVI/WSA dG scoring function. The pose with the ligand most closely aligned with the xanthine ligand was used for comparison, and the $\Delta\Delta G$ value was calculated for ${}^{tz}G_N$ and ${}^{th}G_N$ with native guanine as the reference and for ${}^{tz}X_N$ and ${}^{th}X_N$ with native xanthine as the reference. The equation below was used where $\Delta G_{\rm ref}$ is the GBV1/WSA dG scoring of the reference native ligand and ΔG_{lig} is the GBVI/WSA dG scoring of the ligand analogs (eq 7).

$$\Delta\Delta G = \Delta G_{\rm lig} - \Delta G_{\rm ref} \tag{7}$$

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acschembio.1c00232.

Synthetic schemes, experimental methods, photophysical data, spectra, and X-ray crystallographic data for ${}^{th}G_{N'}$ ${}^{th}X_{N'}$ ${}^{tx}G_{N'}$ and ${}^{tx}X_N$ (PDF)

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The manuscript was written through contributions of all authors. All authors have given approval to the final version of the manuscript.

Notes

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