A Mutant of Uracil DNA Glycosylase That Distinguishes between Cytosine and 5-Methylcytosine

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Abstract

We demonstrate that a mutant of uracil DNA glycosylase (N123D:L191A) distinguishes between cytosine and methylcytosine. Uracil DNA glycosylase (UDG) efficiently removes uracil from DNA in a reaction in which the base is flipped into the enzyme’s active site. Uric was selected over cytosine by a pattern of specific hydrogen bonds, and thymin is excluded by steric clash of its 5-methyl group with Y66. The N123D mutation generates an enzyme that excises cytosine. This N123D:L191A mutant excises C when it is mispaired with A or opposite an abasic site, but not when it is paired with G. In contrast no cleavage is observed with any substrates that contain 5-methylcytosine. This enzyme may offer a new approach for discriminating between cytosine and 5-methylcytosine.

Introduction

Uracil, which is generated by deamination of cytosine [1] producing G.U mispairs, is removed from DNA by uracil-DNA glycosylase (UDG) [2–4]. UDG is highly specific for uracil and shows no activity towards any other base [5]; the base pair partner of the U is not recognised and the enzyme also acts on AU base pairs that arise through misincorporation during DNA replication [6]. Uric is flipped out of the duplex into the enzyme’s active site, followed by cleavage of the N-glycosidic bond [7–9]. This base flipping is aided by L191 that inserts into the DNA duplex [8], pushing out the uracil and increasing its lifetime in the active site [10]. The L191A mutant is less efficient at flipping out uracil [11], though the enzymatic activity can be rescued by pairing uracil with a bulky synthetic nucleoside that occupies the space of the base pair [10,12]. Thymine is excluded from the active site of UDG by steric clash between its 5-methyl group and Y66 [9,13].

UDG’s remarkable specificity for uracil results from specific hydrogen bonding [14,15] and shape complementarity [11,12,16,17]. In particular N123 forms specific hydrogen bonds with O4 and N3 of uracil. Mutation of N123 to aspartate (N123D) alters the hydrogen bond donor-acceptor pattern, allowing for recognition of cytosine thereby generating a cytosine DNA glycosylase (CDG) as shown in Figure 1A [17]. The double mutant (N123D:L191A, designated as CYDG), is unable to excise cytosine from a G.C base pair [18]. It has been reported that this enzyme only shows CDG activity when C is paired with a bulky group such as pyrene, which forces the cytosine into an extrahelical conformation [10,12,18]. This mutant still cleaves uracil at which it is at least 1000-fold less active than UDG. We reasoned that CDG should be able to discriminate between cytosine and 5-methylcytosine (5MeC) by the same mechanism that UDG discriminates between U and T (Figure 1B).

Cytosine methylation, especially at CpG sites, acts as an epigenetic marker which affects gene expression and regulation. The most commonly used methods for detecting 5-methylcytosine are direct sequencing after treatment with bisulphite [19] or protection from cleavage by methylation sensitive restriction enzymes [20,21]. We have therefore explored whether CYDG can discriminate between G and 5MeC, in the same way that UDG discriminates between U and T (Figure 1). We have determined the cleavage selectivity of CYDG and show that it can remove cytosine, but not methylcytosine, when it is mispaired or opposite an abasic site.

Materials and Methods

Preparation of Enzymes

The sequence of Escherichia coli UDG was cloned between the EcoRI and HindIII sites of pUC18. Site-directed mutagenesis generated the L191A mutation, which was followed by the N123D mutation. The sequence was then subcloned into pET28a and inserted between the NdeI and EcoRI sites. The enzyme (CYDG) was expressed in BL21(DE3)/pLysS cells, which were induced with 0.2 mM IPTG for three hours. The cells were lysed by sonication, purified using a Ni-NTA (His Trap FF Crude; GE Healthcare) and eluted in 250 mM imidazole. The enzyme was concentrated and further purified using a 20 mL 10000 MW Vivaspin column (Fisher Scientific). This produced CYDG that was about 95% pure, as estimated by SDS polyacrylamide gel electrophoresis, with a yield of 1.5 mg per litre culture.
Preparation of Oligonucleotides

Oligonucleotides were synthesized on an Applied Biosystems ABI 394 automated DNA/RNA synthesizer on the 0.2 or 1 μmol scale using standard methods. Phosphoramidite monomers and other reagents were purchased from Applied Biosystems or Link Technologies. The pyrrolidine anthraquinone phosphoramidite was purchased from Berry & Associates. Each 31 mer oligonucleotide was radiolabelled at its 5'–end with γ-32P[ATP] using T4 polynucleotide kinase (New England Biolabs), purified by denaturing PAGE, and resuspended in 10 mM MES pH 6.3 containing 25 mM NaCl and 2.5 mM MgCl2. These were mixed with an excess of the unlabelled complementary oligonucleotides and annealed by slowly cooling from 95°C to 4°C.

Enzyme Cleavage

Radiolabelled DNA (approximately 50 nM) was incubated with CYDG (typically 1.25 μM) for up to 24 h, removing samples at various time intervals. The reaction was stopped using 10% piperidine (v/v) and heated at 95°C for 20 min to cleave the phosphodiester backbone. The samples were lyophilised, resuspended in 5 μL loading buffer (80% (v/v) formamide, 10 mM EDTA, 10 mM NaOH and 0.1% (w/v) bromophenol blue) and run on a 12.5% denaturing polyacrylamide gel containing 8 M urea. The gel was then fixed, dried, subjected to phosphorimaging and analysed using ImageQuantTL. Experiments were performed in triplicate; kcat values were determined from plots of percentage cleaved against time, using SigmaPlot, by fitting each set of data to a single exponential rise to maximum. These were then averaged and the rate constants are reported with ± standard deviation. The rate of cleavage of some substrates was very low (less than 10% cleaved after 24 hours incubation). In these instances an estimate of the rate constant was obtained from the fraction cleaved at a given time, assuming a simple exponential process.

Results

Generation of CYDG (N123D, L191A)

Initial attempts to prepare the N123D mutant of E. coli UDG, which should have CDG activity, were unsuccessful, confirming that this enzyme is cytotoxic in E. coli [17,18]. The L191A mutant was therefore first introduced into UDG (generating UYDG [7]), which was followed by the second N123D mutation to produce CYDG. The mutations were generated in pUC18 and then subcloned into pET28a followed by expression of the protein in E. coli.

Excision Properties of CYDG

The activity and specificity of CYDG were tested against a range of double and single stranded DNA templates. Synthetic 31 mer oligonucleotide substrates were designed so as to pair U, T, C or 5MeC with G, A, AP (abasic site), Z (anthraquinone pyrrolidine) or a gap using two 15 mer oligonucleotides (Table 1). Previous studies have used a pyrene nucleoside [10,12,18] as a plug to force the base into the active site; we used anthraquinone pyrrolidine as a similar bulky nucleotide analogue. The results, after incubating all the substrates with an excess of the enzyme, are shown in Figure 2. Most importantly CYDG shows no activity against all the sequences that contain a central methylcytosine, confirming that the 5-methyl group of cytosine is excluded from the active site in a similar fashion to exclusion of the 5-methyl group of T by UDG. In contrast all the sequences with a central cytosine are cleaved, except when this is paired with guanine. As expected, cleavage is observed when C is place opposite the bulky anthraquinone analogue, as previously observed with a pyrene nucleotide [18]. More surprisingly, cleavage is also observed when C is placed opposite an A, an abasic site or a gap, though there is no reaction with a G.C base pair. CYDG has residual activity against uracil, even when this is positioned...
The target bases are shown in bold and underlined; where X = U or C and Q = G, A, AP (abasic site) or Z (anthraquinone pyrrolidine).

doi:10.1371/journal.pone.0095394.t001
base and the 5-methyl group is sterically excluded. Alteration of the hydrogen bonding pattern at N123 changes the base selectivity, but the mutant enzyme is still able to discriminate between pyrimidine and 5-methylpyrimidine. The lack of activity of CYDG against G.C base pairs therefore suggests the possibility of using this enzyme to probe the methylation status of a specific cytosine, by mispairing it with another base such as adenine.

CYDG cleaves cytosine when it is unpaired or mispaired, and the stability of the base pair determines the rate of cleavage [23,24]. CYDG excised cytosine from Z.C faster than uracil from A.U, presumably because the mispaired cytosine is more easily forced into an extrahelical configuration than uracil in the Watson-Crick A.U pair. The faster cleavage of gap.C and AP.C occurs because there is no base opposite the C. If G.C base pairs flank the target cytosine then the rate of cleavage at A.C is dramatically reduced as a result of the increased local DNA stability [25] and the inability of CYDG to flip the base into the active site [10–12]. CYDG retains uracil DNA glycosylase activity despite the N123D mutation since free rotation of the aspartate side chain can still present the correct hydrogen bonding pattern for interacting with U [26]. Although the activity of CYDG is greatly reduced compared with wild type UDG, its catalytic activity is similar to that of many other DNA glycosylases [27–30].

The ability of CYDG to excise uracil from A.U but not cytosine from G.C suggests that this activity is dependent on the stability of the base pair and that the base can move into the enzyme’s active site when it is not involved in a stable base pair. The major role of L191 therefore seems to be to plug the space left after base flipping, rather than to actively assist the mechanism of base flipping itself [11]. The binding of CYDG to the duplex and the distortion it causes to the DNA [11,16,31] appears to be sufficient to destabilise an A.U but not G.C base pairs.

Figure 3. CYDG cleavage of fragments containing a central U or C opposite different bases. In each gel the 32P labelled duplex substrates (~50 nM) were incubated with 1.25 mM CYDG for up to 24 hours and cleaved by boiling in 10% piperidine. The products were resolved on 12.5% denaturing polyacrylamide gels. The graphs are derived from phosphorimage analysis of the gels and show the rate of formation of the cleavage product. These are fitted with single exponential curves.

doi:10.1371/journal.pone.0095394.g003
In summary we have shown that CYDG is able to discriminate between cytosine and 5-methylcytosine. Cytosine-DNA glycosylase activity is observed when C is unpaired or in an unstable (non-Watson-Crick) base pair, while no activity is observed at MeC in any base pair combination. This enzyme may offer an approach for discriminating between cytosine and 5-methylcytosine, in

![Figure 4. Kinetics of CYDG cleavage of fragments containing a central C.](image)

In each gel the $^{32}$P labelled duplex substrates (~50 nM) were incubated with 1.25 μM CYDG for up to 24 hours and cleaved by boiling in 10% piperidine. The products were resolved on 12.5% denaturing polyacrylamide gels. The graph for long gap.C was derived from phosphorimage analysis of the gel and shows the rate of formation of the cleavage product; this is fitted with a single exponential curve.

doi:10.1371/journal.pone.0095394.g004

| Table 2. $k_{cat}$ values for CYDG cleavage of the different DNA substrates. |

| Substrate       | $k_{cat}$ (min$^{-1}$) | Rel |
|-----------------|------------------------|-----|
| A.C             | 0.006±0.001            | 1.7 |
| A.C(G)$^1$      | 0.0001                 | ~0.02 |
| AP.C            | 0.014±0.003            | 4.0 |
| Z.C             | 0.10±0.02              | 29  |
| G.C             | ND                     | <0.001 |
| gap.C$^2$       | 0.016±0.002            | 4.6 |
| Long gap.C      | 0.0072±0.0007          | 2.0 |
| G.U             | 0.36±0.04              | 100 |
| A.U             | 0.020±0.004            | 5.6 |
| ssC(polyA)$^3$  | 0.0003±0.0001          | ~0.07 |
| ssC(GAT)$^1$    | 0.0001                 | ~0.02 |

The sequences of the oligonucleotides are shown in Table 1. No cleavage was observed for any substrate containing methylcytosine. ND - no cleavage detected after 24 hours. Values represent the average of three independent determinations ± standard deviations.

$^1k_{cat}$ values were estimated from single time points at 24 hrs A.C(G), 60 mins ssC(polyA) and 4 hrs ssC(GAT).

$^2$Only 50% of the substrate was cleaved for gap.C. Rel indicates the cleavage rate relative to that of G.U (100).

doi:10.1371/journal.pone.0095394.t002
which the methylation status of a specific cytosine is probed by annealing it with an oligonucleotide that generates a mismatch, such as AC. A cytosine at this position will be cleaved while methylcytosine will not; PCR amplification of the reaction products can then be used to discriminate between the cleaved and uncleaved species.

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Author Contributions

Conceived and designed the experiments: STK TB KRF. Performed the experiments: STK. Analyzed the data: STK KRF. Contributed reagents/materials/analysis tools: STK KRF TB. Wrote the paper: STK TB KRF.