DR0022 from *Deinococcus radiodurans* is an acid uracil-DNA glycosylase

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Uracil-DNA glycosylase (UDG) initiates base excision repair (BER) by removing damaged or modified nucleobases during DNA repair or mammalian demethylation. The UDG superfamily consists of at least six families with a variety of catalytic specificities and functions. *Deinococcus radiodurans*, an extreme radiation resistant bacterium, contains multiple members of UDG enzymes within its genome. The present study reveals that the putative protein, DR0022, is a uracil-DNA glycosylase that requires acidic conditions for its glycosylase activity, which is the first case of such an enzyme within the UDG superfamily. The key residues in the catalytic motifs are investigated by biochemical, enzyme kinetics, and *de novo* structural prediction, as well as molecular modeling analyses. The structural and catalytic roles of several distinct residues are discussed in light of predicted and modeled DR0022 glycosylase structures. The spontaneous mutation rate analysis performed in a *dr0022* deficient *D. radiodurans* strain indicated that the *dr0022* gene plays a role in mutation prevention. Furthermore, survival rate analysis in a *dr0022* deficient *D. radiodurans* strain demonstrated its role in stress resistance, including γ-irradiation. Additionally, the novel acid UDG activity in relationship to its *in vivo* roles is discussed. This work underscores the functional diversity in the UDG superfamily.

**Introduction**

In 1974, Uracil-DNA glycosylase (UDG) was discovered in *Escherichia coli* to be involved in removal of uracil, a deamination product generated from hydrolytic deamination and nitrosative stress-induced oxidative deamination of cytosine in DNA [1–3]. Over decades of investigation, at least six families of glycosylase enzymes have been found in the UDG superfamily. Although the prototypical *E. coli* enzyme is classified as a narrow specificity family 1 uracil N-glycosylase (UNG), UDGs in other families exhibit a variety of substrate specificities, from family 2 MUG/TDG, family 3 SMUG1, family 5 UDGb as broad specificity enzymes, to family 4 UDGa as narrow specificity enzymes, and to family 6 HDG as hypoxanthine DNA glycosylase [4–11]. Within each UDG family, the specificity may also vary depending on the source of organisms [12–19]. A notable expansion of specificity with profound biological significance occurred in family 2 MUG/TDG, in which the human TDG becomes a DNA glycosylase involved in DNA demethylation by removing 5-formylcytosine and 5-carboxycytosine derived from oxidation of 5-methylcytosine by ten-eleven translocation methylcytosine dioxygenase [15,16]. A rather unusual class of UDG enzymes were found originally in *Mycobacterium* that cross links with the DNA after the excision of uracil [20–23]. These results indicate that enzymes in the UDG superfamily are adaptable to acquire new

**Abbreviations**
BER, base excision repair; Dra, *Deinococcus radiodurans*; PDB, Protein Data Bank; Tth, *Thermus thermophilus*; UNG, uracil N-glycosylase; UDG, uracil-DNA glycosylase.
enzymatic functions. In addition to enzymes in the UDG superfamily, endonuclease V and some enzymes in the endonuclease III superfamily are capable of acting on uracil in DNA [24–27].

*Deinococcus radiodurans* (Dra) is an extremely radiation-resistant bacterium initially found in canned meat [28]. The sequenced genome reveals the existence of a multitude of DNA repair enzymes. Four genes for enzymes (DR0689, DR1751, DR0022, DR0715) were found in the sequenced genome as putative UDG enzymes. Previous biochemical investigation determined that DR0689 and DR1751 were authentic family 1 UNG and authentic family 4 UDGa enzymes, respectively, with uracil-DNA glycosylase activity on both double-stranded and single-stranded DNA [29]. DR0715, on the other hand, only showed weak activity on G/U base pair and moderate activity on ethenocytosine-containing DNA [29]. Yet, the UDG activity from DR0715 was later confirmed by a structural and biochemical study [30]. Previous investigation on DR0022 did not find any detectable glycosylase activity, regardless of whether the protein was expressed in *E. coli* or by *in vitro* transcription–translation system [29]. In the present study, we report positive detection of uracil-DNA glycosylase activity in DR0022 and the investigation of its role in mutation repair and stress resistance, and also discuss its potential catalytic mechanism.

**Results**

**Screen for activity and biochemical analyses**

Initially, we tested the DNA glycosylase activity using purified DR0022 protein under our routine assay conditions at pH 7.5 without salt using U, I (hypoxanthine), X (xanthine), and O (oxanine) as substrates (Fig. 1A–D). Consistent with the previous study [29], we failed to detect any glycosylase activity under these assay conditions. We then surmised that this unique UDG may exert its glycosylase activity under different assay conditions. To test whether the lack of activity was a result of the pH, we measured the glycosylase activity in buffers with pH ranging from 4–8 using the U-containing substrate. Unexpectedly, we were able to detect UDG activity around pH 5.5 (Fig. 1D,E). No UDG activity was found at neutral or alkaline pH conditions (Fig. 1D,E). The glycosylase activity was found with three mismatched uracil base pairs in the order G/U > C/U > T/U. No activity was found with the A/U base pair or the single-stranded uracil-containing substrate (Fig. 1D,E). Based on this novel finding, we went back to measure the glycosylase activity on I, X, O, 5-dihydrouracil, 8-oxoG, thymine glycol, AP site, and 5-hydroxyuracil and 5-hydroxycytosine substrates at pH 5.5. We did not find any activity with these substrates (data not shown). However, we found glycosylase activity on hydroxymethyluracil (hmU) in three mismatched substrates in the order G/hmU > C/hmU > T/hmU. Again, no activity was found with the A/hmU base pair or the single-stranded hmU substrate (Fig. 1F). The acidic glycosylase activity was optimal at 0 mM salt and at 37 °C. To define the kinetic properties of the glycosylase activity, we performed time course analyses using the kinetics methodology used previously [10,31,32]. The *K*ₘ values were approximately 1000 nM for the G/U, G/hmU, C/hmU, and T/hmU base pairs, but at lower values for the C/U and T/U base pairs (Table 1). The *k*_2 value was 0.166 min⁻¹ with G/U, which was 3- to 18-fold higher than for other substrates (Table 1). Overall, the catalytic efficiency as defined by *k*_2/*K*ₘ was higher with uracil-containing substrates than that with hmU-containing substrates (Table 1).

In addition to DR0022, we found putative genes from other bacterial species that have similar motif architecture and grouped them together as DR0022-like family (Fig. 2). Among the different families in UDG superfamily, DR0022 is most similar to family 4 UDGa in the conservation of catalytic residues in motifs 1, 2, and 3 (Fig. 2). A notable difference is that motif 1 in DR0022 starts with LEAP rather than a leucine rather than a glycine. On the other hand, motifs 2 and 3 are well conserved with an initiating Asn and His, respectively (Fig. 2). To probe the functional role of these motifs in the catalysis of the acid uracil-DNA glycosylase, we constructed five single-point mutants in five positions (Fig. 2). As shown in Table 2, no glycosylase activity was detected with L43G, E44A in motif 1, N90A in motif 3, and H159A in motif 2. Interestingly, A50E mutant lost glycosylase activity on hmU but exhibited an elevated UDG activity. The higher level of UDG activity on G/U base pair was further investigated by kinetics analysis, which showed approximately 2-fold reduction of *K*ₘ and doubling of *k*_2, resulting in an almost 5-fold increase in *k*_2/*K*ₘ (Table 1). Subsequently, we introduced L43G into the A50E mutant and found that the double mutant had a detectable but lower level of UDG activity compared to A50E mutant and the wild-type enzyme (Table 2). The implication of these results in
relationship to the catalytic mechanism is discussed in detail further below.

**In vivo analysis**

To understand the role of DR0022 mutation prevention *in vivo*, we deleted the *dr0022* gene from the Dra genome using double crossover recombination of a kanamycin resistance cassette into the genome [33]. The deletion of *dr0022* was confirmed by PCR analysis. We first examined the effect of Dra (*dr0022<sup>−</sup>*) on spontaneous mutation by measuring the mutation rates for the rifampicin resistance. In the presence of the *dr0022* gene, the mutation rate was $4.67 \times 10^{-9}$ in Dra (Fig. 3). Once the *dr0022* gene was deleted, the mutation rate jumped to $24.67 \times 10^{-9}$, which is more than a 5-fold increase over the wild-type strain. We then investigated the nature of base pair changes in the *rpoB* gene in Dra (*dr0022<sup>−</sup>*). As shown in Table 3, the majority of the base pair changes was A/T to T/A, followed by G/C to A/T, then by A/T to C/G. The A/T transversion (A/T to T/A, A/T to C/G) represented 82% of the substitutions.

**Response to stresses**

To understand the role of *dr0022* under various stress conditions, we measured survival rates of "dr0022"...
Table 1. Kinetic constants of wild-type and A50E DR0022 glycosylase on U- and hmU-containing DNA substrates. Each kinetic constant was determined as described in the Materials and methods. Data are the average of three independent experiments. NA, no activity detected under the assay conditions.

| Protein | Substrates | $k_{m}$ (nM) | $k_{2}$ (min$^{-1}$) | $k_{2}/k_{m}$ (min$^{-1}$ nM$^{-1}$) |
|---------|------------|--------------|----------------------|----------------------------------|
| WT      | G/U        | 1257 ± 143   | 0.166 ± 0.007        | 1.32 × 10$^{-4}$               |
|         | C/U        | 320 ± 41     | 0.052 ± 0.002        | 1.63 × 10$^{-4}$               |
|         | T/U        | 396 ± 24     | 0.039 ± 0.006        | 9.85 × 10$^{-5}$               |
|         | G/hmU      | 1080 ± 196   | 0.021 ± 0.001        | 1.94 × 10$^{-4}$               |
|         | C/hmU      | 1640 ± 419   | 0.014 ± 0.002        | 8.53 × 10$^{-4}$               |
|         | T/hmU      | 1034 ± 152   | 0.009 ± 0.001        | 8.70 × 10$^{-5}$               |
| A50E    | G/U        | 667 ± 122    | 0.41 ± 0.02          | 6.15 × 10$^{-4}$               |
|         | G/hmU      | NA           | NA                   | NA                              |

Dra strain and the wild-type strain were readily reduced at a NaNO$_2$ concentration of 10 mM (Fig. 4A). At 30 mM, both the dr0022 deficient and the wild-type strains completely lost survival. Notably, the survival rates of the wild-type strain were statistically lower than the dr0022 deficient strain at concentrations of 10 and 20 mM (Fig. 4A). The decrease of survival rates was much slower when treated with H$_2$O$_2$, with a sharp drop at 10 mM and a slow decline up to 80 mM (Fig. 4B). The lower level of survival in the wild-type strain may suggest active removal of nucleobase lesions, which may introduce more AP sites and/or strand breaks and lead to increase lethality. The survival curves of both strains with UV-C treatment were almost identical, with a 50% reduction at 10 min and an almost 100% reduction at 20 min (Fig. 4C). When treated with γ-irradiation, the survival rates of both strains experienced similar levels of reduction up to 3 kGy (Fig. 4D). The responses of both strains then

Fig. 2. Sequence alignment of DR0022-like enzymes with other UDG families. The sequence alignment was performed by CLUSTAL OMEGA (http://www.clustal.org) and adjusted manually. The GenBank accession number is provided after the species name. The sites in DR0022 that are subject to site-directed mutagenesis analysis are marked by an arrow. 0022-like: Dra (DR0022)

Table 2. Kinetic constants of wild-type and A50E DR0022 glycosylase on U- and hmU-containing DNA substrates. Each kinetic constant was determined as described in the Materials and methods. Data are the average of three independent experiments. NA, no activity detected under the assay conditions.

| Protein | Substrates | $k_{m}$ (nM) | $k_{2}$ (min$^{-1}$) | $k_{2}/k_{m}$ (min$^{-1}$ nM$^{-1}$) |
|---------|------------|--------------|----------------------|----------------------------------|
| WT      | G/U        | 1257 ± 143   | 0.166 ± 0.007        | 1.32 × 10$^{-4}$               |
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|         | T/U        | 396 ± 24     | 0.039 ± 0.006        | 9.85 × 10$^{-5}$               |
|         | G/hmU      | 1080 ± 196   | 0.021 ± 0.001        | 1.94 × 10$^{-4}$               |
|         | C/hmU      | 1640 ± 419   | 0.014 ± 0.002        | 8.53 × 10$^{-4}$               |
|         | T/hmU      | 1034 ± 152   | 0.009 ± 0.001        | 8.70 × 10$^{-5}$               |
| A50E    | G/U        | 667 ± 122    | 0.41 ± 0.02          | 6.15 × 10$^{-4}$               |
|         | G/hmU      | NA           | NA                   | NA                              |
started to diverge at higher $\gamma$-irradiation dosages, with the survival rate of the $dr0022$ deficient strain dropping to around 10%, whereas the wild-type strain reduced to around 20% at levels of 5 and 7 kGy, indicating that $dr0022$ plays a role in radiation resistance (Fig. 4D).

**Discussion**

Dra, as an extremely radiation resistant and stress resistant microorganism, has an expanded DNA repair repertoire to process a wide variety of DNA lesions [34]. Previous studies have identified several uracil-DNA glycosylases and confirmed enzymatic activities in several of them [29]. The present study identifies DR0022 as an acid uracil- and hydroxymethyluracil-DNA glycosylase. To our knowledge, this is the first report of such a UDG enzyme in UDG superfamily.

To understand the structural differences between DR0022 and other UDG enzymes, we took advantage of a state-of-the-art protein structure prediction method, ALPHAFOLD [35]. ALPHAFOLD is developed based on deep learning algorithm that demonstrates highly accurate domain structure and amino acid side chain predictions. To validate the accuracy of ALPHAFOLD, we predicted the structures of the Tth family 4 UDGa and the Tth family 5 UDGb using ALPHAFOLD at the Colab online server [36]. Superimposition demonstrated that the predicted structure was highly similar to the corresponding solved crystal structures, with a rmsd of 0.289 Å [Protein Data Bank (PDB): 1UI0] and 0.375 Å (PDB: 2DP6), respectively (Fig. 5). We then used ALPHAFOLD to predict the three-dimensional structure of DR0022 protein. The predicted DR0022 protein structure showed a typical structural fold, as seen in other UDG protein structures solved by X-ray crystallography with a predicted local distance difference test score above 90 (Fig. 6A), an indication of high confidence in the structural prediction. We then used SWISS-MODEL (https://swissmodel.expasy.org) to build a modeled structure based on known crystal structures. Analysis of sequence identity in SWISS-MODEL indicated that DR0022 had a relatively higher sequence identity with the Tth family 5 UDGb (PDB: 2DP6, 25% identity) and the Tth family 4 UDGa (PDB: 1UI0, 20% identity). We therefore built the DR0022 structure using Tth family 5 UDGb (PDB: 2DP6) and Tth family 4 UDGa (PDB: 1UI0) (Fig. 6B,C). The predicted DR0022 structure had a very similar structure to the modeled structures using 2DP6 and 1UI0 structures as the

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**Table 2.** DNA glycosylase activity of DR0022 on U- and hmU-containing DNA. Experiments were performed as described as Materials and methods. Data are the average of three independent experiments.

|        | A | U | T | G | C | A | U | T | G | C |
|--------|---|---|---|---|---|---|---|---|---|---|
| Wild-type | 0 | 12 | 34 | 18 | 0 | 0 | 12 | 23 | 15 | 0 |
| L43G    | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 |
| E44A    | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 |
| A50E    | 0 | 28 | 46 | 26 | 0 | 0 | 0 | 0 | 0 | 0 |
| N90A    | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 |
| H159A   | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 |
| L43G-A50E | 0 | 6 | 16 | 11 | 0 | 0 | 0 | 0 | 0 | 0 |

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**Table 3.** Distribution of mutations leading to Rif in $dr0022$ deficient *Deinococcus radiodurans* strain.

| Mutation site in rpoB (bp) | Amino acid change | Base pair change | Number |
|---------------------------|-------------------|-----------------|--------|
| 1274                      | D425V             | AT-TA           | 39     |
| 1319                      | S440F             | GC-AT           | 9      |
| 1304                      | H435P             | AT-CG           | 2      |

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![Fig. 3. Rif resistance mutation rates from the wild-type and $dr0022$ deficient *Deinococcus radiodurans* strains.](image-url)
template, with rmsd values of 1.517 and 1.115 Å, respectively. As a computational method to predict three-dimensional structures, ALPHAFOLD-based de novo structural prediction has been accessed and evaluated independently as powerful and accurate [37]. As a test case here, the predicted DR0022 structure appears to be highly consistent with the modeled structures.

Despite the similarity in motifs 2 and 3, the most obvious difference is that the first residue in motif 1 of DR0022 is a leucine rather than a glycine residue, which is universally seen in other families of UDG superfamily (Fig. 7). Interestingly, the substitution of Leu with Gly (L43G) rendered DR0022 inactive under both acidic and alkaline pH conditions, suggesting that the enzyme has adapted to the hydrophobic Leu residue in this position. The predicted and modeled DR0022 structures suggest that L43 may stack on the uracil ring to enhance or facilitate other interactions with uracil in the base binding pocket (Fig. 7). Interestingly, the acid UDG activity can be partially rescued by the L43G-A50E double mutant (Table 2), suggesting the correlated nature of certain amino acid changes during evolution.

The base recognition pocket and the catalytic center bear similarities to other UDG enzymes (Fig. 8). In the predicted and modeled DR0022 structures, the backbone of E44 of motif 1 interacts with O2 of uracil as seen in family 4 UDGa [10]. H159 of motif 2 interacts with O2 of uracil to stabilize the departing uracil as seen in Tth UDGa [10]. N90 of motif 3 may engage in extensive interactions with the N3 and O4 moieties of uracil. These important roles in substrate recognition and catalysis are demonstrated by the lack of glycosylase activities in amino acid substitutions in these positions (Tables 1 and 2). These results indicate that DR0022 uses the same general catalytic mechanism as seen in family 4 UDGa to remove uracil or hmU in DNA [10]. An interesting observation is that the substitution of alanine with glutamate at A50 position (A50E) actually increased the acid UDG activity (Table 2), and kinetics analysis demonstrated that $k_2/K_m$ value was almost 5-fold higher than the
Fig. 5. Comparison of predicted Tth family 4 UDGa and family 5 UDGb structures with corresponding crystal structures. (A) ALPHAFOLD2 predicted structure of *Thermus thermophilus* family 4 UDGa protein. (B) Crystal structure of *T. thermophilus* family 4 UDGa (PDB: 1UI0). (C) Superimposition of predicted family 4 UDGa structure with corresponding crystal structure. (D) ALPHAFOLD2 predicted structure of *T. thermophilus* family 5 UDGb protein. (E) Crystal structure of *T. thermophilus* family 5 UDGb (PDB: 2DP6). (F) Superimposition of predicted family 5 UDGbs structure with corresponding crystal structure.

Fig. 6. Modeled and predicted DR0022 protein structures. (A) Structure of DR0022 protein constructed by *de novo* prediction using ALPHAFOLD2. (B) Structure of DR0022 protein modeled after *Thermus thermophilus* family 5 UDGb crystal structure (PDB: 2DP6) using SWISS-MODEL. (C) Structure of DR0022 protein modeled after *T. thermophilus* family 4 UDGa crystal structure (PDB: 1UI0) using SWISS-MODEL.
wild-type enzyme (Table 1). Previously, we compared the structural difference between the Tth UDGa and the *E. coli* family 1 UNG and found that E47 in Tth UDGa can block the entry of thymine into the recognition pocket [10]. The predicted and modeled DR0022 structures indicate that A50E can play the same role in DR0022, which explains the elimination of hmU activity in this mutant. As shown in Fig. 9, the Tth family 5 UDGb has a glycine (G64) and the Tth family 4 UDGa has a glutamate (E47) in the corresponding positions. In comparison with the wild-type structures, the A50E mutant adopts a conformation that is similar to family 4 UDGa. Accordingly, it is likely that A50E may now favorably interact with the uracil base, thus enhancing the UDG activity.

The absolute requirement for an acidic environment allowing DR0022 to act as a UDG is unprecedented. We considered two scenarios to explain this phenomenon. The first scenario relates to the substrate. Uracil is a lactam with two intracyclic amide bonds, which are subject to tautomerization. The lactam form of uracil predominates under normal neutral pH [38]. However, the lactim form of uracil increases under acidic pH conditions. In the Tth UDGa structure, N80, as the first residue in motif 3 forms a bidentate hydrogen bond with N3 and O4 of uracil in the lactam form [10,39]. It is interesting to note that N90 of motif 3 in DR0022 has different conformations. In the predicted structure of DR0022 protein, N90 adopts a conformation that facilitates the formation of a bidentate hydrogen bond with N3 and O4 of uracil in the lactim form (Fig. 8C,F). Our previous study already underscores the important role of K68N of *E. coli* MUG in substrate recognition and catalysis [5]. Interestingly, the sequencing analysis of RifR in *dr0022* deficient Dra strain revealed that the A/T to T/A transversion was the predominant amino acid substitution (Table 3). This led us to speculate that DR0022 may recognize a

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**Fig. 7.** Close-up view of L43 in predicted and modeled DR0022 structures. Uracil is shown as licorice. (A) Crystal structure of *Thermus thermophilus* family 5 UDGb (PDB: 2DP6). G57 is shown as licorice. (B) Crystal structure of *T. thermophilus* family 4 UDGa (PDB: 1UI0). G40 is shown as licorice. (C) Predicted structure of DR0022 protein using ALPHAFOLD2. L43 is shown as licorice. (D) Modeled structure of DR0022 protein with 2DP6 as the template using SWISS-MODEL. L43 is shown as licorice. (E) Modeled structure of DR0022 protein with 1UI0 as the template using SWISS-MODEL. L43 is shown as licorice.
A thymine derivative (T*) that bears some structural similarity to the uracil in the lactim form (Fig. 8F). The thymine derivative may mimic an adenine base to pair with T during DNA replication. As a result, a A/T base pair is converted to a T/A base pair. In the second scenario, we assume the uracil still maintains its lactam form. However, the interaction with the uracil is not possible until one of the key catalytic residues in the active site is protonated at acidic pH conditions.

In addition to the extreme resistance to radiation, Dra is reported to be highly resistant to DNA damaging agents [40]. Most notably, the previous study found that Dra was 62-fold more resistant to nitrous acid treatment, 55-fold more resistant to radiation treatment, and 33-fold more resistant to UV-C treatment than E. coli [40]. We determined the survival rates of dr0022 deficient Dra strain aiming to understand the contribution of dr0022 to resistance to various stresses (Fig. 4). Among the four stress conditions tested, DR0022 appears to contribute to acidified nitrous acid treatment and γ-irradiation (Fig. 4). Acidified nitrous acid can generate nitrating agent, which can deaminate nucleobases. As a uracil-DNA glycosylase, DR0022 would be expected to remove the cytosine deamination product, uracil, during the repair process. Continuous removal of uracil will generate large number of AP sites, which may cause cytotoxicity. As such, the dr0022 strain appeared more resistant to acidified nitrous acid treatment than the wild-type strain.

A previous study found that the expression of dr0022 was increased by over 3-fold when Dra was treated with a high dosage of γ-irradiation [41]. In the survival assay, we found a statistically significant

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**Fig. 8.** Base recognition pocket and catalytic center in DR0022 according to predicted and modeled structures. Uracil is shown as licorice. (A) Crystal structure of *Thermus thermophilus* family 5 UDGb (PDB: 2DP6). L58 and H190 are shown as licorice. (B) Crystal structure of *T. thermophilus* family 4 UDGa (PDB: 1UI0). E41, N80, and H155 are shown as licorice. (C) Predicted structure of DR0022 protein using ALPHAFOLD2. E44, N90 and H159 are shown as licorice. (D) Modeled structure of DR0022 protein with 2DP6 as the template using SWISS-MODEL. E44, N90, and H159 are shown as licorice. (E) Modeled structure of DR0022 protein with 1UI0 as the template using SWISS-MODEL. E44, N90, and H159 are shown as licorice. (F) Lactam-lactim shift of uracil base.
difference at a high dose of γ-irradiation between the wild-type strain and the dr0022 deficient strain (Fig. 4 D). These results suggest that DR0022 can play a role in repairing DNA damage after radiation. In conjunction with the analysis above, we speculate that DR0022 may recognize a base lesion generated after a high dosage of irradiation that bears some structural similarity to uracil in its lactim form. By removing this cytotoxic lesion, DR0022 enhances the survival of Dra after γ-irradiation.

In summary, the present study reports the finding of a new acid uracil-DNA glycosylase in Dra. The biochemical and mutational analyses define the base lesion pocket and suggest a catalytic mechanism. The in vitro and in vivo investigation on mutation prevention raises the prospect that the acidic environment required for the glycosylase activity is related to its role in removing a base lesion that could generate a transversion mutation. The involvement of this glycosylase in radiation resistance is intriguing. More studies are needed to further decipher the fascinating evolution of the UDG superfamily to fulfill a variety of biological functions.

Materials and methods

Reagents, media, and strains

All routine chemical reagents were purchased from Sigma Chemicals (St Louis, MO, USA), Fisher Scientific (Suwanee, GA, USA), or VWR (Suwanee, GA, USA). Restriction enzymes, Phusion DNA polymerase, and T4 DNA

Fig. 9. Close-up view of A50 and A50E in predicted and modeled DR0022 structures. Uracil is shown as licorice. (A) Crystal structure of Thermus thermophilus family 5 UDGb (PDB: 2DP6). G64 is shown as licorice. (B) Crystal structure of T. thermophilus family 4 UDGa (PDB: 1UI0). E47 is shown as licorice. (C) Predicted structure of DR0022 protein using ALPHAFOLD2. A50 is shown as licorice. (D) Modeled structure of DR0022 protein with 2DP6 as the template using SWISS-MODEL. A50 is shown as licorice. (E) Modeled structure of DR0022 protein with 1UI0 as the template using SWISS-MODEL. A50 is shown as licorice. (F) Predicted structure of DR0022(A50E) protein using ALPHAFOLD2. A50E is shown as licorice. (G) Modeled structure of DR0022(A50E) protein with 2DP6 as the template using SWISS-MODEL. A50E is shown as licorice. (H) Modeled structure of DR0022(A50E) protein with 1UI0 as the template using SWISS-MODEL. A50E is shown as licorice.
ligase were purchased from New England Biolabs (Beverly, MA, USA). BSA and dNTPs were purchased from Promega (Madison, WI, USA). The gel DNA recovery kit was purchased from Zymo Research (Irvine, CA, USA). Oligodeoxynucleotides were ordered from Integrated DNA Technologies Inc. (Coralville, IA, USA) and Eurofins Genomics (Huntsville, AL, USA). The LB (Miller) medium was prepared according to standard recipes. Hi-Di Formamide and GeneScan™ 500 LIZ™ dye Size Standard for ABI3130xl were purchased from Applied Biosystems (Waltham, MA, USA). The sonication buffer consisted of 20 mM Tris-HCl (pH 7.5), 1 mM EDTA (pH 8.0), 50 mM NaCl, 2.5 mM dithiothreitol and 0.15 mM phenylmethanesulfonyl fluoride. The TE buffer consisted of 10 mM Tris-HCl (pH 8.0) and 1 mM EDTA.

**Cloning, site-directed mutagenesis, expression, and purification of DR0022**

The dr0022 gene from Dra R1 (GenBank accession number: AE_000513.1) was amplified by PCR using the forward primer DR0022 F (5′-GCTCTAGACCATAGTGG CCGAACCACGTCCCG-3′; the NdeI site is underlined) and the reverse primer DR0022 R (5′-CCGCTCAGGGGA AGACCAACTGGGGCCG-3′; the XhoI site is underlined). The PCR reaction mixture (50 μL) consisted of 20 ng of Dra R1 genomic DNA, 500 nM forward and reverse primers, 1 × Phusion DNA polymerase buffer, 200 μM each dNTP and 0.2 unit of Phusion DNA polymerase (New England Biolabs). The PCR procedure included a predenaturation step at 98 °C for 30 s; 30 cycles of three-step amplification with each cycle consisting of denaturation at 98 °C for 15 s, annealing at 65 °C for 20 s, and extension at 72 °C for 30 s; and a final extension step at 72 °C for 3 min. The PCR product was purified by the gel DNA recovery kit (Zymo Research). The purified PCR product and plasmid pET21a were digested by NdeI and XhoI, purified by the gel DNA recovery kit, and ligated in accordance with the manufacturer’s instructions. The ligation mixture was transformed into E. coli strain HB101 competent cells by electroporation. The sequence of the dr0022 gene in the resulting plasmid (pET21a-DR0022) was confirmed by DNA sequencing.

The resulting plasmid with wild-type dr0022 was used as the template plasmid for all other DR0022 mutants. Amplification of the mutant DNA and DpnI mediated site-directed mutagenesis procedures were modified using primers carrying the desired mutations, as described previously [42]. Briefly, PCR mixtures (25 μL) contained 10 ng of pET-21a (+)−DR0022 as a template, 65 mM each primer pair, 200 μM each dNTP, 1 × Phusion PCR polymerase buffer, and 1 unit of Phusion DNA polymerase. The PCR procedure included a pre-denaturation step at 98 °C for 2 min; 25 cycles of a three-step amplification with each cycle consisting of denaturation at 98 °C for 30 s, annealing at 55 °C for 30 s and extension at 68 °C for 5 min; and a final extension step at 68 °C for 10 min. After treatment with 2 units of DpnI for 1 h at 37 °C, 5-μL PCR products were transformed into E. coli DH5α competent cells. Successful insertion and mutation in the resultant clones were confirmed by DNA sequencing. The pET21a-DR0022 wild-type and mutants were transformed into E. coli strain BH214 (mug+, mug−) by the standard protocol to express the C-terminal His-6-tagged DR0022 protein. Induction, sonication, and purification were carried out as described previously [7]. Briefly, bacterial cells containing pET21a-DR0022 wild-type or mutant plasmid from a 500-mL culture grown to late exponential phase were harvested by centrifugation at 3993 g for 10 min. The cell pellet was suspended in 7 mL of lysis buffer [20 mM Tris-HCl (pH 7.5), 1 mM EDTA (pH 8.0), 50 mM NaCl, 0.1 mM dithiothreitol, and 0.15 mM phenylmethanesulfonyl fluoride] and followed by sonication at output 5 for 3 × 1 min with a 5-min rest on ice between intervals using a Sonifier Cell Disruptor 350 (Branson, Brookfield, CT, USA). The yields from protein expression ranged around from 2–5 mg·500 mL⁻¹. The lysate was clarified by centrifugation at 21 000 g for 20 min and filtered through a 25-mm GD/X syringe filter (Whatman, Little Chalfont, UK). The supernatant was transferred into a fresh tube and loaded onto a 1-mL HiTrap chelating column (GE Healthcare, Chicago, IL, USA), followed by a 1-mL HiTrap SP column (GE Healthcare). The SDS/PAGE analysis of the purified proteins is shown in Fig. 1C. The DR0022 protein was stored in aliquots at −80 °C. Prior to use, the protein was diluted in an equal volume of 2× storage buffer (20 mM Tris–HCl (pH 8.0), 2 mM dithiothreitol, 2 mM EDTA, 400 μg·mL⁻¹ BSA, and 100% glycerol).

**DNA glycosylase activity assay**

The DNA glycosylase cleavage assays for DR0022 were performed at optimized temperature 37 °C for 60 min in a 10-μL reaction mixture containing 10 mM oligonucleotide substrate [43], an indicated amount of glycosylase, 20 mM citric acid and phosphate buffer with the indicated pH (4.0, 4.5, 5.0, 5.5, 6.0, 6.5, 7.0, 7.5, 8.0) [44], 1 mM dithiothreitol, and 1 mM EDTA. The resulting abasic sites were cleaved by incubation at 95 °C for 5 min after adding 1 μL of 1 N NaOH. The reaction mixtures (2 μL) were mixed with 7.8 μL of Hi-Di-formamide and 0.2 μL of GeneScan 500 LI Size Standard (Life Technologies, Carlsbad, CA, USA) and analyzed using a Applied Biosystems 3130xl sequencer with a fragment analysis module. Cleavage products and remaining substrates were quantified using GENEMAPPER (Applied Biosystems).

**Enzyme kinetics analysis**

The uracil-DNA glycosylase assay was performed under optimized reaction conditions at 37 °C in a 10-μL reaction
mixture containing 10 nM oligonucleotide substrate, an indicated excess amount of DR0022 enzyme, 20 mM citric acid and phosphate buffer, pH 5.5, 1 mM dithiothreitol, and 1 mM EDTA. The enzyme concentration ranged from 50 to 4000 nM. Samples were collected at time points from 10, 20, 30, 40, 50, and 60 min. The apparent rate constants for each concentration were determined by curve fitting using the integrated first-order rate:

\[ P = P_{\text{max}} \left(1 - e^{-k_{\text{obs}}t} \right) \]  

where \( P \) is the product yield, \( P_{\text{max}} \) is the maximal yield, \( t \) is time, and \( k \) is the apparent rate constant.

The kinetic parameters \( k_2 \) and \( K_m \) were obtained from plots of \( k_{\text{obs}} \) against the total enzyme concentration ([\( E_0 \)]) using a standard hyperbolic kinetic expression [10,31,32]:

\[ k_{\text{obs}} = \frac{k_2 [E_0]}{K_m + [E_0]} \]  

\[ (2) \]

**Construction of the dr0022 deficient strain**

The dr0022 deletion mutant was constructed by double crossover recombination of a kanamycin resistance cassette into the genome, with some modifications of the method as described previously [33]. Briefly, all primers were designed using the genome sequence information of Dra R1 available on NCBI (https://www.ncbi.nlm.nih.gov). The upstream fragment was amplified by PCR, with the primer set DR0022 up \( \text{XbaI} \) F (5'-GCTCTAGACCCAGAATGCC GAGGGTGGC-3'; the \( \text{XbaI} \) site is underlined) and DR0022 up \( \text{BamHI} \) R (5'-CGCCGATCCGAGCTTAG GAGGGGCGCT-3'; the \( \text{BamHI} \) site is underlined) and DR0022 up \( \text{BamHI} \) R (5'-CGCCGATCCGAGCTTAG GAGGGGCGCT-3'; the \( \text{BamHI} \) site is underlined). Similarly, the downstream fragment was PCR-amplified with the primer set DR0022 down HindIII F (5'-CCCAAGCTTC GTTACTAGG GCAATTGAC-3'; the HindIII site is underlined) and DR0022 down \( \text{XhoI} \) R (5'-CCGCTCGAGGGGCGATACCTGCCCCCGAT-3'; the \( \text{XhoI} \) site is underlined). The upstream and downstream fragments were digested with \( \text{BamHI} \) and HindIII, respectively, and ligated to the \( \text{BamHI} \)-HindIII HindIII predigested fragment of the kanamycin resistance cassette obtained from a shuttle plasmid pRADK bearing the groEL promoter and the kanamycin resistance gene [33]. The ligation product was used as a template for PCR amplification of the full-length PCR product, with the primer set DR0022 up \( \text{XbaI} \) F and DR0022 down \( \text{XhoI} \) R. The resulting PCR product was digested by \( \text{XbaI} \) and \( \text{XhoI} \) and ligated into the pET21a vector that had been digested with the same enzymes to yield a pET21a-DR0022 deletion, which was then transformed into Dra R1. Mutant strains were selected on TGY agar plates supplemented with kanamycin (25 \( \mu \)g/mL\(^{-1}\)). The dr0022 gene mutant was designated as Dra (dr0022\(^{-}\)).

**Spontaneous mutation rate**

The spontaneous mutation rates of Dra deficient mutants were tested using the rpoB/Rif\(^{+}\) system [45]. Briefly, the freshly growing Dra cells were diluted and tittered by plating on TGY plates. Spontaneous Rif\(^{+}\) mutants were obtained by inoculating a certain number (2 \( \times \) 10\(^3\)) of Dra cells on TGY plates containing 50 \( \mu \)g/mL\(^{-1}\) rifampicin (Sigma Chemicals). The Rif\(^{+}\) mutation rate (\( f \)) was determined by the number of clones with rifampicin-resistance divided by the total number of Dra cells plated on TGY plates supplemented with 50 \( \mu \)g/mL\(^{-1}\) rifampicin.

**Sequencing the rpoB gene for mutations**

Using genomic DNA as a template, the rpoB gene was amplified with the following two primer pairs for direct sequencing. Primer pair 1: 5'-AAACCTGTGGCGAT GGTGGGAC-3' (\( \text{pos} \) 1058) and 5'-TAGCCTCGCAG GCCATTCCAC-3' (\( \text{pos} \) 1945). Primer pair 2: 5'-TCTTTCCCCTCGAGGCAGTCC-3' (\( \text{pos} \) 173) and 5'-CACGATGCGGCGGT TGTT-3' (\( \text{pos} \) 1224). The PCR reaction included 1 x Phusion HF buffer (Thermo Scientific, Walthem, MA, USA), 50 pmol each PCR primer, 40 nmol dNTP, 3% dimethylsulfoxide, 0.5 units of Phusion DNA polymerase (Thermo Scientific), 10 ng of genomic DNA, and double-distilled H\(_2\)O. The DNA was denatured at 95\(^°\)C for 4 min, amplified for 30 cycles of 95\(^°\)C for 30 s, 57\(^°\)C for 30 s, and 72\(^°\)C for 1 min and extended for 7 min at 72\(^°\)C. PCR products were purified with the Gel DNA recovery kit (Zymo Research) and sequenced. Primer pair 1: 5'-CGCATTACAAAAGACACTGG CGT-3' (\( \text{pos} \) 323), respectively.

**Survival rates of Dra (dr0022\(^{-}\)) under different stresses**

To induce the nitrosative stress, Dra (\( \text{wt} \)) and Dra (dr0022\(^{-}\)) cells were treated at the indicated concentration of \( \text{NaNO}_2 \) (pH 5.0) for 20 min. Next, cells were washed twice with TGY medium and plated on the TGY plates. To induce \( \text{H}_2\text{O}_2 \) stress, Dra (\( \text{wt} \)) and Dra (dr0022\(^{-}\)) cells were treated with the indicated concentration of \( \text{H}_2\text{O}_2 \) for 20 min. Then, the cells were washed twice with TGY medium and cells were plated on the TGY plates. To induce UV stress, Dra (\( \text{wt} \)) and Dra (dr0022\(^{-}\)) cells were treated under UV-C light for the indicated time. Then, the cells were washed twice with TGY medium and plated on the TGY plates. The irradiation with \( \gamma \)-rays assay was performed as described previously [46]. Briefly, cell suspensions of Dra (wild-type) and Dra (dr0022\(^{-}\)) strains were irradiated at room temperature for 24 h with \( ^{137}\text{Cs} \) \( \gamma \)-rays at various distances from the source, which correspond to various doses (0, 1, 2, 3, 4, 5, 6, 7, and 8 kGy). After
irradiation, cells were plated on TGY plates. All of the plates from the four assays were incubated at 30 °C for 3 days prior to colony enumeration. Survival rates are calculated as a percentage of the number of colonies obtained with untreated cells. All results are the average of three independent experiments.

**De novo prediction and modeling of DR0022 protein structures**

The structures of the wild-type and mutant DR0022 proteins were predicted by ALPHAFOLD2 at the Colab server without templates or homologous structures [36]. The predicted local distance difference test scores calculated by Colab-ALPHAFOLD were used to evaluate the confidence in structural prediction. We also modeled the DR0022 wild-type and mutant protein structures using two other known uracil-DNA glycosylase structures as a template (PDB: 2DP6 and 1UI0) in SWISS-MODEL [47]. The predicted and modeled structures were analyzed with PYMOL (The PyMOL Molecular Graphics System, version 2.0; Schrödinger, LLC, New York, NY, USA).

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**Conflict of interest**

The authors declare no conflict of interest.

**Author contributions**

WC conceived and designed the project. JL designed and performed experiments. JL and WC analyzed data. YY participated in mutation rate experiments. CC assisted with structure prediction and modeling. WC and JL wrote the manuscript.

**Data availability statement**

All data generated or analyzed during this study are included in the published article.

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