Guanine to Inosine Substitution Leads to Large Increases in the Population of a Transient G·C Hoogsteen Base Pair

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Supporting Information

ABSTRACT: We recently showed that Watson–Crick base pairs in canonical duplex DNA exist in dynamic equilibrium with G(syn)-C′ and A(syn)-T Hoogsteen base pairs that have minute populations of ~1%. Here, using nuclear magnetic resonance (NMR) R_{i;p} relaxation dispersion (RD)6–7 and single-atom substitution experiments that G-C and A-T Watson–Crick (WC) base pairs (bps) transiently form Hoogsteen (HG) bps with populations of ~0.4 and 0.7% and lifetimes of 1.7 and 0.3 ms at pH ~5.4 (Figure 1A). HG bps form through a 180° rotation of the purine base around the glycosidic bond from an anti to a syn conformation (Figure 1A). By modifying the structural and chemical presentation of DNA and modulating nucleobase accessibility, HG bps can play unique roles (reviewed in ref 9) in DNA–protein recognition,10–13 DNA damage induction14 and repair,15,16 and replication.17,18

We provided evidence of transient HG bps based on analysis of R_{i;p} RD experiments, which quantify the degree of line broadening due to chemical exchange with a transient low-population species.7 A two-state analysis of the RD data measured on a variety of G-C and A-T Watson–Crick bps in canonical duplex DNA was consistent with the existence of a transient species that features downfield-shifted purine base C8 and sugar C1′ chemical shifts. These chemical shifts are a hallmark of syn purine, where the base flips 180° about the glycosidic bond. Such a syn purine can be paired with pyrimidine via HG pairing. By using chemical modification, including N1-methyladene (N1-Me-A) and N1-methylguanine (N1-Me-G), which are naturally occurring damaged forms of purine bases, we successfully trapped HG bps in duplex DNA and showed that they have the characteristic downfield-shifted carbon chemical shifts.1 As further support for the assignment of HG bps for the observed transient state, we subsequently showed that the single-atom substitution, 7-deazaguanine and 7-deazaadenine, which specifically knocks out an HG H-bond between guanine N7 and cytosine N3 (N7···H···N3) without affecting WC H-bonds, destabilizes the transient state, such that it can no longer be detected by NMR RD.2

Here, we sought to shift the WC–HG equilibrium in the other direction, toward the HG state. In G-C bps, a transition toward HG bps preserves the O6–H···N4 H-bond present in WC pairing but leads to the loss of the N1–H···N3 and N2–H···O2 H-bonds, which are replaced with a single N7···H···N3 H-bond that further requires protonation of cytosine N3 (Figure 1). Therefore, one would expect that, in inosine, the loss of the exocyclic amino group at position C2 of guanine would lead to the loss of a WC H-bond without affecting HG...
pairing and, consequently, lead to an increase in the population of the transient state observed using NMR RD.

To examine the impact that replacement of guanine in the CA step with inosine would have on the WC–HG equilibrium, we prepared an A_{C9}-DNA duplex (A_{C9}-DNA\textsuperscript{10}) (Figure 2A), in which G10 was replaced with I10. We previously reported detailed NMR characterization of transient HG bps in A_{C9}-DNA as well as variants that bear chemical modifications and mutations at the same position.\textsuperscript{1–3} Because of difficulties in preparing isotopically \textsuperscript{13}C/\textsuperscript{15}N-labeled oligonucleotides with a modified inosine base that are required for obtaining high-quality off-resonance RD data, the strand complementary to the inosine-containing strand was \textsuperscript{13}C- and \textsuperscript{15}N-labeled. The two-dimensional \textsuperscript{13}C–\textsuperscript{1}H heteronuclear single-quantum coherence NMR spectra of this selectively labeled strand were in very good agreement with spectra of A_{C9}-DNA indicating formation of a WC-like I-C bp that does not impact neighboring bps (Figure S1 of the Supporting Information). Because I10 could not be isotopically enriched and the unlabeled strand concentration was not high enough (~0.5 mM), making it difficult to measure reliable R_{1ps} RD data at natural abundance, the complementary labeled C15 residue was used to probe chemical exchange at the I-C bp. In particular, formation of a G-C H G bp is accompanied by a significant downfield shift of the C-C6 chemical shift due to protonation of C-N3, and this can be used to probe the WC–HG equilibrium.\textsuperscript{3} Here, replacement of G10 with I10 resulted in a measurable downfield chemical shift change toward the transient HG state as well as significant line broadening at C15-C6, consistent with increased WC–HG exchange at the I-C bp.

To quantify transient HG bp formation, we collected on- and off-resonance \textsuperscript{13}C RD dispersion on C15-C6 in A_{C9}-DNA\textsuperscript{10}. We observed substantially larger RD for C-C6 in the I-C bp than for C-C6 or G-C8 in the G-C bp under similar conditions (Figure 2B).\textsuperscript{1} A two-state analysis of the RD data allowed for determination of all chemical exchange parameters, including the transient state chemical shift $\Delta \omega_{RB(C6)} = \omega_{RB(C6)} - \omega_{AL(C6)}$, population ($p_B$), and lifetime ($\tau_B = 1/k_B$) (Figure S2 and Table 2 of the Supporting Information). The transient state chemical shift for C15-C6 is ~2.3 ppm downfield-shifted relative to the G8, consistent with formation of a HG bp, and is in excellent agreement with the downfield shift (~2.3 ppm) obtained when the G10-C15 bp in A_{C9}-DNA is changed to a HG bp using guanine N1-methylation.\textsuperscript{7}

The magnitude of the C15-C6 downfield chemical shift is similar to that previously seen in C-C6 (~2.2 ppm) or G-C8 (~3.7 ppm) in transient G-C HG bps\textsuperscript{1} and cannot explain the RD observed in the I-C bp being much larger than that in the G-C bp. Rather, the larger RD arises primarily from a substantial 17-fold increase in the population of HG bps in I-C ($p_B \approx 11.8\%$) as compared to G-C ($p_B \approx 0.7\%$). This is in accord with our expectations based on the loss of WC H-bonds without significantly affecting HG H-bonds. In particular, the relative stability of the WC bp is decreased by ~1.8 kcal/mol in I-C compared to that in G-C. This is consistent with previous studies showing an ~2 kcal/mol loss of B-DNA stability from a single G-C to I-C substitution\textsuperscript{9,20}, and could be explained by the loss of a single N-H–O bond or replacement of the N-H–O bond with a weaker C-H–O bond.\textsuperscript{21} The increase in the HG population is accompanied by an increase in chemical exchange rate $k_{ex}$ ($k_{ex} = k_1 + k_2$ increases ~6-fold from ~600 s\textsuperscript{-1} in G-C to 3500 s\textsuperscript{-1} in I-C), which corresponds to an ~100-fold increase in the forward rate constant $k_1$ and a much smaller ~5-fold increase in the reverse rate constant $k_2$. This translates into a simultaneous decrease in the free energy difference between WC and HG states ($\Delta G_{WC-HG}$) and the free energy required to reach the transition state ($\Delta G^2_{WC-HG}$), which is in turn consistent with selective destabilization of WC due to the inosine substitution. We recently reported a similar trend based on $\psi$ value analysis of WC-to-HG transitions measured across diverse DNA sequence and position contexts, where $k_1$ increased in sequences with more abundant HG bps and $k_2$ exhibited a much weaker sequence dependence.\textsuperscript{3} These results implied that the variations in the relative energetic stabilities of HG bps are dominated by variations in the stabilities of WC bps, as demonstrated by a direct correlation between $\Delta G_{WC-HG}$ and $\Delta G^2_{WC-HG}$.\textsuperscript{4} Importantly, the observed correlation of sequence-dependent $\Delta G$ values for WC bps extrapolates to the $\Delta G$ values for the modified I-C bp (Figure 2C) and strongly supports our earlier proposal that WC bp stabilities, unlike HG bp stabilities, are highly sequence-dependent.

The 5-fold increase in $k_2$ could arise from destabilization of the HG bp, possibly because of the loss of H-bonding with solvent\textsuperscript{11,21} or, alternatively, stabilization of the TS due to removal of the bulky amine group and more facile rotation of the purine base within the helix. While the structure of the TS remains unknown, most crystal structures of HG bps display H-bonds between G-NH$_2$ and the backbone phosphate or water molecules. The loss of the NH$_2$ group in inosine could contribute to the small (<1 kcal/mol) decrease in the stability of the transient HG I-C$^\ast$ versus G-C$^\ast$ state.

As a control, we conducted on-resonance \textsuperscript{13}C RD experiments at natural abundance on inosine C8 in an I-T bp,

Figure 2. Increased level of $R_{1ps}$ RD and HG bp formation at I-C. (A) A_{C9}-DNA highlighting the A-T or G-C bp substituted with inosine at the purine base. (B) On-resonance $\textsuperscript{13}C$ $R_{1ps}$ RD profiles comparing chemical exchange in G-C, I-C, and I-T bps (pH 5.2 for G-C C-C6 and pH 5.4 for all others). Solid lines represent best global fits to eq S1 of the Supporting Information. (C) Correlation between $\Delta G_{WC-HG}$ and $\Delta G_{WC-HG}$ for WC bps\textsuperscript{4} (data for the I-C bp colored blue). Shown are the best-fit line and corresponding Pearson coefficient ($R$) with (---) or without (—) inclusion of the I-C bp.
which should not be capable of stably forming HG bps. This is because only one H-bond can form in the I-T HG bp without additional tautomerization and/or protonation. Indeed, we observed no evidence of chemical exchange at I-C8 (Figure 2B), supporting the idea that the observed transient state results from specific pairing of inosine with an appropriate HG pairing partner.

The results presented here provide additional evidence in support of transient HG bps in a canonical duplex and also show that HG bps can exist in much greater abundance in I-C bps. Inosine is a rare form of mutagenic damage that is recognized and excised by conserved repair enzymes such as DNA glycosylases and endonuclease V. It is conceivable that the HG-type bps participate more broadly in damage recognition and repair, as could be the case for G-A and A-C mispairs where the partially exposed HG face of the syn-purine is specifically recognized by the DNA mismatch repair enzyme MutS or for the common oxidative damage 8-oxoguanine, which can form a stable HG-type bp with adenine. HG-type bps involving inosine have already been implicated as the source of mutagenic substitutions, most frequently from A-T to G-C. For example, syn-inosine has been observed to form an HG-type bp with protonated adenine at near-neutral pH, which affords a possible mode of recognition for I-A mispairs. The observation of more abundant HG bps in I-C also raises the possibility that other forms of damage and chemical modifications may increase the abundance of these bps in genomes where they can potentially conduct unique biological functions.

ASSOCIATED CONTENT

Supporting Information
Methods, Tables 1 and 2, and Figures S1 and S2. This material is available free of charge via the Internet at http://pubs.acs.org.

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Funding
This study was supported by National Institutes of Health Grant GM089846 awarded to H.M.A.-H.

Notes
The authors declare no competing financial interest.

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