| Title | Insights into the Structures of DNA Damaged by Hydroxyl Radical: Crystal Structures of DNA Duplexes Containing 5-Formyluracil |
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Hydroxyl radicals are potent mutagens that attack DNA and convert them to many different kinds of base and ribose derivatives [1, 2]. Every aerobic organism possesses several enzymes to remove such toxic oxides, as well as to recover the damaged DNA. However, when an excess amount of the radicals attacks DNA, the thymine base is oxidized at the 5-methyl group to form 5-formyluracil base (hereafter 2′-deoxy-5-formyluridine residue is referred to as fU) as a major product. (The four characters, A, T, G, and C, represent the respective nucleotide residues in DNA sequence.) It was demonstrated that 2′-deoxy-5-formyluridine triphosphate (dUTP) was incorporated against both A and G templates, possibly forming fU:A and fU:G base pairs during in vitro DNA replication [3, 4]. On the other hand, it was reported that dUTP-induced pyrimidine transitions, G:C → A:T and A:T → G:C, as well as a gene transversion from G:C to T:A, could occur in vivo [5, 6]. These results suggest that fU can behave as C, A, and G in addition to its original property of T.

1. Introduction

Hydroxyl radicals, activated from hydrogen peroxide and hydrogen peroxide anion under light radiation, are well known as potent mutagens that attack DNA and convert them to many different kinds of base and ribose derivatives [1, 2]. Every aerobic organism possesses several enzymes to remove such toxic oxides, as well as to recover the damaged DNA. However, when an excess amount of the radicals attacks DNA, the thymine base is oxidized at the 5-methyl group to form 5-formyluracil base (hereafter 2′-deoxy-5-formyluridine residue is referred to as fU) as a major product. (The four characters, A, T, G, and C, represent the respective nucleotide residues in DNA sequence.)
In order to reveal the interaction geometry of the modified base fU, we performed X-ray analyses on fU-containing DNA duplexes. The fU residues were introduced into the self-complementary Dickerson-Drew-type dodecamer sequence, which is expected to be easy to crystallize. The DNAs used in this study are d(CGCGATfUCGCG) and d(CGCGGATfUCGCG) and will be referred to as fUA and fUG, respectively. The fU base faces either an adenine or a guanine at the two sites in each duplex. Four fUA crystals and three fUG crystals were obtained under different conditions. Their crystal structures have been successfully determined at resolutions ranging from 1.5 to 3.0 Å. In the preliminary papers [7, 8], magnesium ion effects on crystallizations of fUA were discussed, but the detailed structure of base-pair formations and its biological significance were not published. In this paper we describe the structures of the base pairs formed between fU and G and between fU and A, based on which the pyrimidine transition induced by the oxidized thymine base will be discussed.

2. Materials and Methods

2.1. Oligodeoxyribonucleotide Synthesis. fUA and fUG were synthesized by the solid phase phosphoramidite method as described previously in [9] and were purified by reverse-phase column chromatography and reverse-phase and anion-exchange HPLCs. After NaIO₄ treatment, the oligonucleotides with fU were further purified by reverse-phase and anion-exchange HPLCs.

2.2. Crystallization and Data Collection. Initial screenings of crystallization conditions were performed using the hanging drop vapor diffusion method, equilibrating 2 μl droplets against 1 ml of the reservoir solution. The optimized conditions for growing the four different crystals of fUA (fUA₁, fUA₂, fUA₃, and fUA₄) and three different crystals of fUG are given in Table 1. As the fUG₁ crystal was too small, two kinds of dyes, Hoechst 33258 and DAPI, were added to stabilize the duplex formation (fUG₂ and fUG₃).

Crystals suitable for X-ray data collections were picked up from their mother liquors using a nylon loop (Hampton Research) and transferred into liquid nitrogen. All X-ray experiments for the seven crystals were performed with synchrotron radiation at the Photon Factory in Tsukuba. Diffraction patterns of the fUA crystals, which were recorded on imaging plates, were processed subsequently using the programs DENZO and Scalepack [10] and those of the fUG crystals, recorded on Quantum 4 CCD, with the program DPS/MOSFLM [11]. Low-resolution data around the 10 Å resolution shell were truncated because they did not fit well into the diffraction profile. The crystal data and the statistics of data collection are summarized in Table 2.

2.3. Structure Determination and Refinement. Initial phases were derived by molecular replacement with the program AMoRe [12] using the atomic coordinates of the corresponding unmodified DNA duplex (PDB ID 355D, see [13]) as structural probes. The molecular structures were constructed and modified on a graphic workstation with the program Coot [14] in the CCP4 program suite [15]. The atomic parameters were refined with the program Refmac [16] in CCP4 [15] with maximum-likelihood techniques, followed by interpretation of an omit map at every nucleotide residue. Newly defined patches were prepared for partially applying...
Figure 1: The final $2|F_o|-|F_c|$ maps around the fU residues in the fUA (a), the fUG (b), and the fUG (c) crystals. The maps are contoured at 1σ level by the program DINO [18]. Those of the fUG crystal are omitted due to similarity to (c). Values indicate possible hydrogen bond distances. Characters N, O, and W indicate nitrogen, oxygen, and water oxygen atoms, respectively.
### Table 2: Crystal data and statistical information on data collection and processing and on structure determination.

| Crystals | fUA¹ | fUA² | fUA³ | fUA⁴ | fUG¹ | fUG² | fUG³ |
|----------|------|------|------|------|------|------|------|
| Wavelength (Å) | 1.00 | 0.90 | 1.00 | 1.00 | 1.00 | 1.00 | 1.00 |
| Beamline (at PF) | BL-18B | BL-6B | BL-18B | BL-6B | BL-18B | BL-18B | BL-18B |
| Oscillation ranges(°) | 5 | 3 | 5 | 5 | 3 | 2 | 2 |
| Frames | 72 | 60 | 34 | 36 | 60 | 90 | 90 |
| Space group | P2₁2₁2₁ | P2₁2₁2₁ | P2₁2₁2₁ | P2₁2₁2₁ | P2₁2₁2₁ | P2₁2₁2₁ | P2₁2₁2₁ |
| a (Å) | 25.8 | 26.0 | 25.2 | 25.3 | 24.6 | 25.3 | 25.0 |
| b (Å) | 39.3 | 39.5 | 41.2 | 41.7 | 40.0 | 40.3 | 40.5 |
| c (Å) | 65.0 | 65.7 | 65.4 | 66.0 | 68.5 | 66.0 | 66.8 |
| Z⁺ | 1 | 1 | 1 | 1 | 1 | 1 | 1 |
| Matthews coefficient (Å³ Da⁻¹) | 2.15 | 2.20 | 2.22 | 2.28 | 2.28 | 2.28 | 2.29 |
| Solvent content (%) | 42.9 | 44.2 | 44.6 | 45.9 | 46.0 | 46.1 | 46.3 |
| Resolution (Å) | 1.57 | 1.55 | 1.85 | 1.80 | 3.0 | 1.95 | 2.6 |
| Completeness (%) | 97.1 | 97.7 | 99.3 | 99.3 | 99.9 | 99.9 | 99.9 |
| R_merge (%) | 3.1 | 4.8 | 2.9 | 2.8 | 11.4 | 4.0 | 6.3 |
| Observed reflections | 87588 | 66227 | 31872 | 45728 | 16654 | 72307 | 38399 |
| Unique reflections | 9502 | 10301 | 7779 | 6881 | 1531 | 5312 | 2342 |
| Redundancy | 9.22 | 6.43 | 4.10 | 6.65 | 10.9 | 13.6 | 16.4 |

*Structure refinement¹*

| Resolution range (Å) | 10.0−1.57 | 10.0−1.55 | 10.0−1.85 | 10.0−1.80 | 15.0−3.02 | 10.0−1.95 | 10.0−2.70 |
| R-factor (%)² | 18.3 | 18.3 | 21.6 | 19.3 | 18.8 | 20.3 | 21.1 |
| Rfree (%)³ | 23.2 | 22.5 | 26.0 | 23.1 | 22.9 | 26.1 | 27.8 |

| Bond distances (Å) | 0.027 | 0.028 | 0.019 | 0.021 | 0.005 | 0.019 | 0.007 |
| Bond angles (°) | 3.8 | 3.6 | 3.2 | 3.4 | 3.4 | 0.9 | 3.1 | 1.0 |
| No. of ions | 1Mg²⁺, 1K⁺ | 1Mg²⁺ | — | — | — | 1Mg²⁺ | 1Mg²⁺ |
| No. of additive molecules | — | — | — | — | — | 1Hoechst33258 | 1DAPI |
| No. of water molecules | 150 | 185 | 82 | 142 | 27 | 123 | 45 |
| PDB-ID | 1G75 | 1G8N | 1G8U | 1G8V | 3AJJ | 3AJK | 3AJL |

¹Number of duplexes in the asymmetric unit. ²R_merge = 100 × ∑ₕ |Iₕ| − ⟨|Iₕ|⟩/ ∑ₕ |Iₕ|, where ⟨|Iₕ|⟩ is the jth measurement of the intensity of reflection h and |Iₕ| is its mean value. ³Two data sets were taken for a crystal by changing exposure time to compensate overloaded reflections. ⁴R-factor = 100 × ∑|Fₒ| − |F_c|/ ∑|F_c|, where |F_c| and |Fₒ| are the observed and calculated structure factor amplitudes, respectively. ⁵Calculated using a random set containing 10% of observations that were not included throughout refinement [17].

2.4. Coordinates. Atomic coordinates and structure factors have been deposited in the Protein Data Bank with accession codes 1G75, 1G8N, 1G8U, 1G8V, 3AJJ, 3AJK, and 3AJL for fUA¹, fUA², fUA³, fUA⁴, fUG¹, fUG², and fUG³, respectively.

3. Results

3.1. Quality of X-ray Analyses. All the crystals are isomorphous to the orthorhombic form of the unmodified duplex crystal. Crystallization of fUA was easy as expected, and the crystals obtained diffracted within the 1.5−1.8 Å resolution range. In every crystal, the formyl group of the fU8 residue on one of the two chains was disordered between the two alternative conformers, anti and syn, while that in the other chain adopted only the syn conformation. Relative occupancies of the disordered oxygen atoms, estimated structural restraints to the modified residue. Water, ion, and dye molecules were assigned and included in the refinements. The program CNS [17] was used in the final refinements of fUG¹ and fUG³ to stabilize the base pairs containing X with hydrogen bonds. The statistics of structure refinements are summarized in Table 2. Examples of the quality of the final electron density maps are depicted in Figure 1. Helical and local base-pair parameters [19], as well as the torsion angles and pseudorotation phase angles of sugar rings [19], were calculated using the program 3DNA [20]. Some of them are shown in Table 3 and Figure 4.
Table 3: A comparison of the local base-pair parameters at the modified pairs, calculated with the program 3DNA [20].

| Pair  | fUA1 (Å) | fUA2 (Å) | fUA3 (Å) | fUA4 (Å) | fUG1 (Å) | fUG2 (Å) | fUG3 (Å) | Ave (Å) | Unm (Å) |
|-------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|---------|---------|
| Shear (Å) fUA:R | 0.01 | −0.03 | −0.05 | −0.08 | −2.00 | −2.24 | −2.19 | −0.94 | −0.11 |
|       R:fUA20 | 0.05 | 0.08 | −0.04 | −0.01 | 1.90 | −2.08 | 2.15 | 0.29 | −0.04 |
| Stretch (Å) fUA:R | −0.19 | −0.13 | −0.08 | −0.10 | 0.15 | 0.00 | 0.08 | −0.04 | −0.12 |
|       R:fUA20 | −0.10 | −0.11 | −0.16 | −0.16 | 0.13 | −0.58 | 0.10 | −0.13 | −0.11 |
| Stagger (Å) fUA:R | 0.10 | 0.11 | −0.06 | 0.08 | −0.33 | −0.08 | 0.03 | −0.02 | −0.00 |
|       R:fUA20 | 0.09 | 0.00 | 0.00 | 0.10 | −0.06 | 0.10 | −0.05 | 0.03 | 0.01 |
| Buckle (°) fUA:R | −6.04 | −4.89 | −2.68 | −4.97 | −5.62 | −10.56 | −9.96 | −6.39 | −1.56 |
|       R:fUA20 | 8.11 | 6.02 | 9.91 | 9.34 | 6.78 | 8.68 | 4.50 | 7.62 | 4.72 |
| Propeller (°) fUA:R | −12.4 | −12.4 | −12.5 | −11.9 | −9.1 | −16.3 | −13.3 | −12.6 | −16.4 |
|       R:fUA20 | −14.3 | −14.5 | −15.2 | −15.3 | −18.8 | −14.6 | −13.9 | −15.2 | −15.3 |
| C1′···C1′ (Å) fUA:R | 10.4 | 10.5 | 10.4 | 10.4 | 11.0 | 11.2 | 11.0 | 10.5(WC) | 10.5 |
|       R:fUA20 | 10.5 | 10.6 | 10.3 | 10.4 | 11.1 | 10.6 | 11.1 | 11.1(rw) | 10.5 |

R: a purine residue, A or G, Ave: average, Unm: unmodified duplex [14], WC: Watson-Crick type and rw: reversed wobble.

Figure 2: The overall structures of DNA duplexes containing fU, (a) fUA1, (b) fUA2, (c) fUA3, (d) fUA4, (e) fUG1, (f) fUG2, and (g) fUG3. The fU residues are colored red. Hoechst33258 and DAPI are omitted for clarity.
from the electron densities, were assumed in the structure refinements. (The residue numbering is 1 ~ 12 for one chain and 13 ~ 24 for another chain.)

On the other hand, fUG was difficult to crystallize. Attempts to crystallize fUG under conditions similar to those for fUA, that is, at neutral or slightly acidic pH, were unsuccessful. Small single crystals of fUG2 appeared at basic pH (pH 8.1) but they poorly diffracted X-rays. Cocrystallization with several dyes was then attempted with the hope of stabilizing duplex formation. This approach led to the successful crystallization of fUG3 and fUG5 although the resolution of the fUG3 crystal was still quite low. Hoechst 33258 and DAPI were found to be bound in the central region of the minor grooves of fUG2 and fUG3, respectively, in a manner similar to the other DNA duplexes containing such dyes [21–27]. Interestingly, during refinements of fUG2, it was found that the two fU bases moved to different directions, one toward the minor groove side and the other toward the major groove side from the canonical Watson-Crick-type pairing position, and the resulting electron density also showed the same movements. In the case of fUG3, however, the two fU bases moved to the same direction toward the minor groove side, and the electron densities, though poor, also supported the movements.

3.2. Overall Structures. The local helical parameters show that all the fUA and the fUG duplexes adopt the B-form conformation even in complex with dyes, as shown in Figure 2. Superimpositions of the present structures onto the unmodified duplex structure are shown in Figure 3 and yield an average root mean square deviation of 1.4 Å.

Closener inspection of the superimposed structures reveals no drastic differences between the modified and the unmodified duplexes. However, the minor groove widths (see Figure 4) indicate that fUG2 and fUG3 are wider at the center compared with those in fUA1, fUA2, fUA3, fUA4, and fUG1 because Hoechst33258 and DAPI are bound in the grooves.

3.3. Hydrogen Bonding Schemes of Base Pairs. In every crystal, the two DNA strands are associated to each other through base-pair formations, and all the bases, except for the modified bases and their counter bases positioned on the opposite strands in the duplexes, form the canonical Watson-Crick base pairs. All the pairing geometries of fU residues are shown in Figure 5. As depicted in Figure 1(a), the electron densities clearly show that the fU residues in fUA2 are paired with the opposite A residues in the Watson-Crick geometry. Those of other fUA crystals also show the same paired structures. These results indicate that the oxidized T residue (fU) still has an ability to form a Watson-Crick-type base pair.
Figure 5: The pair formation geometry of fU residues found in (a) fUA1, (b) fUA2, (c) fUA3, (d) fUA4, (e) fUG1, (f) fUG2, and (g) fUG3. Broken and dotted lines indicate possible hydrogen bonds and CH···O interactions, respectively, and values indicate atomic distances in Å. The pictures are drawn by the program RASMOL [28].
Figure 6: Hydrogen bonding schemes with chemical structures of the observed base pairs: (a) Watson-Crick type, (b) wobble type, and (c) reversed wobble type. M indicates a hydronium or sodium ion.

Figure 7: Models of the Watson-Crick-type fU:A (gray) and the reversed wobble fU:G (blue) pairs bound in DNA polymerase I (PDB-ID 3EZ5, see [30]), constructed by fitting the paired bases to the observed pair. The template residues and incoming NTP molecules are bound in the binding pocket of the enzyme, the hydrophilic residues of which may interact to the atoms at the edges of the two bases. The guanine N² and C² atoms of the wobble fU:G pair (red) are too close to the protein atoms.

with A. The formyl groups of fU could either adopt the anti or syn conformations depending on the surrounding water structure.

On the other hand, the most interesting features can be seen in the interaction geometries between fU and G in the fUG crystals. In the fUG² crystal, the fU20 residue forms a typical wobble pair with G5 through the N³H...O⁶ and O²...HN¹ hydrogen bonds, and the formyl group is disordered between syn and anti conformations. However, the fU8 residue forms a new type of pair with the opposite G17 residue through the two hydrogen bonds, O⁴...HN¹ and N³...HN². In the pairing, the G base moves to the major groove side and the fU8 base moves to the minor groove side. Compared to the wobble pair, these bases move in the reverse direction. An atom was located on the electron density map (see Figure 1), and it is within hydrogen distance of the O⁴ and O⁵ atoms of fU8 and the O⁶ atom of G17. As the density of the atoms was of almost the same level as those of other water molecules, a water molecule was temporarily positioned at the peak for further structure refinements. In other words, it seems that a water molecule is trapped in the space surrounded by the three oxygen atoms to stabilize the pair formation. In the fUG³ crystal, the two fU residues at the different sites also form the respective pairs with the opposite G residues. The geometries are, however, both in the new reversed wobble type, as shown in Figure 1(c). Water molecules are also assigned at the positions similarly surrounded by the three oxygen atoms, as described above, though the corresponding electron density at the fU20 site is rather poor due perhaps to low resolution. Furthermore, in the fUG¹ crystal obtained at pH 8.1 without the help of dyes, the electron density, though again at low resolution, suggests that the two fU residues form pairs with the opposite G residues in a similar manner to those found in fUG³ obtained with DAPI. Therefore it is considered that the presence of dyes stabilizes duplex structure but does not affect the pairing modes.

4. Discussion

Figure 6 summarizes the pairing modes with their chemical structures, found in the fUA and fUG crystals. It is noteworthy that an oxygen atom always exists at the center almost equidistant from the three surrounding oxygen atoms (O⁴ and O⁵ of fU and O⁶ of G) in the reversed wobble geometry. Because of this pairing, the N¹ atom of fU should be deprotonated and the O⁴ atom of fU might be ionized so that the central pocket must accept a hydronium ion instead of a water molecule to stabilize the pair formation. This is consistent with the fact that the fUG¹ crystal was obtained in alkaline state without the help of dyes. The pKₐ value decreases from 9.7 to 8.1 in response to the formylation by T oxidization [29]. A sodium ion but not a potassium could also be accommodated, judging from the size of the pocket.

A comparison of the local base-pair parameters at the modified pairs is given in Table 3. As the sequence is basically palindromic, the duplex has a twofold axis at the centre of the duplex perpendicular to the helical axis. Therefore,
the base pairs at fU8 and fU20 exhibit buckle angles and shear distances with signs (+/-) different between the positions related by the twofold symmetry, but propeller twist angles with the same sign though their absolute values are affected by crystal packing. All of the fU:A pairs satisfy these conditions. (The shear values fluctuate around zero in the standard B-form duplex.) The fUG duplexes also adopt this geometrical rule, except for the shear parameter. In the fUG2 duplex, the two fU:G pairs have the shear values with the same sign (fU8 = 2.24 Å and fU20 = 2.08 Å). This means that the G17 and G5 bases which are paired with fU8 and fU20 move in the different directions along the X-axis [see the definition in (19)], that is, G17 shifts up toward the major groove side and G5 shifts down toward the minor groove side (or fU8 moves down toward the minor groove and fU20 moves up toward the major groove). These movements are just ascribed to the reversed wobbling and the normal wobbling, respectively. Another feature of the fUG reversed wobbling can be seen in the C1′-C4′ distance which is longer by 0.6 Å, as compared with those of the Watson-Crick pairing. This expansion is also reflected in the wider minor groove widths found in every fU:G duplexes.

The present work has clarified a total of three interaction modes, a Watson-Crick type for fU:A pairing and two wobble types (wobble and reversed wobble) for fU:G pairing. It has been believed that the DNA polymerase accepts base pairs only with the Watson-Crick geometries. The wobble pairing easily occurs between G and T, but the DNA polymerase only with the Watson-Crick geometries. The wobble pairing can also be accepted by the enzyme the pairing geometry was modeled into the enzyme in silico. As seen in Figure 7, the reversed wobble pair is reasonably accommodated without atomic collisions by slightly rotating the pair. On the contrary, a wobble pair, rotated in the opposite direction, would still collide with atoms of the enzyme. Thus, it could be concluded that G forms a pair with fU in the ionized form and A forms a Watson-Crick pair with fU and that the reversed wobble pair is allowed to be bound in the DNA polymerase. This tolerance explains the occurrence of mis-incorporation during replication and leads to pyrimidine transition mutagenesis.

Methoxyamine is also well known as a potent mutagen. N6-methoxyl-adenine [31, 32] and N4-methoxyl-cytosine [33-35] residues in the damagedDNAs adopt the imino tautomer to form mismatched base pairs, which mimic the Watson-Crick pair geometries. In contrast, fU derived from thymine oxidation forms a reversed wobble pair, which is also acceptable to the DNA polymerase.

The focus of the present work is on pyrimidine transition mutation and not gene transversion. The atomic mechanism of the latter will be revealed by similar X-ray studies.

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