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A dual function of the CRISPR-Cas system in bacterial antiviral immunity and DNA repair

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Summary

Clustered Regularly Interspaced Short Palindromic Repeats (CRISPRs) and the associated proteins (Cas) comprise a system of adaptive immunity against viruses and plasmids in prokaryotes. Cas1 is a CRISPR-associated protein that is common to all CRISPR-containing prokaryotes but its function remains obscure. Here we show that the purified Cas1 protein of *Escherichia coli* (YgbT) exhibits nuclease activity against single-stranded and branched DNAs including Holliday junctions, replication forks, and 5'-flaps. The crystal structure of YgbT and site-directed mutagenesis have revealed the potential active site. Genome-wide screens show that YgbT physically and genetically interacts with key components of DNA repair systems, including *recB*, *recC* and *ruvB*. Consistent with these findings, the *ygbT* deletion strain showed increased sensitivity to DNA damage and impaired chromosomal segregation. Similar phenotypes were observed in strains with deletion of CRISPR clusters, suggesting that the function of YgbT in repair involves interaction with the CRISPRs. These results show that YgbT belongs to a novel, structurally distinct family of nucleases acting on branched DNAs and suggest that, in addition to antiviral immunity, at least some components of the CRISPR-Cas system have a function in DNA repair.

Keywords

Cas1; CRISPR; DNA recombination; DNA repair; nuclease; YgbT

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Supporting information

Supporting information is available in the online version of this article.

Introduction

Clustered Regularly Interspaced Short Palindromic Repeats (CRISPRs) represent the most widely distributed family of DNA repeats in prokaryotes (Makarova *et al.*, 2006; Marraffini and Sontheimer, 2010a; Sorek *et al.*, 2008; van der Oost *et al.*, 2009). The CRISPRs and their associated proteins (CRISPR-associated, Cas) appear to comprise a novel microbial defense (immune) system which functions to some extent analogously to the eukaryotic RNA silencing machinery (Makarova *et al.*, 2006; Sorek *et al.*, 2008; van der Oost *et al.*, 2009). CRISPRs are widespread among prokaryotes and are present in approximately 90% of archaeal and approximately 40% of bacterial genomes (Grissa *et al.*, 2007; Horvath and Barrangou, 2010; Karginov and Hannon, 2010; van der Oost *et al.*, 2009).

Most CRISPR-containing prokaryotes possess multiple CRISPR clusters (from 2 to 20 loci), each of which is organized as a tandem array of up to 100 identical repeats of ~25–50 base pairs (Sorek *et al.*, 2008). A unique feature of CRISPRs is the separation of the direct repeats by non-repetitive spacers of similar size (Fig. 1A). The CRISPR clusters are transcribed as multi-unit precursors that are subsequently cleaved into smaller units that consist of one spacer flanked by two partial repeats (Brouns *et al.*, 2008; Hale *et al.*, 2009; Tang *et al.*, 2002; Tang *et al.*, 2005).

Most of the CRISPR spacers lack detectable sequence homologues, but in some organisms a varying, often small fraction of the spacers are homologous to sequences from bacteriophage and plasmid genomes. This key finding suggests that the spacer elements of the CRISPRs represent traces of past invasions by phages and plasmids (Bolotin *et al.*, 2005; Marraffini and Sontheimer, 2010a; Mojica *et al.*, 2005; Pourcel *et al.*, 2005). Recently, a role for CRISPR spacers in defense against specific foreign DNA has been demonstrated in two gram-positive bacteria, *Streptococcus thermophilus* and *Staphylococcus epidermidis* (Barrangou *et al.*, 2007; Marraffini and Sontheimer, 2008 – 2010b), and in an engineered gram-negative species, *Escherichia coli* (Brouns *et al.*, 2008). However, in other studies, the presence of phage-specific spacers in CRISPR clusters of various bacteria failed to confer immunity against the cognate phage (Semenova *et al.*, 2009; van der Ploeg, 2009; Zegans *et al.*, 2009), suggesting that at least some of the CRISPRs may perform cellular functions other than phage immunity. Recently, it has been shown that some of the CRISPR-containing genomes also carry self-targeting CRISPR spacers which are homologous to chromosomal genes and might represent a form of autoimmunity or a regulatory mechanism (Aklujkar and Lovley, 2010; Stern *et al.*, 2010).

CRISPR loci are associated with a large number of Cas protein-coding genes, which have been classified into approximately 45 protein families and 8 types of CRISPR/Cas systems (CRISPR sub-types) (Haft *et al.*, 2005; Makarova *et al.*, 2006). Six Cas protein families (Cas1–6) represent the core group of CRISPR-associated proteins with Cas1 and Cas2 proteins found in all CRISPR-bearing organisms (Haft *et al.*, 2005; Makarova *et al.*, 2006). *Pseudomonas aeruginosa* Cas1 (PaCas1) has a metal-dependent DNase activity but its role in CRISPR function and cell biology remains unknown (Wiedenheft *et al.*, 2009). In contrast, no nuclease activity has been detected in the Cas1 protein SSO1450 from *Sulfolobus solfataricus* (in the presence of 1 mM Mg²⁺ or Ca²⁺) (Han *et al.*, 2009). The small Cas2 proteins possess endoribonuclease activity (Beloglazova *et al.*, 2008; Han and Krauss, 2009), whereas the Cas3 protein is predicted to be a helicase that in many prokaryotes also contains a predicted nuclease domain (Haft *et al.*, 2005; Makarova *et al.*, 2002; Makarova *et al.*, 2006; van der Oost *et al.*, 2009). The biochemical activities of Cas4 and Cas5 are unknown. Cas6 proteins belong to a diverse class of proteins (15 families), named RAMP (Repeat Associated Mysterious Protein) (Haft *et al.*, 2005; Makarova *et al.*, 2006). The RAMP proteins are predicted to function as RNA-binding modules (Makarova *et al.*

al., 2006). Cas6 from the archaeon *Pyrococcus furiosus* (PF1131) can cleave long precursor CRISPR transcripts into small guide RNAs (Carte *et al.*, 2008). In *P. furiosus*, four different RAMP proteins form a complex, which binds to the CRISPR-derived guide RNAs to specifically target and cleave the invader RNA, but not DNA (Hale *et al.*, 2009). By contrast, two studies in *S. epidermidis* and in the engineered *E. coli* suggest that the CRISPR/Cas systems destroy invading DNA rather than RNA (Brouns *et al.*, 2008; Marraffini and Sontheimer, 2008). Thus, the CRISPR systems of archaea and bacteria show a great diversity in spacer and protein composition and appear to use various molecular mechanisms for protection against alien genetic elements.

The *E. coli* K12 strain W3110 contains three CRISPR loci with the spacers showing no homology to known phage or plasmid sequences. The CRISPR locus-1 has 13 spacers (14 repeats) and 8 *cas* genes which encode three core Cas proteins Cas1 (*ygbT*), Cas2 (*ygbF*), and Cas3 (*ygcB*) and five non-core Cas proteins, Cse1 (*ygcL*), Cse2 (*ygcK*), Cse3 (*ygcH*), Cse4 (*ygcJ*), and Cas5e (*ygcI*) (Fig. 1A) (Diez-Villasenor *et al.*, 2010). In *E. coli*, the five non-core Cas proteins have been shown to form a complex, “Cascade”, which processes long CRISPR RNA transcripts into short guide RNAs (~57 nt) (Brouns *et al.*, 2008). The “Cascade” complex and Cas3 (YgcB) can confer phage resistance when the *E. coli* strain is engineered to incorporate CRISPR spacer sequences homologous to an infecting phage λ (Brouns *et al.*, 2008). By contrast, co-expression of Cas1 (YgbT) with Cascade in this strain had no effect on the sensitivity to phage λ , leaving the role of this universal Cas protein enigmatic (Brouns *et al.*, 2008).

To gain insight into the function(s) of Cas1, we characterized the *E. coli* Cas1 protein YgbT using biochemical, genetic and structural approaches, and found that it is a multifunctional nuclease that cleaves Holliday junctions (HJs) and other intermediates of DNA repair and recombination. We found that YgbT interacts physically and genetically with multiple proteins involved in DNA recombination and repair, and that strains lacking YgbT show defects in DNA repair and chromosome segregation. Similar defects are caused by the deletion of the CRISPR repeats. Taken together, these results indicate that, in addition to their role in antiviral immunity, at least some CRISPR-Cas systems function in DNA repair.

Results

YgbT is a nuclease that can cleave linear substrates and Holliday junctions

Recently, it has been shown that the *P. aeruginosa* Cas1 protein is a metal-dependent DNase (Wiedenheft *et al.*, 2009); however, no nuclease activity was found in the Cas1 protein SSO1450 from *S. solfataricus* (in the presence of 1 mM Mg²⁺ or Ca²⁺) (Han *et al.*, 2009). To characterize the biochemical activity of the *E. coli* Cas1 protein YgbT, we purified this protein to homogeneity and tested its activity on a wide range of the ³²P-labeled linear DNA and RNA substrates (Suppl. Table 1). We found a prominent divalent metal cation-dependent nuclease activity against single-stranded (ss) DNAs and a lower activity against ssRNAs (Fig. 1B, 1C; Suppl. Fig. 1A, 1B). YgbT also cleaved short linear double-stranded (ds) DNAs (34 nt), but no cleavage activity was observed with dsRNAs (16 – 39 nt) or longer dsDNAs (60 nt) (Fig. 1D, 1E; Suppl. Fig. 1C).

An activity common to DNA integration and recombination that also requires the cleavage of ssDNA is the resolution of Holliday Junctions (HJs), a cruciform-like DNA intermediate produced by reciprocal strand exchange between two dsDNA molecules (Garcia *et al.*, 2000; Sharples, 2001). HJs are formed in all organisms during DNA integration, recombination, and recombinational DNA repair, as well as the regression or restart of a replication fork. The purified YgbT was tested for the HJ resolving activity using the general HJ substrate HJ1, which contains a homologous core formed by four partially complementary

oligonucleotides, one of which is 5'-³²P-labeled (Lilley and White, 2001). The presence of a homologous core allows the junction point to move freely by branch migration (up/down or left/right) permitting the HJ resolving enzymes to select an optimal sequence or site for cleavage. As shown in Figure 1F, purified YgbT displayed significant activity against the HJ1 substrate, producing a nicked duplex with electrophoretic mobility similar to that of the RuvC product. The cleavage of HJ1 was proportional to the concentration of YgbT (Fig. 1G) with the highest activity at pH 8.5 in the presence of Mg²⁺ (10 mM) (Figure 1H) and K⁺ (100 mM) (data not shown).

To characterize the cleavage pattern of YgbT, the products of HJ1 resolution were analyzed by denaturing PAGE and compared with the products of RuvC which cleaves this substrate at one major site in strands A and C and two major sites in strands B and D (Fig. 1I and J). In contrast to RuvC, YgbT did not show any pronounced sequence preference in the cleavage of HJ1 and introduced multiple symmetrical nicks in strands A and C, as well as in B and D (Fig. 1I, 1J). Strands A and C were cleaved preferentially on the 5'-side of the homologous core, whereas a distinct preference for cleavage at the center of the substrate was apparent in strands B and D (Fig. 1I, 1J). The HJ cleavage pattern of YgbT is similar to that of the human HJ resolvases MUS81-EME1 and SLX1 which likewise exhibit low sequence selectivity and introduce multiple nicks close to the 5'-side or at the center of the substrate (Constantinou *et al.*, 2002 ; Svendsen *et al.*, 2009). To determine whether YgbT, similarly to RuvC, resolves HJs through the introduction of ligatable nicks, we used the asymmetric substrate HJ2. The treatment of RuvC cleavage products with T4 DNA ligase yielded two main ligation products, whereas re-ligation of the YgbT cleavage products generated several products (indicated by the arrows) (Fig. 1K). Formation of several re-ligatable products was also observed with human HJ resolvase MUS81-EME1 (Constantinou *et al.*, 2002). Hence, analogously to other HJ cleaving nucleases, YgbT cleaves HJs with the formation of ligatable products.

YgbT is a branched DNA nuclease

In addition to HJs, many known HJ cleaving nucleases can cleave other branched DNA substrates such as replication forks, Y-junctions, splayed arms, and 3'- and 5'-flaps (Abraham *et al.*, 2003; Benson and West, 1994 ; Ciccio *et al.*, 2008; Sharples, 2001; Svendsen *et al.*, 2009). The substrate specificity of YgbT was further characterized using a series of branched 5'-³²P-labeled DNA substrates containing sense and antisense sequences of the *E. coli* CRISPR repeat, including a static HJ (HJ3), replication fork, 5'-flap, 3'-flap, and splayed arm duplex, structures that mimic various DNA repair and recombination intermediates (Fig. 2A). At pH 8.5, YgbT cleaved all these substrates with variable efficiency, whereas at pH 7.0 it was more specific and showed significant activity only against HJ3 (Fig. 2A). Purified YgbT also bound to 5'-flaps structures, HJs, ssDNAs and ssRNAs, and produced oligomeric complexes, as detected using mobility shift assays (Fig. 2B, only 5'-flap and HJ are shown). Like RuvC (Benson and West, 1994), YgbT cleaved the three-way junction substrate (3wHJ) but showed no activity against a Y-junction or a heterologous loop (data not shown). Analysis of the cleavage products by denaturing PAGE revealed that YgbT introduced a limited number of cuts into the replication fork and 5'-flap structures, but produced multiple cleavage products with the HJ3, 3'-flap, and splayed arm substrates (Fig. 2C, 2D).

In *E. coli* and many other CRISPR-containing genomes, CRISPRs encompass palindromic repeats that can potentially generate cruciform (CF)-like structures. To test whether these structures represent possible substrates for YgbT, we designed two artificial CF-like substrates (CF-1 and CF-2) using oligonucleotides that correspond to the sequence of the *E. coli* CRISPR repeat-1 (Suppl. Table 1). YgbT efficiently cleaved both CF-like substrates and generated multiple cleavage products (Fig. 2E, 2F; only CF-1 is shown). With the CF-1

substrate, the five main cleavage products had the lengths of 5 nt, 6 nt, 29 nt, 31 nt, and 34 nt (Fig. 2E, 2F).

Crystal structure and site-directed mutagenesis of YgbT

The recently published crystal structure of *PaCas1* revealed the presence of a small N-terminal β -strand domain (residues 1–106) connected by a flexible linker to a larger α -helical domain (residues 113–324) (Wiedenheft *et al.*, 2009). We cloned and purified the C-terminal domain of YgbT without the N-terminal domain (residues 96–278) and found that the C-terminal domain retained all the activities observed in the intact YgbT protein (HJ cleavage, ssDNase and ssRNase), indicating that this domain contains the active site of YgbT (Suppl. Fig. 1D, 1E). We then determined the crystal structures of both the YgbT C-terminal domain (residues 96–278; 1.40 Å resolution; PDB code 3NKE) and the full-length protein (1.95 Å resolution; PDB code 3NKD) (Fig. 3, Suppl. Table 3). The structure of the full-length YgbT showed that this protein is a homodimer (Fig. 3A), consistent with our gel-filtration results ($M_r = 70$ kDa; predicted monomer $M_r = 33.2$ kDa). Like *PaCas1* and the Cas1 protein aq_369 from *Aquifex aeolicus* (PDB code 2YZS), the YgbT monomer consists of two domains: a small N-terminal domain (amino acids 1–89) with a beta-sandwich-like structure connected by a flexible linker (aa 90–95) to a larger, mostly α -helical, C-terminal domain (aa 96–305) (Fig. 3B). Our gel-filtration experiments showed that the purified C-terminal domain of YgbT (predicted monomer M_r 20 kDa) exists as a mixture of the dimeric (M_r 38 – 46 kDa; 5% to 20%) and monomeric (M_r 22 – 27 kDa; 80% to 95%) forms in solution.

A Dali search (Holm and Sander, 1998) for structures similar to that of YgbT identified three other Cas1 proteins as the best hits including *PaCas1* (3GOD, Z-score 21.9, rmsd 3.4 Å) and two unpublished structures of Cas1 proteins from *A. aeolicus* (aq_369, PDB code 2YZS, Z-score 15.3, rmsd 3.4 Å) and *Thermotoga maritima* (TM1797, PDB code 3LFX, Z-score 16.6, rmsd 4.6). However, these proteins share relatively low overall sequence similarity (16–23% sequence identity) and belong to different CRISPR subtypes (CASS-2: YgbT; CASS-3: *PaCas1*; CASS-7: aq_369 and TM1797) (Makarova *et al.*, 2006) (Suppl. Fig. 2). Surface charge analysis of YgbT revealed the presence of several large patches of positively charged residues which represent potential DNA-binding sites (Fig. 3C). In *PaCas1*, several basic residues surround the negatively charged metal-binding site creating a potential nucleic acid binding site in proximity to the catalytic metal cation (Wiedenheft *et al.*, 2009). YgbT contains a larger cluster of highly conserved basic residues (R112, R123, R138, R146, R163, K211, and K224) positioned around the potential metal-binding site (E141, H208, D221) (Fig. 3C, 3D). In addition, YgbT contains another cluster of positively charged residues located at the same protein side near the connection of the two domains (K37, R59, R84, R245, R248, and R252); this cluster has no counterpart in *PaCas1*, suggesting that these proteins might differ in the details of the substrate binding (Fig. 3C). Surface screen analysis (Binkowski and Joachimiak, 2008) showed that the surface of the YgbT main basic patch is similar to that of the ssDNA-binding site of *E. coli* topoisomerase III (PDB code 1I7D), which binds ssDNA through the direct or water-mediated interactions with the phosphate groups of the DNA phosphodiester backbone (Changela *et al.*, 2001). Fig. 3E shows the possible position of ssDNA modeled on the YgbT surface based on the superposition of the YgbT and topoisomerase III surfaces.

Previous sequence analysis of Cas1 proteins showed that four strictly conserved residues (three carboxylates and one histidine) represent the signature motif of the Cas1 family (Makarova *et al.*, 2002). Two of these residues have been shown to be required for the nuclease activity of *PaCas1* (D265 and D268) (Wiedenheft *et al.*, 2009). In YgbT, the signature residues (E141, H208, D218, and D221) form a cluster of closely positioned side chains (2.5 to 6.6 Å) located at the bottom of the most prominent cleft of the C-terminal

domain (Fig. 3F). These four residues are surrounded by the positively charged side chains of 7 highly conserved residues (R112, R123, R138, R146, R163, K211, and K224) which form the main basic patch of the YgbT surface and are potentially involved in the coordination of the phosphate backbone of bound DNA. In the structure of the YgbT C-terminal domain, three of these residues (R112, R138, and R163) interact with two bound sulfates which can be construed as mimicking the phosphates of the DNA backbone (Fig. 3F). The third sulfate molecule in this structure is bound to three residues from the second basic surface patch (R245, R248, and R252) (Fig. 3F).

Alanine replacement mutagenesis of YgbT showed that its nuclease activity against all substrates was abolished in the purified mutant proteins E141A, H208A, D218A, and D221A (Fig. 2G: 5'-flap and HJ; Suppl. Fig. 1F: ssRNA; Suppl. Fig. 1G), in accord with the prediction that these residues contribute to the active site. Mutations of other residues near the potential active site (T184A, K211A, and K224A) also had strong inhibitory effects on the activity of YgbT, whereas the E135A, Y165A, and Y188A mutant proteins retained significant activity (Fig. 2G), suggesting that the active site of YgbT is located on its C-terminal domain close to the potential ssDNA-binding site.

Taken together, these results indicate that YgbT is a nuclease that can cleave HJs and other branched DNA substrates, as well as linear ss/dsDNAs and ssRNAs. The ability of YgbT to cleave branched DNA substrates potentially could contribute to the addition of new spacers to CRISPR clusters. Moreover, the identified activity of YgbT against branched DNAs suggests that this protein might also participate in some of the DNA repair or recombination pathways. This possibility is consistent with previous studies which proposed a function for the CRISPR system in DNA repair or in chromosomal segregation (DeBoy *et al.*, 2006; Makarova *et al.*, 2002; Mojica *et al.*, 1995).

YgbT is associated with proteins involved in DNA repair

To probe the potential role of YgbT in DNA repair and recombination, we performed genome-wide assays for physical and genetic interactions between YgbT and other *E. coli* proteins. Endogenous YgbT was C-terminally tagged using a cassette encoding a calmodulin binding peptide and a triple FLAG-tag (Zeghouf *et al.*, 2004). Proteins stably associated with the tagged YgbT were treated with DNase and affinity-purified in two steps on anti-FLAG and calmodulin resins and then identified using tandem mass spectrometry (LC-MS) (Butland *et al.*, 2005). Among the proteins co-purifying with YgbT (Suppl. Table 3) were RecB and RecC, two subunits of the RecBCD nuclease-helicase (recombinase) complex, which is a major player in the recombinational repair of DNA double-strand breaks (DSBs) (Kowalczykowski, 2000). In addition, the DNA repair proteins that co-purified with YgbT included RuvB, a recombinational DNA helicase which, together with RuvA, binds to HJs and catalyzes HJ branch migration to facilitate the cleavage of HJs by the RuvC resolvase (West, 1996). Although no physical interactions between YgbT and other Cas proteins in *E. coli* have been reported previously (Brouns *et al.*, 2008), we reproducibly detected two subunits of the CRISPR-associated Cascade complex, YgcJ (CasC) and YgcH (CasE) (Fig. 4A, Suppl. Table 3). Interestingly, purified recombinant YgcH (a Cascade subunit) inhibited the HJ cleavage activity of YgbT in a concentration-dependent manner (Fig. 4B), suggesting that YgbT and the Cascade complex might interact and regulate each other's activities.

The observed physical interactions of YgbT with DNA repair/recombination proteins (RecB, RecC, RuvB, and UvrC) and with two other Cas proteins (YgcJ and YgcH) were validated by reciprocally purifying C-terminally tagged RecB, RecC, RuvB, UvrC, YgcJ and YgcH proteins. In each case, mass spectrometry analyses of the affinity-purified protein confirmed its association with YgbT (Fig. 4A, Suppl. Table 3). We also confirmed the physical association of YgbT with RecB, RecC, RuvB, YgcH and YgcJ by co-immunoprecipitation

using cell lysates from exponential phase cultures of strains expressing C-terminal affinity-tagged YgbT; a strain expressing tagged Fis, a nucleoid associated DNA-binding protein which modulates gyrase (Cho *et al.*, 2008) and topoisomerase I production (Weinstein-Fischer and Altuvia, 2007), was used as a control. The tagged YgbT was immunoprecipitated from the cell lysates, and the presence of RecB, RecC and RuvB was probed by immunoblotting using anti-RecB, -RecC, or - RuvB antisera, or anti-(His₆)-tag monoclonal antibody (to detect the (His₆)-tagged YgcH or YgcJ) (Fig. 4C, 4D). As expected, all these proteins (RecB, RecC, RuvB, YgcH and YgcJ) co-precipitated from extracts containing tagged YgbT but not from control extracts prepared from an untagged parental *E. coli* strain or the Fis-tagged strain. Thus, our results reveal a stable physical association of the *E. coli* Cas1 protein YgbT with some of the key proteins involved in DNA repair pathways and with two other Cas proteins, YgcH and YgcJ.

YgbT genetically interacts with DNA repair-recombination pathways

To identify *E. coli* genes that genetically interact with *ygbT*, we used the recently developed *E. coli* Synthetic Genetic Array (eSGA) approach (Butland *et al.*, 2008). Double mutants were constructed by conjugating a *ygbT* donor strain (marked with $\Delta::\text{Cm}^R$) with single gene deletions of almost all other non-essential *E. coli* genes (recipients marked with $\Delta::\text{Kan}^R$) or hypomorphic alleles of selected essential *E. coli* genes (marked with Kan^R) (Suppl. Table 4, Suppl. Information). The colony growth and relative fitness of the resulting double mutants that survived double drug selection were examined by digital imaging. Using a statistical interaction score (*S*) to quantify both the strength and confidence of the interactions between each *E. coli* mutant gene pair, we detected cases of both synthetic sick or lethal (SSL; i.e. negative or aggravating interactions) and alleviating (i.e. positive or suppressing) growth phenotypes (Suppl. Table 4, Suppl. Information). In congruence with the results of the protein-protein interaction experiments, we observed genetic interactions of *ygbT* with DNA repair and recombination genes, including SSL interactions with *recBC* and *recN* and strong alleviating interactions with the *ruvABC* genes (Suppl. Table 4). Over-representation analysis (Irizarry *et al.*, 2009) using Gene Ontology annotation terms (Suppl. Information) showed that the interacting genes identified in the *ygbT* deletion screen were significantly enriched (*q*-value < 0.005) for genes involved in DNA repair, DNA recombination, cell division, and chromosomal segregation and condensation (Suppl. Table 5).

Because the eSGA method depends on recombination in the recipient cells and genes like *recBC* are important for recombination, we validated the observed genetic interactions of *ygbT* with *recABCD* and *ruvABC* in reciprocal conjugation experiments using *recABCD* and *ruvABC* single gene deletion mutants as donors and $\Delta ygbT$ as recipient (Fig. 5A, 5B). Consistent with the *ygbT* genetic interaction results, the recipient *ygbT* deletion strain exhibited a striking suppression (alleviating) phenotype when combined with individual gene deletions of $\Delta ruvABC$ donor strains. Conversely, a significant SSL growth defect was consistently observed between the $\Delta ygbT$ recipient and the $\Delta recB$ and $\Delta recC$ donor mutants, whereas hardly any growth deficiency was seen with *recA* and *recD* donor mutants (Fig. 5B). These genetic interactions were not a consequence of recombination suppression resulting from gene proximity because no detectable effects on growth were observed when *recBC* or *ruvABC* deletion donors were combined with deletions of functionally unrelated genes flanking *ygbT*. Moreover, the use as donors of other, functionally unrelated genes from the same genome region, namely *csdA* and *rpoS*, did not reveal synthetic genetic interaction with *ygbT*, *recABCD* or *ruvABC*, except in the case of *csdA* and *recB* which are too close to one another (Fig. 5A, 5B). Recently, Pougach *et al.* (Pougach *et al.*, 2010) have demonstrated substantial readthrough transcription from the kanamycin resistance cassette inserted into several *cas* genes (*ygL*, *ygK*, or *ygJ*) located upstream of *ygbT* (Fig. 1A) using *E. coli* strains from the Keio collection, which was also used in our work. However,

the readthrough transcription from the kanamycin resistance cassette inserted into *ygbT* is unlikely to have a substantial effect on the observed genetic interactions, because only one *cas* gene (*ygbF*) is located downstream of *ygbT* (Fig. 1A). Thus, our results suggest that the observed genetic interactions between *ygbT* and *recBC* and *ruvABC* represent *bona fide* functional relationships. The detection of a SSL association in the first case and an alleviating interaction in the second case indicates that the putative function of YgbT in repair might be redundant with the function of RecBC but cooperates with the function of RuvABC.

In addition, *ygbT* exhibited a strong SSL interaction with *recN* and weak SSL interactions with other *rec* genes (*recF*, *recO* and *recQ*) in the RecF recombination pathway (Suppl. Table S5). In *E. coli*, the major homologous recombination mechanism is the RecBC pathway, which is involved in both conjugal and transductional recombination, as well as in the repair of DSBs and the degradation of foreign DNA, whereas the RecF recombinational repair pathway is implicated preferentially in the repair of UV-induced daughter strand gaps (Kuzminov, 1999; Tseng *et al.*, 1994). Thus, the SSL interactions of *ygbT* with *recBC* and *recF* genes suggest that YgbT could be involved in both pathways or yet a third parallel pathway of DSB repair.

The $\Delta ygbT$ screen also identified genetic interactions outside the Rec and Ruv systems, including synthetic lethality following inactivation of the site-specific recombinase XerD, the chromosome partitioning protein MukB, and the essential cell division protein FtsK (Suppl. Table 4). These proteins function in chromosome segregation and the separation of replicated chromosome dimers, which involves the dynamic formation and resolution of HJs and other branched DNA intermediates (Barre *et al.*, 2001; Gordon and Wright, 2000.; Sherratt *et al.*, 2001). Aggravating genetic interactions were also observed with other genes involved in cell division (e.g., *dinF*, *envC*, *gida*, *gidB* and *dicB*) and in methyl-directed mismatch repair (MMR) (e.g., *mutH*, *mutL* and *mutS*) (Suppl. Table 4). Thus, the genetic interaction data appear to support the functional link between YgbT and DNA repair-recombination.

YgbT is required for the resistance of *E. coli* to DNA damage

Our biochemical results together with physical and genetic interaction data pointed to a role of YgbT in DNA repair-recombination pathways and to a cooperation between YgbT and RuvABC. To investigate the biological implications of these results, we compared the sensitivity of the *E. coli ygbT* deletion strain to DNA damage induced by either the genotoxic agent mitomycin C (MMC) or UV light with the sensitivities of wild-type or *ruvABC* deletion strains. Both MMC and UV introduce an array of lesions, including pyrimidine dimers (UV) and inter-strand cross-links (MMC), leading to the formation of DSBs in DNA (De Silva *et al.*, 2000; Garinis *et al.*, 2005). In *E. coli*, UV- or MMC-induced DNA damage is repaired by a multitude of pathways, including homologous recombination and nucleotide excision repair (Cole, 1973; Keller *et al.*, 2001). Strains with individually deleted *ruvABC* genes are known to be substantially more sensitive than the wild-type to UV- or MMC-induced DNA damage (Le Masson *et al.*, 2008; Lloyd *et al.*, 1984), and there is little difference in the sensitivity among the strains carrying mutations in each of the *ruv* genes (Lloyd *et al.*, 1984; Otsuji *et al.*, 1974; Sargentini and Smith, 1989). In our experiments, knockout of *ygbT* increased the sensitivity of *E. coli* cells to both MMC and UV to approximately the same extent as the knockout of each of the *ruv* genes (Fig. 5C, 5D). The sensitivity of double deletion combinations of *ygbT* with the *ruvABC* genes to MMC and UV was similar to each of the single knockout mutants (Fig. 5C, 5D). Moreover, the sensitivity of *ygbT-ruvABC* double mutants to cisplatin (40 μ M), another genotoxic agent known to induce DSBs that are repaired by excisional and recombinational repair pathways (Zdraveski *et al.*, 2000), was also comparable with the sensitivities of the single

mutants (data not shown). Thus, the lack of synergy in the *ygbT-ruv* double knockouts suggests, in agreement with the eSGA results, that YgbT functions in the same DNA repair pathway(s) with RuvABC.

To ascertain that the observed phenotype was caused solely by the *ygbT* deletion, we showed that the MMC and UV resistance of the *ygbT* null strain could be restored to the wild type level by introducing a pBAD-plasmid expressing the wild-type *ygbT* gene under the control of an arabinose-inducible promoter but not *ygbT* mutants with replacements of the predicted catalytic amino acid residues (E141A, H208A and D218A) (Fig. 5C, complementation experiment with UV not shown). Similarly, the resistance of *ygbT-ruv* double mutants to MMC and UV-irradiation was fully restored to wild-type levels by ectopic expression of both YgbT and the respective Ruv proteins (data not shown). Thus, the nuclease activity of YgbT appears to be critical for the resistance of *E. coli* to DNA damage.

YgbT is recruited to DNA damage sites in MMC-treated cells

In bacteria, MMC treatment induces the formation of DSBs at replication forks, which then rapidly recruit proteins involved in DNA repair (e.g., RecN) leading to the formation of discrete foci detectable using fusions with appropriate tags (Kidane *et al.*, 2004; Sanchez *et al.*, 2006). To test whether YgbT is likewise recruited to DSBs in *E. coli*, we constructed chromosomal yellow fluorescent protein (YFP) fusions (Taniguchi *et al.*, 2010) in which YgbT as well as RuvA, RuvB, and RuvC genes were tagged with the YFP coding sequence at their C-terminal ends. A strain expressing a YFP fusion of RecN was used as a positive control. Fluorescence microscopy showed that YFP-labeled YgbT, as well as RuvABC and RecN, formed discrete fluorescent foci on the nucleoids after 120 min of MMC treatment. In YgbT-YFP and RuvABC-YFP strains, 13–17% of the > 250 analyzed cells contained a single focus per nucleoid (Fig. 6A, 6B), whereas in the RecN-YFP strain, the fusion protein localized to a discrete focus in the majority (~70%) of cells. Although it remains unclear why a smaller fraction of MMC-treated cells had YgbT or RuvABC localized to foci compared with RecN, no foci were detected in the absence of MMC in YgbT-YFP or RecN-YFP expressing cells (Suppl. Fig. 3A). Thus, YgbT appears to be specifically recruited to DNA DSBs, in agreement with the other indications of a role of this protein in DNA repair.

YgbT is required for cell division in MMC-treated cells

Strains showing hypersensitivity to MMC, such as *ruvABC*, *recN*, *recF* and *recO* loss-of-function mutants, exhibit impaired cell division and chromosomal segregation in the presence of unrepaired DNA damage, leading to the formation of abnormally long, non-septate, multi-nucleate cells (Ishioka *et al.*, 1998; Kidane *et al.*, 2004; Meddows *et al.*, 2005; Otsuji *et al.*, 1974; Suzuki *et al.*, 1967; Zahradka *et al.*, 1999). This unusual morphology might be a consequence of the inability of the cells to resolve chromosome dimers. The demonstration of the HJ cleavage activity of YgbT and the physical and genetic interactions between YgbT and several repair proteins described above suggest that YgbT could also be involved in resolving chromosomes during cell division. A role of YgbT in chromosome segregation is also suggested by its aggravating genetic interactions with several genes known to be involved in cell division and chromosome segregation factors (e.g., FtsK, MukB, and XerD) (Suppl. Table 4). Should YgbT participate in the resolution of sister chromosomes, the $\Delta ygbT$ strain would be expected to form abnormal filaments in the presence of MMC. Indeed, we found that the *ygbT* deletion strain formed greatly elongated cells containing one long nucleoid after 120 min of MMC treatment (Fig. 6C). The average length of *ygbT* mutant cells ($\sim 5.8 \pm 0.3 \mu\text{m}$) in the presence of MMC was almost twice that of wild type cells (from ~ 2.5 to $3.9 \mu\text{m}$) (Suppl. Fig. 3B). This extent of cell elongation was similar to that seen in *recN*, *recF*, or *recO* deletion strains (from ~ 4.7 to $6.1 \mu\text{m}$) or *ruvABC* single mutants (from $\sim 5.0 \pm 0.6$ to $5.9 \pm 0.3 \mu\text{m}$). Double mutants of *ygbT* with *ruvABC*, but

not with *recN*, *recF*, or *recO*, produced even longer filamentous cells in the presence of MMC, with average cell lengths ranging from $\sim 8.6 \pm 0.8$ to 9.7 ± 0.3 μm after 120 min of MMC treatment (Fig. 6C, Suppl. Fig. 3B). Thus, our findings indicate that YgbT might be involved in chromosome segregation following DNA damage. This result is consistent with observations on the archaea *Haloflex volcanii* and *H. mediterranei* which implicated the CRISPR repeats in replicon partitioning (Mojica *et al.*, 1995). Surprisingly, when it comes to chromosome segregation, YgbT appears to function in cooperation with the RecF pathway and redundantly with RuvABC.

Association of YgbT with CRISPR clusters

The CRISPR repeats of *E. coli* and many other organisms contain a 5–7 nt palindromic repeat (Jansen *et al.*, 2002; Kunin *et al.*, 2007; Sorek *et al.*, 2008). Such palindromes could potentially form cruciform structures (in dsDNA) or hairpins (in ssDNA), whereas direct tandem repeats can form slipped-strand DNA (S-DNA) with mispaired complementary repeats (Lilley, 1989; Mirkin and Mirkin, 2007). These unusual DNA structures can interfere with DNA replication, resulting in replication fork stalling and genomic instability (Lindsey and Leach, 1989; Mirkin and Mirkin, 2007). As the *cas1* gene is present only in bacteria and archaea that also possess CRISPRs, functional coupling between the Cas1 protein and CRISPRs appears most likely (Brouns *et al.*, 2008; Makarova *et al.*, 2006). The *E. coli* K12 laboratory strain W3110 carries three CRISPR clusters (Cluster-1 with 14 repeats, Cluster-2 with 7 repeats, and Cluster-3 with 3 repeats), with the *cas1* (*ygbT*) gene located close (~ 300 bp) to CRISPR cluster-1 (Fig. 1A). Taking into account the function of YgbT in DNA recombination-repair that is strongly suggested by the results of this work and the obligate association of *cas1* with CRISPR clusters, we speculated that the repair function of YgbT might involve alleviating the potential deleterious effect of CRISPR. Should that be the case, the requirement for YgbT in DNA repair would be lifted by deletion of the CRISPR clusters. To test this possibility, we constructed *E. coli* strains lacking *ygbT* and either CRISPR cluster-1, or cluster-2, or both CRISPR clusters, and compared their MMC sensitivities with that of the *ygbT* mutant strain (Fig. 6D). The results showed that the requirement of *ygbT* for MMC resistance was unrelated to the prevention of the potential deleterious effect of CRISPRs because the triple deletion of both CRISPR clusters and *ygbT* exhibited MMC sensitivity similar to that of the *ygbT* mutant (Fig. 6D).

Interestingly, deletion of one or both CRISPR clusters itself increased the sensitivity of *E. coli* to MMC (Fig. 6D). One possible explanation of this observation could be that CRISPR clusters are required for the expression of YgbT (Pul *et al.*, 2010). However, this appears not to be the case because the assembly of YgbT at DSBs was independent of the presence of either CRISPR cluster (Fig. 6A, 6B). Thus, CRISPR clusters are apparently required for the function of YgbT in DNA repair but not for YgbT recruitment to MMC-induced DSBs. To elucidate the molecular mechanism of the CRISPR-dependent role of YgbT in repair, additional experiments are obviously required.

Discussion

The results presented here indicate that the *E. coli* Cas1 protein YgbT is a novel nuclease that can cleave HJs and other branched DNA substrates, as well as linear ss/ds DNAs and ssRNAs. To date, *E. coli* is known to encode one functional HJ resolvase, RuvC, and one cryptic resolvase, RsaA, which is normally not expressed (Benson and West, 1994; Sharples *et al.*, 1994). In contrast to RuvC, YgbT shows no apparent target sequence specificity. In addition, YgbT cleaves *in vitro* a broad range of branched DNA substrates (asymmetrical Holliday junction, replication fork, 5'-flap, 3'-flap and splayed arm substrates) which represent various intermediates of DNA recombination and repair (Fig. 2). Like human MUS81-EME1 and SLX1, YgbT produces multiple HJ cleavage products and

can also cleave replication fork and 3'-flap structures; however, in addition, YgbT is active with 5'-flap and splayed arm substrates which are not cleaved by MUS81 (Constantinou *et al.*, 2002). The 5'-flap endonucleases, which are required for the removal of RNA primers during replicative and repair DNA synthesis and typically can cleave both RNA and DNA, are encoded as distinct proteins in eukaryotes (FEN-1), archaea and some DNA viruses, whereas their bacterial homologs are N-terminal domains of DNA polymerase I (Shen *et al.*, 1998). Given that YgbT cleaves both ssRNA and 5'-flap DNAs but is unrelated to the FEN-1 family it might represent a new group of stand-alone flap endonucleases.

When compared to one another, the three experimentally characterized Cas1 proteins display related but different biochemical properties: SSO1450 binds DNA/RNA but appears not to possess nuclease activity, *PaCas1* cleaves ss/dsDNA, whereas YgbT is also active against branched DNA substrates and linear ssRNAs. These proteins belong to different CRISPR subtypes (Makarova *et al.*, 2006) and share rather low overall sequence similarity (18.9 – 21.6 % of sequence identity), suggesting that they might possess genuinely different substrate preferences. This possibility is consistent with the presence of several (2 to 5) distinct paralogous Cas1 proteins in many bacteria and archaea. Further, detailed biochemical studies of Cas1 proteins from diverse organisms are required to delineate the functional versatility of this protein family.

The ability of YgbT to cleave branched DNA substrates *in vitro* suggests that this activity might contribute to the addition/removal of CRISPR spacers, which is proposed to proceed through DNA recombination (Mojica *et al.*, 2009). Four strictly conserved amino acid residues of YgbT (E141, H208, D218 and D221) are critical for activity and represent its active site, which is located close to the potential DNA-binding site in the C-terminal domain. Further detailed analysis of the role of YgbT in the CRISPR mechanism requires the use of a natural experimental model of CRISPR spacer addition/removal, which has yet to be established in *E. coli*.

The incorporation of a novel CRISPR spacer and the accompanying repeat apparently involves the recognition of foreign DNA followed by processing steps and insertion into the CRISPR cluster. Given that YgbT showed no obvious sequence selectivity in the cleavage of branched DNAs, we hypothesize that other Cas (and non-Cas) proteins are also involved in the formation of novel CRISPR spacers, whereas YgbT might contribute to one of the final steps in the integration of the spacers into the chromosomal or plasmid DNA (e.g., HJ resolution or flap substrate cleavage). The physical interaction between YgbT and two components of the Cascade complex (Cse4/CasC and Cse3/CasE) reported in this work also suggests that Cascade might contribute to the integration of new spacers in *E. coli*.

The key conclusion of the present work is that YgbT and CRISPR are involved in one or more repair-recombination pathways and contribute to the resistance of *E. coli* to at least some types of DNA damage and chromosome segregation. This conclusion is consistently supported by several lines of genetic and biochemical evidence: (1) knockout of the *ygbT* gene results in a substantial increase in the sensitivity of *E. coli* to DNA damage caused by UV or MMC; (2) the rescue of the knockout mutants requires catalytically active YgbT, indicating that the apparent role of YgbT in repair-recombination depends on its demonstrated endonuclease activity towards the characteristic intermediates of several DNA repair pathways, including recombinational (HJs and splayed arms), base excision (5'-flaps), and nucleotide excision (3'-flaps) pathways; (3) YgbT physically interacts with several key repair proteins including RecB, RecC and RuvB; (4) the *ygbT* gene genetically interacts with repair genes, including synthetic-sick interactions with *recB*, *recC* and *recN*, indicative of functional redundancy, and alleviating interactions with *ruvABC*, indicative of involvement in the same repair-recombination pathway(s); (5) YgbT is recruited to DSBs in MMC-

treated cells; (6) YgbT is required for cell division in MMC-treated cells, as evidenced by the unusual morphology of *ygbT* knockouts that resembles the morphology of knockout mutants of other repair genes upon treatment with DNA-damaging agents. In addition, we found that the function of YgbT in DNA repair apparently requires interaction with CRISPRs because deletion of the CRISPRs or the double deletion of the CRISPRs and *ygbT* led to the same phenotype as the *ygbT* knockout. The specific role of CRISPRs in repair remains to be elucidated, but the involvement of their recombinogenic potential seems plausible; previously, it has been suggested that CRISPRs mediate genome rearrangements in *Thermotogales* via recombination (DeBoy *et al.*, 2006).

Thus, YgbT appears to be a multifunctional nuclease that can cleave various branched DNA intermediates produced by DNA repair pathways, chromosome segregation mechanisms, and (potentially) by the CRISPR system. More generally, our results suggest an intrinsic connection between the function of the CRISPR-Cas system in the acquired antiviral immunity and its emerging role in DNA repair in prokaryotes. These findings reconcile the latest observations on the antiviral functions of this system that depend on unique spacers homologous to viral genes (Horvath and Barrangou, 2010; Karginov and Hannon, 2010) and the earlier hypothesis on a repair function of the Cas proteins (Makarova *et al.*, 2002) that was proposed before the discovery of the virus-specific spacers, solely on the basis of the predicted enzymatic activities of the Cas proteins (a helicase, a polymerase and multiple nucleases). Furthermore, a dual function of the CRISPR-Cas system in defense and repair is compatible with the typically small fraction of virus-specific CRISPR spacers and the inability of some of these spacers to protect the host against infection (Semenova *et al.*, 2009; van der Ploeg, 2009; Zegans *et al.*, 2009). It is conceivable that the array of diverse enzymes and nucleic acid-binding proteins associated with the CRISPRs evolved at early stages of the evolution of bacteria and archaea within the intrinsically interlinked contexts of DNA damage repair and antiviral defense. Further research into the functions of various CRISPR-Cas systems should reveal the relationships between its apparently diverse functions and more specific roles of the individual components.

Experimental procedures

Bacterial strains, gene cloning, and protein purification

All parental and mutant bacterial strains are listed in Suppl. Table 6 and are described in Suppl. Methods. For enzymatic assays and crystallization, the proteins were over-expressed in the *E. coli* BL21 (DE3) strain (Novagen) and affinity purified as previously described (Proudfoot *et al.*, 2008; Zhang *et al.*, 2001) (Suppl. Methods). This *E. coli* strain has been shown to contain no *cas* genes (Brouns *et al.*, 2008).

Nuclease assays

DNA substrates used in this work and nuclease assays are described in Suppl. Table 1 and Suppl. Methods. The reaction products were analyzed by native (8% PAAG) or denaturing (15% polyacrylamide/8M urea PAAG) electrophoresis and visualized by autoradiography.

Crystallization and structure determination

The Se-Met-substituted C-terminal domain of YgbT (aa 92 – 281) and native two-domain YgbT (aa 7–281) were crystallized by the hanging drop vapor diffusion method as described in Suppl. Methods. The structure of the YgbT C-terminal domain was determined using the single anomalous scattering dispersion (SAD) method; whereas the structure of the two-domain YgbT was solved by molecular replacement using the structure of the YgbT C-terminal domain as a model (Suppl. Table 2 and Suppl. Methods).

SPA-tagging, purification and immunoprecipitation

YgbT, Ruv, Rec and Cas proteins were C-terminally tagged and purified essentially as previously described (Butland *et al.*, 2005). The co-purifying proteins were identified using SDS-PAGE fractionation followed by peptide mass fingerprinting or using gel-free liquid chromatography/tandem mass spectrometry (LCMS) essentially as previously described (Babu *et al.*, 2009). The physical association of YgbT with RecB, RecC, RuvB, YgcH and YgcJ proteins that were confirmed by co-immunoprecipitation are described in detail in Supplementary Methods.

Genome-wide eSGA screen and gene set enrichment analysis

A genome-wide eSGA screen using *ygbTA::Cm^R* in Hfr Cavalli as donor was carried out and analyzed as previously described (Butland *et al.*, 2008), as were various mini-array crosses. Gene set enrichment analysis (Irizarry *et al.*, 2009) was performed on the interaction *S* scores to identify the significantly enriched GO terms (Suppl. Table S6) spanning various biological processes (see Suppl. Methods).

DNA damage sensitivity assays

UV irradiation experiments were performed as described previously (Nair and Finkel, 2004). Diluted cultures were plated onto LB plates and colonies formed after overnight incubation at 32 °C were counted. Cell survival results were derived from the mean of three independent experiments. For the MMC and cisplatin sensitivity assays, exponentially growing cells were serially diluted in LB medium and pinned onto LB plates in the absence or presence of the indicated concentration of the DNA damaging agent. Sensitivity to DNA damage was assessed after 36 to 48 hrs of incubation at 32 °C.

Cell morphology and protein localization analysis

The C-terminal fusions of chromosomal YgbT, RuvA, RuvB, RuvC, RecN, and MutH to YFP proteins were constructed in the DY330 background by converting the SPA tags and kanamycin resistance cassette in these strains to YFP tags via the λ -RED recombination system (Yu *et al.*, 2000). PCR amplification of a YFP-chloramphenicol resistance cassette was performed using ~40 base primers with homology to the insertion site (SPA) and the kanamycin resistance cassette. The deletion mutants or YFP-fusion strains were grown exponentially in LB medium at 32 °C. Prior to imaging, the cultures ($A_{600}=0.5$) were incubated for 120 min with MMC (0.15 μ M), pelleted, and 1.5 μ l of suspension was spotted onto a glass slide for image analysis. The images were captured using the Quorum WaveFX Spinning Disc Confocal System, and the cell length was measured using the Volocity program (Improvision Ltd) as previously described (Sydorsky *et al.*, 2010).

Coordinates

The atomic coordinates and structure factors have been deposited in the Protein Data Bank (<http://www.rcsb.org>) with accession codes 3NKD and 3NKE for the structures of YgbT and its C-terminal domain, respectively.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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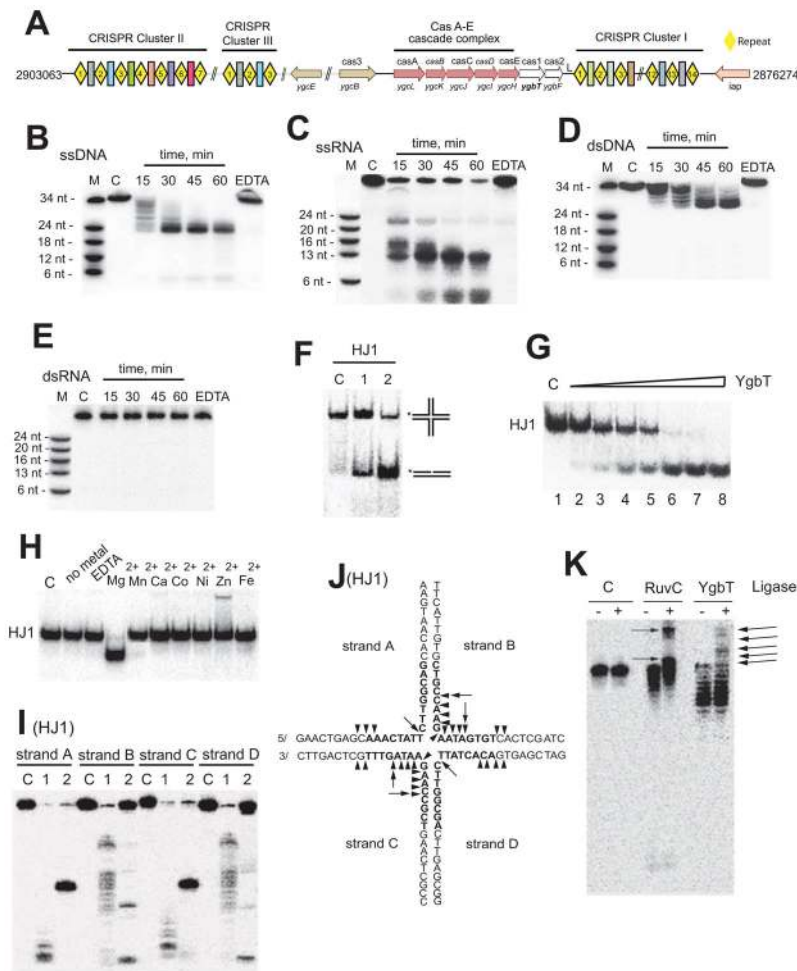


Fig. 1. *E. coli* CRISPR system and nuclease activity of YgbT. (A) CRISPR-Cas system of *E. coli* K12 W3110. Repeats are shown as yellow diamonds, and spacers as colored rectangular boxes. (B, C, D, E) Cleavage of ssDNA (B), ssRNA (C), dsDNA (D) or dsRNA (E) by YgbT (denaturing PAGE/autoradiography). The 5'-[³²P]-labeled ssDNA (34 nt), ssRNA (37 nt) or in a complex with complementary oligonucleotides was incubated without protein (C), with YgbT for the indicated times (37 °C) or in the presence of 10 mM EDTA for 60 min (EDTA). (F) Cleavage of HJ1 by YgbT (native PAGE/autoradiography). The 5'-[³²P]-labeled (*) HJ1 substrate was incubated at 37 °C for 45 min without protein (C) or with YgbT (1) or RuvC (2). (G) Dependence of HJ1 cleavage on YgbT concentration (native PAGE). The 5'-[³²P]-labeled HJ1 substrate (20 nM) was incubated in the absence (1) or in the presence of YgbT (Lanes 2–8: 0.05, 0.1, 0.2, 0.4, 0.8, or 1.0 μg) at 37 °C for 30 min. (H) Effect of divalent metal cations on HJ1 cleavage by YgbT (30 min, 37 °C, 5 mM metals or 10 mM EDTA; native PAGE). (I, J) Denaturing gel analysis of the cleavage of HJ1 by YgbT (triangles) or RuvC (arrows). The HJ1 substrate labeled by [³²P] on different strands (A, B, C, or D) was incubated without protein (C) or with YgbT (1) or RuvC (2). (K) Religation assay of the HJ cleavage products generated by YgbT or RuvC. The [³²P]-labeled asymmetric HJ2 was incubated (30 min at 37°C) without protein (C) or with RuvC or YgbT, and the reaction products were then incubated with T4 DNA ligase prior to PAGE. Religation products are indicated by arrows.

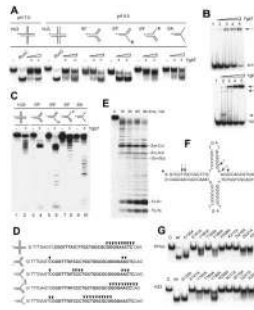


Fig. 2. Nuclease activity of YgbT against branched DNA substrates containing sequences of the *E. coli* CRISPR. **(A)** Cleavage of branched DNA substrates by YgbT (native PAGE). The substrates including static HJ (HJ3), replication fork (RF), 5'-flap (5/F), 3'-flap (3/F), and splayed arm duplex (SA) were 5'-[³²P]-labeled on the indicated strand (*) and incubated (30 min at 37 oC) in the absence or in the presence of 0.2 or 0.4 μg of YgbT at pH 7.0 or pH 8.5. **(B)** Gel-shift assay showing the binding of YgbT to 5'-flap or HJ3 (native PAGE). The 5'-[³²P]-labeled substrates were pre-incubated for 15 min at 25 oC without YgbT (1) or with YgbT (Lanes 2–5: 0.2 μg, 0.4 μg, 0.6 μg, and 0.8 μg). Arrows indicate the position of the DNA-YgbT complexes. **(C, D)** Denaturing PAGE analysis of YgbT cleavage sites. The 5'-[³²P]-labeled (*) substrates were incubated without or with YgbT (30 min, 37 oC). Panel D shows the sequences of the labeled strands and major cleavage sites of YgbT (arrowheads). The sequence of the *E. coli* CRISPR repeat-1 is shown in bold letters. **(E, F)** Cleavage of the CRISPR cruciform-like substrate CF1 by YgbT (denaturing PAGE analysis). The 5'-labeled CF1 (shown in panel F) was incubated with YgbT (0.2 μg) at 37 oC. **(G)** Alanine replacement mutagenesis of YgbT: cleavage of 5'-flap and HJ3 substrates by purified mutant proteins (30 min at 37 oC; native PAGE/autoradiography).

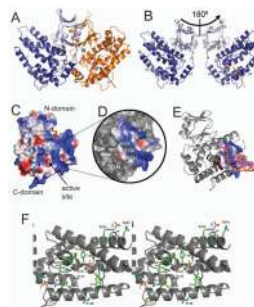


Fig. 3. Crystal structure of YgbT and the potential active site. **(A)** Overall structure of the YgbT dimer. **(B)** Two views of the YgbT monomer (related by a 180° rotation) showing the presence of two domains: the N-terminal β -sandwich-like domain (light-blue) and the C-terminal all- α domain (dark-blue). The position of the potential active site is indicated by the side chains of three conserved residues (shown as sticks). **(C)** Surface charge presentation of the YgbT monomer showing the presence of several large basic patches representing potential DNA-binding sites (colored in blue). **(D)** Close-up view of the main basic patch of YgbT located close to the potential active site (E141, H208, and D221) and showing the position of several conserved residues. **(E)** Superposition of the ssDNA fragment from the *E. coli* topoisomerase III DNA-binding site (1i7d) onto the potential DNA-binding site of YgbT. The DNA docking was performed using the Surface Screen analysis (Binkowski and Joachimiak, 2008). **(F)** Close-up stereo view of the YgbT active site from the structure of the C-terminal domain showing the position of three bound sulfate molecules shown as sticks and labelled (1, 2, and 3). The YgbT residues are shown as green sticks along a YgbT ribbon (grey). Two arrows in A and B indicate the protein side and position of the active site shown in C.

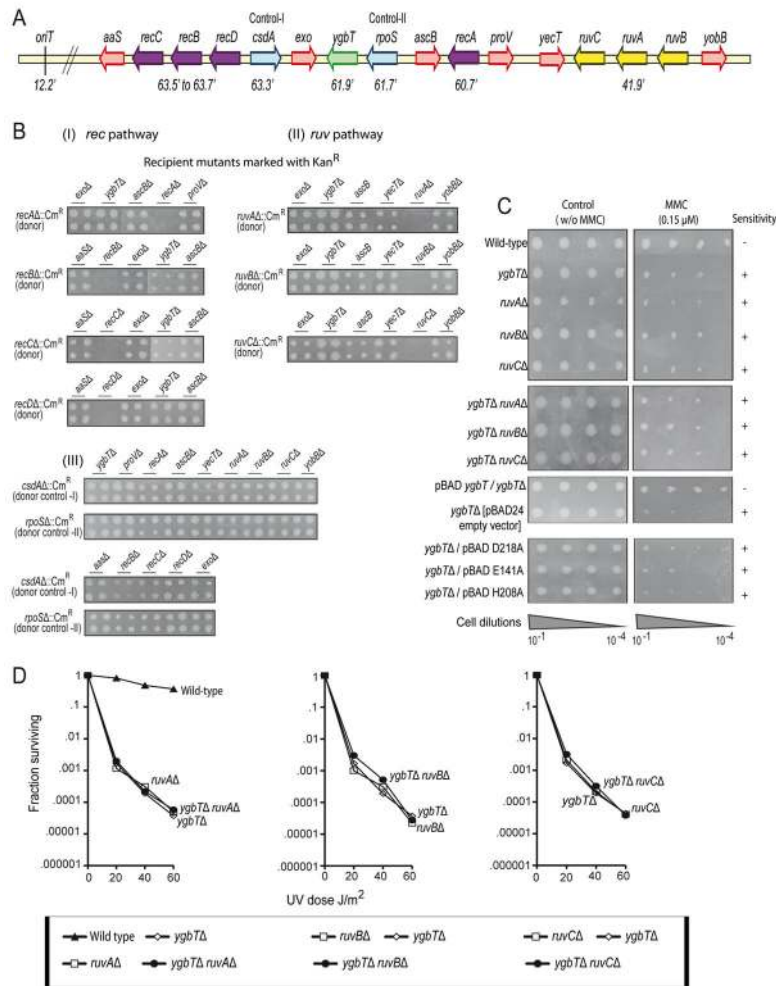


Fig. 5. Genetic interactions of *ygbT* and DNA damage sensitivity of the *ygbT* deletion strain. **(A)** Chromosomal positions of *ygbT* (green arrow), *rec* (purple arrows), *ruv* (yellow arrows), and other gene (red arrows) deletions used as donors (marked with *Cm^R*) or recipients (marked with *Kan^R*) in eSGA experiments. *csdA* and *rpoS* donor deletions (blue arrows) located close to the *rec* and *ygbT* genes served as controls for effects of proximity on recombination. The numbers indicate the gene coordinates on the *E. coli* chromosome (in minutes). *OriT* is the F origin of transfer in Hfr Cavalli (12.2 minutes). **(B)** Validation of *ygbT-rec* and *ygbT-ruv* synthetic genetic interactions. *rec* (Panel I), *ruv* (Panel II), and control (*csdA* and *rpoS*, Panel III) Hfr Cavalli donor deletion strains were crossed with the indicated *F⁻* recipient deletion strains (deleted for *ruv*, *ygbT*, or various genes flanking *ruv* and *ygbT*), followed by selection on plates containing chloramphenicol and kanamycin. **(C)** Serial cell dilution assay showing growth inhibition of *ygbT-ruvABC* double mutants and their single mutants by MMC (0.15 μM). Growth is classified as sensitive to MMC (+) or not sensitive compared with wild-type (-). Also shown are the effects of complementing $\Delta ygbT$ with pBAD vectors expressing wild-type or inactive mutant YgbT. **(D)** Effects of the indicated doses of UV irradiation on the survival of *ygbT-ruvABC* double mutants and their single mutants. The values plotted are averages from three independent experiments.

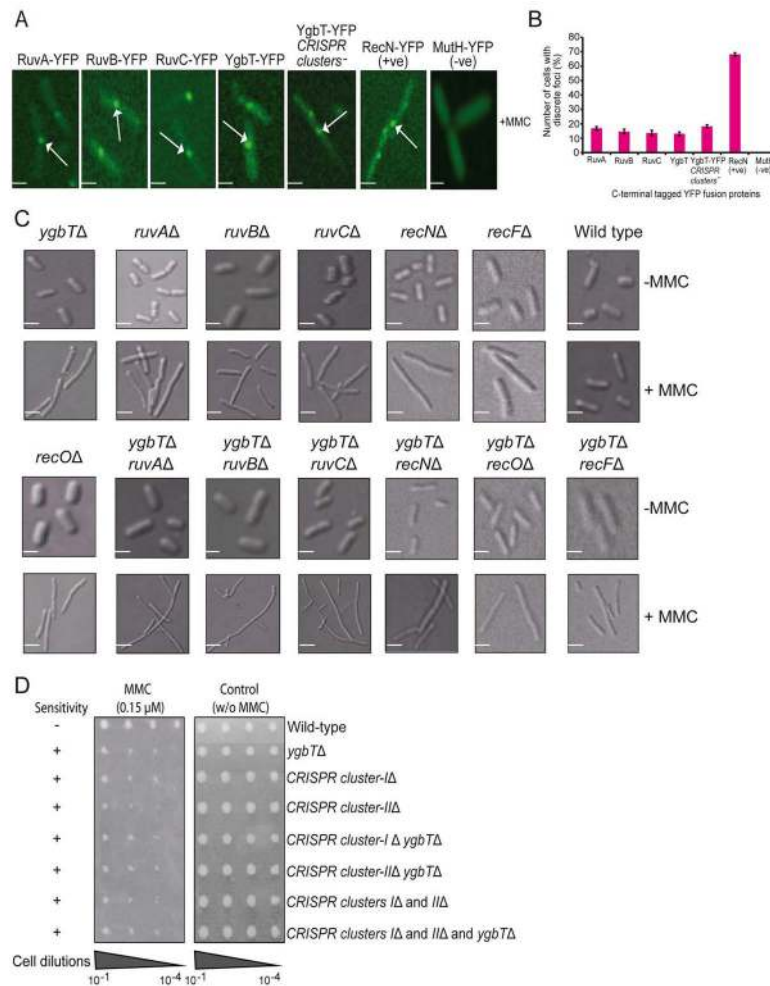


Fig. 6. Cell morphology of the *ygbT* deletion strain and intracellular localization of YgbT in response to DNA damage. **(A)** Intracellular localization of RuvA-YFP, RuvB-YFP, RuvC-YFP, YgbT-YFP, RecN-YFP and MutH-YFP fusion proteins in exponentially growing *E. coli* cells treated with MMC for 120 min. Scale bar, 2 μm. **(B)** The percentage of *E. coli* cells with discrete foci of various YFP fusion proteins in their nucleoids after MMC treatment. In each case, 250 different cells from five independent biological replicates (50 cells per replicate) were examined. Error bars indicate standard deviation from mean. **(C)** Differential interference contrast images of MMC-induced changes of cell morphology in *ygbT-ruvABC*, *ygbT-recN*, *ygbT-recO*, and *ygbT-recF* double mutants and their respective single mutants in the presence (+) and in the absence (-) of MMC. Scale bar, 2 μm. The average cell length of these cells is shown in Suppl. Fig. 4. **(D)** *E. coli* strains with deleted CRISPR clusters exhibit increased sensitivity to MMC. Serial cell dilution assays showing the effects of MMC (0.15 μM) on the growth of *E. coli* strains with deletions of one or both CRISPR clusters and/or *ygbT*. Growth is classified as sensitive to MMC (+) or not sensitive (-) in comparison to wild-type.