Artificial Genetic Systems: Exploiting the “Aromaticity” Formalism To Improve the Tautomeric Ratio for Isoguanosine Derivatives

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Abstract: The tautomerism of 2′-deoxy-7-deaza-isoguanosine (2) was studied and compared to that of 2′-deoxy-isoguanosine (1). The fixed N-methyl (8) and O-methyl (4) derivatives were synthesized to represent the pure extremes of each tautomer. The replacement of the imidazole ring in 1 with a pyrrole ring in 2 makes the keto form in the latter more favored by 2 orders of magnitude (K\text{TAUT} for 2 ≈ 10^3, as opposed to K\text{TAUT} for 1 ≈ 10).

It has been a decade since it was shown that the geometry of the Watson–Crick nucleobase pair can accommodate as many as 16 nucleobases forming eight mutually exclusive pairs, to yield an artificially expanded genetic information system (AEGIS).1 By providing molecular recognition on demand in aqueous solution, similar to nucleic acids but with a coding system that is orthogonal to the system in DNA and RNA, AEGIS today enables clinical assays such as Bayer’s branched DNA diagnostics tool that monitors the load of viruses in patients infected with HIV and hepatitis C viruses.2 Both assays are FDA-approved and are widely used to provide clinical assays such as Bayer’s branched DNA diagnostics tool that monitors the load of viruses in patients infected with HIV and hepatitis C viruses.2 Both assays are FDA-approved and are widely used to provide personalized patient care in the clinic. Further, AEGIS components enable an assay for the early detection of the SARS virus.3 With the emergence of the first six-letter PCR,4 AEGIS now supports the development of a synthetic biology where higher order processes of living systems, including reproduction and Darwinian evolution, are duplicated by artificial, designed chemical systems.

One issue remaining before a synthetic biology based on AEGIS is fully implemented arises from the tautomeric form displayed by isoguanosine (and 2′-deoxyisoguanosine, disoG, 1). In its major keto form (Figure 1), disoG implements a hydrogen bond donor–donor–acceptor pattern (proceeding from the major to the minor groove) on a purine heterocycle (Figure 1). A minor enolic tautomter of disoG is known to be present in aqueous solution to perhaps 10%,5 however. The enolic tautomer presents the donor–acceptor–donor hydrogen bonding pattern that is complementary to the thymidine and uridine heterocycles.6 Although the presence of the minor tautomer does not adversely affect the use of isoguanosine in clinical diagnostics, it does inconvenience some polymerases that prefer to place thymidine (T) and uridine (U), rather than isocytidine (isoC), opposite isoguanosine in a template.

Given that the keto forms of pyrimidinones normally dominate over enolic forms, we viewed the enolic tautomer of disoG as an exception in need of explanation. We reasoned that the enolic form was present in appreciable concentration in disoG because it restores formal aromaticity to the five-membered imidazole ring, which must be cross-conjugated in the keto tautomer (Figure 1). It is well-known that pyrrole is less “aromatic” than imidazole, with the former being 59% as aromatic as benzene, the latter being 64% (compared to pyridine at 86%, thiophene at 66%, and furan at 43%).8 This suggested that replacement of the imidazole ring of disoG by a pyrrole ring to give 7-deaza-isoguanine might generate a purine analogue that would also implement the pUDDA hydrogen bonding pattern (like disoG), but with less of the enolic tautomter present at equilibrium.

7-Deaza-isoguanine (C7isoG) and its 2′-deoxy nucleoside analogue (dC7isoG, 2) are both known.9-11 The crystal structure of dC7isoG12 and its base-pairing ability have both been studied.13-15 The tautomisation of both C7isoG and dC7isoG remain undetermined, however.

FIGURE 1. Isoguanosine (1), a purine (pu), can generate two possible tautomers, an enol tautomer (giving a puDAD form, which will pair with T or U) or a keto tautomer (giving a pUDDA form, which will pair with isoC). The extent of this tautomisation has been reported to be 10:1 in favor of the keto form.

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To test our hypothesis, the O-methyl-7-deaza-isoguanine derivative was prepared by the procedure of Davoll.\textsuperscript{16} Reaction with 1-(\(\alpha\))-chloro-3,5-di-O-(p-toluoyl)-2-deoxy-o-ribose\textsuperscript{17} gave the 3',5'-ditoluoyl-protected 2'-deoxyriboside of 2-methoxy-6-amino-7-deaza-purine (3). Seela's procedure\textsuperscript{9} for the removal of the 2-methoxy group (HCl, 1,4-dioxane, 100 °C) did not work well in our hands (Scheme 1). We therefore used thiophenol in ethylene glycol\textsuperscript{18} to generate the di-O-(p-toluoyl)-derivative of 2 (5). This compound was converted to the 2'-deoxynucleoside using NaOMe in MeOH to afford 2, which had spectroscopic properties identical to those reported for this compound by Seela et al.\textsuperscript{9} The structure of this compound was further confirmed by treatment with diazomethane, which generated 4'. The structure of 4' was identical in all aspects to that of 4. The N-methyl derivative was prepared by protection of the exocyclic amino group as the diisobutylformamidine\textsuperscript{19,20} (generating 6), reaction with CH\textsubscript{3}I (to give 7), and immediate deprotection by treatment with methanolic NH\textsubscript{3}/NaOMe, affording 8. N-Methylation at the 1N position was confirmed by long-range (HMBC\textsuperscript{21}) and short-range (HMQC\textsuperscript{22}) \(^1\)H--\(^{13}\)C NMR correlation spectroscopy.

The pK\textsubscript{a} of the nucleoside analogue (2) was determined by spectrophotometric titration (pH 2.8 to 13.7)\textsuperscript{23} at 240–350 nm to give values for pK\textsubscript{1} and pK\textsubscript{2} as 4.3 (±0.1) and 9.9 (±0.2), respectively (Supporting Information). These are comparable to the pK\textsubscript{a} values of natural nucleobases and ensure that the C7isoG heterocycle is largely uncharged in neutral water.

Four tautomeric forms are worthy of consideration for 7-deaza-isoguanine (Scheme 2). On the basis of literature precedent, as well as precedent in other heterocycles, the imino form was considered to be the least likely.\textsuperscript{5,24–26}

\begin{figure}
\centering
\includegraphics{scheme1.png}
\caption{Scheme 1}
\end{figure}

\begin{figure}
\centering
\includegraphics{scheme2.png}
\caption{Scheme 2}
\end{figure}

This was confirmed by NMR: the -NH₂ group was easily identified in the NMR by integration and D₂O exchange.

The UV spectra of 2'-deoxy-7-deaza-isoguanosine (2) in aqueous solution resembles closely that of the N-methylated derivative (8) and not the O-methylated derivative (4) (Figure 2). This similarity suggests but does not prove that the keto tautomer predominates, following the same logic used by Sepiol et al. to infer that the keto tautomer predominates for isoguanosine.⁵

The multiwavelength analysis described by Dewar and Urch²⁷ was used to suggest that, at most, one additional tautomer was present as a major contributor to the mixture (Supporting Information).

The procedure of Voegel et al.²⁸ was then used to estimate a value of the keto:enol tautomers in pure water for dC₇isoG. This procedure exploits the fact that in nonpolar solvents the UV spectra for both disoG and C₇isoG shift from their form in pure water (which resembles the N-methylated derivatives) to a form that resembles the UV spectra of the O-methylated derivatives. This is interpreted as evidence that the tautomeric equilibrium shifts from one favoring the keto tautomer in pure water to one favoring the enol tautomer in pure dioxane.

To obtain quantitative data, the ratio of the extinction coefficients at two wavelengths (λ = 296 and 255 nm for disoG and λ = 305 and 254 nm for dC₇isoG; Supporting Information), chosen to maximize the difference between the keto and enol forms, was chosen as a metric for the tautomeric equilibrium constant in dioxane/water mixtures in varying proportions. This was plotted against the Dimroth E₁(30) value, which provides a measure of the local dielectric constant.²⁹–³³

Even qualitatively, disoG and dC₇isoG behave differently in these experiments. The UV spectrum of disoG changes well before the water is completely removed. In contrast, the UV spectrum of dC₇isoG is not identical to that of the O-methylated derivative even at the highest concentrations of dioxane tested. Further, the UV spectrum of disoG continues to change even in mixtures approaching pure water; this suggests that the conversion of the enolic tautomer to the keto tautomer of disoG is not complete even in pure water (the conclusion of Sepiol et al.⁵). With dC₇isoG, however, the UV spectrum ceases to be solvent-dependent when the fraction of water is greater than 50%. These results suggest that our hypothesis at outset had manipulative value: the keto tautomer of dC₇isoG is more stable relative to its enolic tautomer than the keto tautomer of disoG is relative to its enolic tautomer.

We then estimated the Kᵣᵤₜₑᵦ (= [keto]/[enol]) for dC₇isoG using the method of Voegel et al.²⁸ Here, the fraction of enol form was estimated over a range of E₁(30) values where the Kᵣᵤₜₑᵦ ≈ 1, regions where an

FIGURE 2. UV spectra of compounds 2, 4, and 8 in water. (Inset) UV spectra of the same compounds, in 99:1 1,4-dioxane/water.

estimate could be made with some reliability. This is the region where \( E_{T(30)} \approx 40-50 \). The log fraction of enol was plotted against \( E_{T(30)} \) (Figure 3), and the line was extrapolated to the \( E_{T(30)} \) of pure water (63.1). This generated a value of \( K_{\text{TAUT}} = 10^3 \). Recalculating the \( K_{\text{TAUT}} \) for isoG gave a value of \( 10^{10} \), consistent with that previously reported. This can be compared with the value for the natural guanosine nucleoside (\( K_{\text{TAUT}} = 10^{4-10^5} \)).

With this result, we have now fixed each of the problems identified in AEGIS components exploiting the full range of hydrogen bonding patterns. Adjusting the electron-withdrawing groups has generated a pyDDA derivative that is free of epimerization. Adjusting side chains of the pyAAD system has managed deamination issues. Adjusting the ring nitrogens has corrected the problematic pKₐ values of bases of acidic purines. In the work presented here, adjustment of the aromaticity of rings has resolved a tautomerism issue. Thus, with this paper, we have the ingredients for a fully functional 12-letter genetic alphabet that meets the specifications met by natural nucleic acids.

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**Supporting Information Available:** Experimental procedures, characterization data, HPLC traces, and experimental details for the pKₐ measurements and tautomerization data. This material is available free of charge via the Internet at http://pubs.acs.org.

(45) It is clear that the \( E_{T(30)} \) value at the interior of the DNA helix is not known; our current work is looking into whether the behavior of dC7isoG in water will correlate to its behavior in DNA.