Two cDNAs from the plant Arabidopsis thaliana that partially restore recombination proficiency and DNAdamage resistance to *E.coli* mutants lacking recombination-intermediate-resolution activities

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ABSTRACT

Escherichia coil ruvC recG mutants lack RuvC endonuclease, which resolves crossed-strand joint molecules (Holliday junctions) formed during homologous recombination into recombinant products, and an activity (RecG) thought to partially replace RuvC. They are therefore highly deficient in homologous recombination, and sensitive to UV light and chemical DNA-damaging agents, presumably because of inability to tolerate unrepaired DNA damage by recombinational mechanisms (Lloyd, R.G. (1991) J. Bacteriol. 173:5414-5418). We transformed these mutants with plasmids expressing cDNAs from the plant Arabidopsis thaliana. Selection for bacteria with increased resistance to methylmethanesulfonate yielded two cDNAs, designated DRT111 and DRT112 (DNA-damage-repair/toleration). Expression of these plant cDNAs, especially DRT111, restored conjugal recombination proficiencies in ruvC and ruvC recG mutants to nearly wild-type levels. Both plant cDNAs significantly increased resistance of both mutants to UV light and several chemical DNA-damaging agents, but did not fully correct the mutant phenotypes. Drt111 activity, but not Drt112, also increased, to nearly wildtype levels, resistance of recG single mutants to UV plus mitomycin C. The predicted Drt111 and Drt112 polypeptides, 383 and 167 amino acids respectively, show no similarity with one another or with prokaryotic Holliday resolvases. Both appear chloroplast targeted; Drt112 is highly homologous to Arabidopsis plastocyanin. DRT111 and DRT112 probes hybridize only to DNA from closely related plants.

INTRODUCTION

Projected depletion of the stratospheric ozone layer is expected to significantly increase terrestrial UV-B irradiation at DNAdamaging wavelengths (1,2). This has heightened interest in the mechanisms by which green plants, which will necessarily be exposed continually to increased UV fluxes, resist DNAdamaging agents. Studies with yeast and, especially with the bacterium E. coli, have demonstrated that removal of photoproducts and recombinational toleration of unrepaired DNA lesions are both important resistance mechanisms (3). Excision repair and photoreactivation of UV photoproducts have been described for several plant species, including the model green plant Arabidopsis thaliana (4). However, there has been no strong evidence for homologous-recombination-dependent toleration processes, such as daughter-strand- gap filling (5).

The E.coli RecA protein mediates homologous pairing and strand exchange during recombination, yielding a crossed-strand intermediate (Holliday junction). Mutants lacking this activity are highly sensitive to DNA damage, as well as recombinationdeficient. Recently, we isolated four Arabidopsis cDNAs that partially complemented the UV-sensitivity phenotypes of E. coli mutants lacking all repair and toleration responses (Pang, Q., Hays, J.B., Rajagopal, I. and Schaefer, T.S., manuscript submitted). One of these, DRT100 (DNA-damage-repairtoleration) proved to partially complement RecA⁻ DNAdamage-sensitivity and recombination-deficiency (Rec-) phenotypes (6). The size of the predicted DrtlOO protein was similar to that of bacterial RecA proteins, but there was little global homology. Simultaneously, Jagendorf and coworkers (7) isolated an Arabidopsis cDNA with considerable recA homology by a hybridization approach, but did not test it for activity in E.coli. Both DrtlOO, the putative RecA analog, and the Arabidopsis RecA homolog appear to be chloroplast-targeted proteins. The existence of these genes argues strongly for the importance of DNA-damage-toleration processes in plants, at least for chloroplast genomes.

Activities that resolve crossed-strand intermediates into recombinant products have been demonstrated in E. coli phages T4 (8) and T7 (9), in *E. coli* itself (10), and in yeast $(11-13)$. The E. coli resolvase active in extracts has been identified as the product of the $ruvC$ gene (14), purified to homogeneity, and

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characterized biochemically (15). Surprisingly, $ruvC$ mutants are only slightly Rec-, but are highly sensitive to UV light and other DNA-damaging agents (16); double mutants, ruvC recG, are highly Rec⁻, and even more DNA-damage-sensitive. This suggests that recG encodes (or controls) a resolvase-like activity, not readily detectable in crude extracts, that substitutes fairly well for RuvC in conjugal and transductional recombination, but not in DNA-damage toleration. Further evidence for the more demanding nature of the latter process is its requirement for the RuvA and RuvB proteins as well (16).

The lack of amino-acid conservation among the phage and bacterial resolvases (17) suggested that hybridization approaches were not likely to yield the corresponding plant genes. Instead, we have selected for Arabidopsis cDNAs that apparently complement E. coli RuvC⁻ RecG⁻ phenotypes. The two cDNAs isolated in this way restore recombination nearly to wild-type levels, and increase resistance to DNA-damaging agents.

MATERIALS AND METHODS

Bacteria and bacteriophages

All strains used are derivatives of *Escherichia coli* K-12. Strain AB1157 is wild-type with respect to the markers of interest here (RuvC⁺ Rec⁺ phenotype) and is also F⁻ thi-1 his-4 Δ (gpt-proA) $argE3$ thr-1 leuB6 kdgK1 rfbD(?) ara-14 lacY1 galK2 xyl-5 mtl-1 $txx-33$ supE44 $rpsL31$ (18). Isogenic with AB1157, except as indicated, are the RuvC⁻ strain CS85, $ruvC53$ eda-51::Tn10, and the RuvC⁻ RecG⁻ strain N3398, recG258::Tn10 mini-kan ruvC53 eda-51 (19), and the RecG⁻ strain N2731 (20), recG 258::Tn10 mini-kan. Strain EG333 is HrfC pyrA::Tn10 mstB cysG303 $\Delta (lac-pro)$ XIII (21). The vector phage λ YES is cl857; it incorporates a cloning/expression site and yeast-E. coli shuttlevector-plasmid elements between two phage P1 lox sites (25). Phage λ CRE cI(ind⁻) red3 xis1 overexpresses the P1 Cre protein via a bacterial lac promoter (6,25). Neither λ YES nor λ CRE cI(ind⁻) prophages are inducible by DNA-damaging agents.

Plasmids

Plasmid pSE936, the product excised from λ YES by Cre-lox recombination, encodes ampicillin-resistance and plasmid ori elements for selection and propagation in E. coli, as well as URA3 and other elements for function as a yeast plasmid (25). Depending on their orientation, cDNAs inserted at the unique XhoI site are transcribable via the bacterial p_{lac} or yeast p_{GAL1} promoters. In plasmids pQP1110 and pQP1120, Arabidopsis cDNAs DRT111 and DRT112 are transcribed via p_{lac} . Plasmids pQP1112 and pQP1122, in which DRT111 and DRT112 are inverted with respect to p_{lac} , were constructed by digestion of plasmids pQP1110 and pQP1120 with XhoI endonuclease and re-ligation of the products, and were identified by restriction analysis of plasmids from transformed bacteria.

Media and solutions

TBY-broth, LB-broth-plates, and M9-minimal-plates have been described (26,27). TBY-Ap broth and LB-Ap plates contain 50 μ g/ml ampicillin (Ap).

Selection and isolation of DRT111 and DRT112 cDNA

We infected about 10^{10} RuvC⁻ RecG⁻ bacteria (strain N3398), lysogenic for λ CRE *cI*(ind⁻) red3 xis1, with an aliquot (5 × 10¹⁰) plaque-forming units) of an Arabidopsis cDNA library in the

vector XYES. This library had been obtained from R. Davis, Stanford University (25) and amplified once, as described previously (6). We grew the infected cells for one hour in TBY broth, at which point there were 5×10^9 total Ap-resistant bacteria (as determined by plating ^a small aliquot). We selected for cells containing excised plasmids, by growth for three hours in 100 ml TBY-Ap broth, plus 2 mM isopropylthio- β -Dgalactopyranoside (IPTG), yielding about 10^{11} bacteria. These were harvested by centrifugation and resuspended in 10 ml TBY. We spread the entire culture on twenty LB plates containing 0.06% methylmethane-sulfonate (MMS) and IPTG, and incubated them 40 h at 30 $^{\circ}$ C. Although none of 10^{11} bacteria in a parallel N3398(pSE936) culture survived on these plates, the cDNA-library-containing culture yielded 25 survivors. These were streaked across LB plates and tested for resistance to 5, 10, 15, and ²⁰ J/m2 of UV light. Four isolates were UVresistant, and plasmids extracted from each of these conferred resistance upon naive RuvC- RecG- bacteria. When digested with EcoRI endonuclease, one active plasmid released insert fragments of about 1.1 kb and 0.3 kb; we designated the cDNA as DRTJJI. The other three plasmids released 0.8-kb inserts; based on their apparently identical sizes, and complementation phenotypes in preliminary experiments (data not shown), we designated all three cDNAs as DRT112, and arbitrarily picked one for further study. (Their identity was subsequently confirmed by DNA sequence determinations.) We designated the respective plasmids pQP1110 and pQP1120.

Measurement of bacterial resistance to DNA-damaging agents

Cells were grown to late log phase in TBY-Ap broth containing IPTG, harvested by centrifugation, and resuspended to 1.5×10^8 colony-forming units (CFU) per ml, as described (6). Cell suspensions were treated with 254-nm UV light at ^a rate of ¹ w/m^2 and spread on LB-Ap plates, or spread on LB-Ap plates containing mitomycin C or methylmethanesulfonate (MMS) or 4-nitroquinoline-N-oxide (NQO). All manipulations were performed under room lighting, so that all cells were phenotypically Phr+. Plates were incubated overnight at 30°C.

Measurement of conjugal recombinant frequencies

Procedures were essentially as described by Miller (27). Overnight cultures of the donor strain (EG333), grown in TBY broth containing tetracycline (12.5 μ g/ml), and of recipient strains harboring various plasmids, grown in TBY-Ap broth with or without 2mM IPTG, were subcultured in fresh broth, grown to about 2×10^9 cells per ml, mixed at a ratio of three donors to one recipient, and incubated at 37'C. After ¹ hr we stirred mixtures vigorously, harvested the cells by centrifugation, and resuspended them in one volume of 0.O1M MgSO4. After 30 min at room temperature, cells were spread on LB-Ap plates and incubated overnight at 37°C, to score total recipient colonyforming units, or spread on M9-glucose-minimal plates containing ampicillin (50 μ g/ml), histidine (0.5 mM), arginine (0.06 mM), and proline (2 mM), to score Ap-resistant Leu+ Thr+ recombinants, or on M9-galactose-(casamino acids) plates containing ampicillin to score Ap-resistant Gal⁺ recombinants, or on M9-glucose-minirnal plates containing arginine (0.06 mM), histidine (0.5 mM), proline (2 mM), ampicillin and 13 μ g per ml tetracycline (Tc) to score Ap-resistant (Leu+ Thr+) recombinants.

Figure 1. Resistance to DNA-damaging agents of bacteria expressing Drt111 and Drt112. Indicated bacteria were (A) irradiated with 254-nm UV light to fluences indicated, or (B) spread on plates containing indicated mitomycin C concentrations, and surviving bacteria scored, as described under 'Materials and Methods'. Surviving fraction equals CFU surviving divided by CFU on LB-Ap plates (no UV). Strains [phenotypes] employed: (×), AB1157 (pSE396) = [Ruv+ Rec+]; (0), CS85 (pSE396) [= RuvC⁻ Rec⁺]; (\triangle), CS85 (pQP1110) = [RuvC⁻ Rec⁺(Drt111)]; (\square), CS85 (pQP1120) = [RuvC⁻ Rec⁺(Drt112)]; (\bullet), N3398 (pSE396) = [RuvC- RecG-]; (A), N3398 (pQP1llO) = [RuvC- RecG-(Drtlll)]; (U), N3398 (pQP1120) = [RuvC- RecG-(Drtll2)]. Data correspond to averages for two trials. Standard deviations were generally less than symbol sizes and were almost always 10% of values shown or less.

Determination and analysis of DNA sequences

The Oregon State University Central Services Laboratory determined DNA sequences on double-stranded DNA by an automated technique, using an Applied Biosystems Model 373A DNA sequencer and ^a Taq Dye Primer Cycle Sequencing Kit, with dideoxy chain termination, thermal cycling and primercoupled dyes, according to instructions supplied by the manufacturer (bulletin No. 237605). M13mplac Universal and Reverse primers (United States Biochemical Corp.) and internal primers were employed. For each sample, sequences were determined in both directions at least once each. In some regions we also determined DNA sequences manually, by dideoxy sequencing of duplex DNA using ^a Sequenase kit (United States Biochemical Corp). We compared the predicted amino-acid sequences for Drt111 and Drt112 to protein sequences in the SWISSPROT data base, using Intelligenetics Suite release 5.4 programs QUEST, PEP, SEQ, and GENALIGN. The last program, developed by Dr. H. Martinez, is a copyrighted product of Intelligenetics, Inc. We searched GENBANK release 69, and NBRFIPIR and EMBL protein-sequence libraries using the Intelligenetics FASTDB search program.

Hybridization analyses

DNA was extracted from Arabidopsis, broccoli and cabbage (Brassica pekinensis) tissues as described (4). DNA from bean (Phaseolus vulgaris) and maize (Zea mays) were gifts respectively of David Mok and Carol Rivin, Oregon State University. DNA

digested with EcoRI endonuclease was analyzed by electrophoresis, blotting, hybridization with a [32P]-labeled [random-primer-method (28)] EcoRI fragment isolated from plasmids QP1111 and pQP1121, and autoradiography, as described (29), with minor modifications. Hybridization, at 37°C, employed solutions containing 50% formamide and 0.6 M salt. Reduced-stringency hybridization employed 32% formamide. Aqueous washes (0.03 M salt) were at 37°C; filters were autoradiographed for 48 hr.

RESULTS

Selection and isolation of cDNAs

E. coli ruvC recG bacteria lack activities that resolve recombination intermediates (Holliday junctions), and therefore are deficient in homologous recombination (Rec⁻) and DNAdamage-toleration functions (19). We established an Arabidopsis cDNA plasmid expression library in RuvC- RecG- bacteria harboring a λ Cre prophage, by infecting 10¹⁰ cells of strain N3398 with a phage λ YES Arabidopsis cDNA library, at a multiplicity of 5 phage per cell. The endogenous Cre activity expressed by the λ Cre prophage (25) excised plasmids from the λ YES phage via site-specific recombination at the lox sites in XYES; plasmids were established in about 50% of the bacteria.

We amplified cDNA-plasmid-containing bacteria and plated them in the presence of 0.06% methylmethanesulfonate, a concentration which killed all non-cDNA-containing bacteria, and

Table I. Effects of Drt111 and Drt112 on resistance of E.coli mutants to UV light

Relevant bacterial (plasmid) phenotype ^a	Relative survival (%) after indicated UV fluences $(J/m^2)^b$			
	20	40	60	
Ruv^+ Rec^+ (none)	(100)	(100)	(100)	
Ruv^+ RecA ^{$-$} (none)	0.3	0.05	0.05	
Ruv^+ RecA ⁻ (Drt111)	0.4	0.05	0.05	
Ruv^+ RecA ⁻ (Drt112)	0.4	0.06	0.05	
Ruv^{+} RecB ⁻ C ⁻ F ⁻ (none)	0.2	0.08	0.05	
Ruv^+ RecB ⁻ C ⁻ F ⁻ (Drt111)	0.2	0.08	0.08	
Ruv^+ RecB ⁻ C ⁻ F ⁻ (Drt112)	0.3	0.07	0.06	
$RuvC^-$ Rec G^- (none)	0.1	0.01	0.03	
$RuvC^-$ Rec G^- (Drt111)	2.7	0.16	0.67	
$RuvC^-$ Rec G^- (Drt112)	1.4	0.06	0.27	

^aRespective bacterial strains (plasmids) employed, lines 1 through 10, were ABI 157 (pSE936), QP3070(pSE936), QP3070(pQP1 110), QP3070(pQP1 120), JH312(pSE936), JH312(pQP1110), JH312 (pQP1120), N3398 (pSE936), N3398 (pQP1120), N3398 (pQP1120).

bFractions of bacterial suspensions surviving indicated UV doses were measured as described under 'Materials and Methods,' and divided by surviving fractions for Ruv⁺ Rec⁺ (none) bacteria. These latter values (relative survival of 100%) were 0.32 at 20 J/m², 0.043 at 40 J/m², and 0.0027 at 60 J/m². Data correspond to averages for two plates (range typically \pm 10%).

we tested the 25 survivors for resistance to $10-30$ J/m² UV light. All four UV-resistant isolates harbored plasmids containing putative cDNA inserts-one of 1.4 kb, three of 0.8 kb. On the basis of the apparent identity of size and correction activity (see below) of the latter three cDNAs, we identified two unique DNAdamage-repair/toleration cDNAs, DRT111 and DRT112. We tested these for their effect on various DNA-damage-sensitivity and recombination-deficiency phenotypes of E. coli ruvC, ruvC recG, and recG mutants.

Partial correction of DNA-damage-sensitivity phenotypes by Drtlll and Drtll2

We measured the effect of Drt111 and Drt112 activity on the resistances, to ultraviolet light (Fig. IA) and to the DNAcrosslinking agent mitomycin C (Fig. 1B), of $RuvC^-$ single and RuvC⁻ RecG⁻ double mutants. Drt111 and Drt112 increased the resistance to UV light of both RuvC⁻ single mutants (Fig. 1A, open symbols) and $RuvC^-$ Rec G^- double mutants (filled symbols). Although the factors by which survival was increased were greater for the double mutants, apparent correction efficiencies relative to wild-type resistance ranged from 0.1 % to 1%. Resistance of $RuvC^-$ and $RuvC^-$ Rec G^- bacteria to mitomycin C (Fig. 1B; open, filled symbols) was also increased by the presence of Drtl 11 and Drt 112. Here resistances were increased by as much as 400-fold $[RuvC - RecG - (Drt111)$ at 0.3 μ g/ml], and apparent correction efficiencies were as high as 10 to 40% (at 0.1 μ g/ml). The two plant cDNAs also increased resistance to methylmethanesulfonate, at concentrations of 0.015 to 0.045%, by factors of about 2-fold for RuvC⁻ and up to 20-fold for RuvC- RecG- mutants, corresponding to complementation efficiencies of 0.1% or less; resistance of RuvC⁻ RecG⁻ mutants to 10 μ M nitroquinoline oxide was increased 7-fold by Drtlll and 3-fold by Drti12 (data not shown).

Drtl 11 and Drtl 12 might partially correct the DNA-damagesensitive phenotypes of $RuvC^ RecG^-$ bacteria by resolving intermediates generated by normal (RecA-dependent) E.coli recombinational toleration processes, or suppress the phenotypes, tested the effects of $Drt111$ and $Drt112$ on a bacterial

Table II. Resistance of RecG⁻ mutants expressing Drt111 or Drt112 to UV light plus mitomycin C.

Relevant bacterial	Survival (%) of bacteria treated as indicated	
(plasmid) phenotype ^a	UV only	UV plus mitomycin C
Rec^+ (none)	15 ± 2	$13 + 0.5$
$RecG^-$ (none)	1.7 ± 0.3	0.26 ± 0.13
$RecG^{-} (Drt111)$	8.5 ± 0.7	6.1 ± 0.7
$RecG^-$ (Drt112)	$2.7 + 0.2$	0.45 ± 0.06

^aRespective strains (plasmids) employed, lines 1 through 4, were AB1157 (pSE936), N2731 (pSE936), N2731 (pQP11lO), N2731 (pQP1120). bBacteria were grown in broth containing ampicillin and IPTG, treated with 30 J per m² UV light, as described under 'Materials and Methods,' spread on LB plates with or without $0.2 \mu g$ per ml mitomycin C (20), and incubated overnight. Data represent averages for two trials (two plates per trial), with ranges indicated.

Table III. Conjugal recombinant frequencies

Relevant bacterial (plasmid) phenotype of recipients ^a	Number of recipients $(CFU/ml \times 10^{-7})^b$	Number of Thr^+ Leu ⁺ Ap ^r recombinant $(CFU/ml \times 10^{-7})^b$	Relative recombinant frequency ^c
Rec^+ Ruv ⁺ (none) $Rec^+ RuvC^-$ (none) $Rec^+ RuvC^- (Drt111) 21 \pm 2$ Rec^+ RuvC ⁻ (Drt112) $RecG^- RuvC^-$ (none) $RecG^{-}$ RuvC ⁻ (Drt111) 38 \pm 6 $RecG^{-}$ RuvC ⁻ (Drt112) 44 \pm 8	35 ± 4 32 ± 4 24 ± 2 37 ± 2	5.6 ± 0.6 1.0 ± 0.4 2.5 ± 0.2 $1.7 + 0.2$ 0.019 ± 0.004 0.67 ± 0.02 0.41 ± 0.7	(1.0) 0.21 0.75 0.44 0.003 0.110 0.058

^a Respective recipient bacterial strains (plasmids) employed lines ¹ to 7, were: 1), AB1157 (pSE936), CS85 (pSE936), CS85 (pQPIIIO), CS85 (pQP1120), N3398 (pSE936), N3398 (pQP1110), N3398 (pQP1120).

^b Conjugal matings and measurements of total recipients and Ap-resistant Thr⁺ Leu⁺ transconjugants were performed as described under 'Materials and Methods'. Hfr donor was strain EG333. Data are averages and standard deviations for two trials.

^c Relative recombinant frequency equals ratio of recombinant frequency to recipient frequency for indicated bacteria, divided by ratio for Rec⁺ Ruv⁻ bacteria.

by mediating new repair or recombinational toleration pathways that did not require RuvC or RecG function, for example. We tested the ability of plasmids expressing Drt111 or Drt112 to increase the resistance of other Rec⁻ mutants to UV light (Table I). Under conditions where Drt 111 and Drt112 significantly increased survival of RuvC⁻ RecG⁻ bacteria (Table I, lines $8-10$), there was no effect on survival of RecA⁻ (lines $2-4$) or RecB⁻C⁻F⁻ (lines 5-7) bacteria.

We also tested for correction of Rec G^- single mutations, using the UV-plus-mitomycin-C assay of Lloyd and Buckman (20) (Table II). $RecG^-$ (Drt111) bacteria were about half as resistant as wild-type, but RecG⁻ (Drt112) bacteria were not significantly more resistant than RecG⁻.

Correction of recombination deficiencies

Although Drt111 and Drt112 both significantly increased the resistance of both $RuvC^-$ and $RuvC^ RecG^-$ mutants to a variety of DNA-damaging agents, the apparent correction efficiencies were only several percent or less, in all but a few cases. Furthermore, these data provide no direct evidence that recombination-enhancing activities are involved. Therefore, we

C. DRT112 (low stringency)

Figure 2. Hybridization of DRTIII and DRTII2 cDNAs to plant DNA. Lanes contain DNA from Arabidopsis leaves and stems (5 μ g), broccoli florets (10 μ g), Chinese cabbage (Brassica pekinensis) leaves (10 μ g), bean (Phaseolus vulgaris) seedlings (10 μ g), and whole maize (Zea mays) plants (10 μ g). Extraction of DNA, digestion with EcoRI endonuclease and hybridization were as described under 'Materials and Methods'. A. DRT111 probe. B. DRT112 probe. C. DRT112 probe at low stringency.

recombination process. Homologous recombination is needed for formation of stable transconjugants during mating of Hfr and Fbacteria. Although recA mutations drastically reduce conjugal recombination frequencies (30), a single $ruvC$ mutation in a wildtype background reduces recombination only 5-fold [ref. (19) and Table I, line 2], perhaps because recombination intermediates are resolved in other ways. Drt111 and Drt112 corrected this $RuvC^-$ phenotype, restoring recombination to nearly wild-type levels in the former case (Table HI, lines 3,4). The recombinant frequency in ruvC recG double mutants is drastically reduced, to just 0.3% of wild-type levels (Table III, line 5). Here Drt111 increased the frequency 32-fold, to 11% of wild-type levels (about half the restoration by RecG alone), and Drt112 was slightly less effective (Table III, lines $6,7$).

Neither Drt111 nor Drt112 corrected recombination deficiencies in other Rec⁻ mutants tested: The frequency of tetracycline-resistant transconjugants, in matings of the Hfr $pyrA::Tn10$ strain EG333 with RecA- and RecB-C-Frecipients, were reduced respectively to 0.01% and 1% of rec⁺ levels, in the presence or absence of DRTIll or DRT112 (data not shown); in parallel experiments Drt111 and Drt112 increased recombinant frequencies in $RuvC^ RecG^-$ bacteria from 0.17% of wild-type frequencies to 7.4% and 2.5%, respectively. Neither Drt111 nor Drt112 affected the efficiency of transfer of conjugal F' episomes in any bacteria tested [Rec+, RecA-, RecB-Fand RuvC⁻ RecG⁻ (data not shown)], i.e. the apparent increases in recombinant frequency are not due to increased mating efficiencies in the presence of Drt111 or Drt112.

To determine whether correction of the RuvC⁻ RecG⁻ recombination deficiency required expression of DRTI11 and DRT112, rather than being the result, for example, of induction of new E. coli activities by the presence of the plant DNA

Table IV. Effects of gene orientation and induction of expression on activity of DRTIII and DRTII2 in recombination-deficient bacteria

Bacterial (plasmid) phenotype of recipient ^a	Frequency $(\%)$ of Ap ^r Gal ⁺ transconjugants per recipient ^b		
	IPTG-induced	no IPTG	
Ruv^+ Rec ⁺ (none)	5.0	6.2	
$RuvC^-$ Rec G^- (none)	0.008	0.006	
$RuvC^ RecG^-$ (Drt111)	0.462	0.080	
$RuvC^ RecG^-$ (Drt111 INV)	0.008	0.008	
$RuvC^-$ Rec G^- (Drt112)	0.180	0.060	
$RuvC^ RecG^-$ (Drt112 INV)	0.008	0.007	

aRespective bacterial strains (plasmids) employed, lines 1 to 6, were AB1157 (pSE936), N3398 (pSE936), N3398 (pQP111O), N3398 (pQP1112), N3398 (pQP1120), N3398 (pQP1122).

^oConjugal matings, with EG333 as Hfr donor and scoring for Ap^r Gal⁺ recombinants, were performed as described under 'Materials and Methods.' Data correspond to average for two plates; range was less than $\pm 10\%$ in almost all cases.

sequences themselves, we measured requirements for transcription (Table IV). Neither DRT111 nor DRT112, when inverted with respect to the plasmid P_{lac} promoter, showed any correction activity (Table IV, lines 4, 6). In the absence of induction by IPTG, the activity of P_{lac} -transcribed DRT111 and DRT112 sequences was decreased but still significant (Table IV, lines 3, 5), presumably because multiple copies of the lac operator titrated out endogenous levels of even lacIq-expressed lac repressor (31).

DNA and protein sequence analyses

DNA and predicted protein sequences for DRT111 and DRT112 are available via GENBANK access numbers M98455 and

M98456, respectively. We used the Intelligenetics FAST DB program to search protein sequences in the GENBANK, NBRFIPIR, and EMBL libraries for amino-acid-sequence similarities. The DRT111 open reading frame predicts a 383-residue polypeptide without significant global similarity to any known protein, including the prokaryotic Holliday-junction resolvases-phage T4 gene 49 protein (32), phage T7 gene 3 protein (9), E.coli RuvC (17) and RecG proteins (16)—and the yeast cruciform-cutting enzyme (33). However, the Drt111 amino-acid sequence $Q_{225}GqGlGKS$ strongly resembles the RecG sequence QGdvGsGKT (16), and thus the nucleotide binding motif (Walker box) GxxGxGKS (34) characteristic of many ATP-hydrolyzing DNA repair and recombination proteins. Surprisingly, the predicted Drtl 12 polypeptide, 167 amino acids, is 75% identical with Arabidopsis plastocyanin (35). The homology extends over most of the sequences, except for nine plastocyanin residues between Drtl12 amino acids 24 and 27. Drt112 shows no significant homology with any other proteins in the data bank, or with other resolvases. The N-terminal portions of both polypeptides show features characteristic of chloroplast transit peptides (36): high frequency of serine and threonine residues, large numbers of small hydrophobic amino acids, net positive charge. We did not find ^a consensus chloroplast processing site $(36,37)$ in Drt111, but the plastocyanins (35) are cleaved at a sequence that ends in DrtI 12 at amino acid 68. The apparent chloroplast-targeting domain of DrtI 11 occupies about 90 N-terminal residues.

Hybridization of DRTIll and DRTJ12 probes to plant DNA

We hybridized, at high stringency, *DRT111* and *DRT112* probes to bulk DNA from Arabidopsis, the closely related Brassicaceae broccoli and Chinese cabbage, and from bean and the monocot plant maize (Fig. 2). The DRT111 probe hybridized strongly to DNA from Arabidopsis and very weakly to broccoli DNA, but not to DNA from other plants (Fig. 2A). At high stringency, the DRT112 probe yielded only ^a single strong DNA signal (Fig. 2B) which presumably corresponds to DRT112 itself. The additional two lower-molecular-weight bands that appear at lower stringency may correspond to the Arabidopsis plastocyanin gene (35), which encodes an EcoRI restriction site not in the DRT112 sequence. Vorst et al. (35) detected only a single (intron-less) plastocyanin gene by hybridization analysis. Thus, despite the high degree of homology, the nucleotide differences (about 30%) apparently prevent DRT112-plastocyanin hybridization at high stringency.

DISCUSSION

We have isolated two *Arabidopsis* cDNAs that appear to increase recombinant progeny in conjugal crosses involving E.coli mutants lacking ability to resolve intermediates (Holliday structures). These cDNAs were originally selected by virtue of their ability to promote survival of $ruvC$ recG mutants, on plates containing methylmethanesulfonate (MMS), but they proved to significantly increase resistance to other DNA-damaging agents as well. We have considered three explanations, other than genuine complementation-replacement of RuvC or RecG resolution activities-for these observations. Informational suppression can be ruled out: $recG258$, which is strongly corrected by DRT111, is an insertion mutation; the $ruvC53$ allele encodes a highly temperature-sensitive but full-length protein (R. Lloyd, personal communication). Second, the requirement for P_{lac} -initiated

transcription of DRTIll and DRT112 excludes the possibility that either DNA sequence in and of itself provokes ^a phenotypesuppressing response in $E. coli$. Third, the inability of Drt111 or Drtl 12 to promote conjugal recombination or UV-resistance in E. coli recA or recB recC recF mutants argues strongly against the notion that either activity mediates a novel recombination pathway that does not depend on resolution via RuvC or RecG activity.

Correction efficiencies, for DRTIll and DRT112 relative to wild-type cells, were high for conjugal recombination $(9-70\%$ and $4-25\%$, respectively), for resistance to lower levels of mitomycin C $(10-40\%$ and $3-10\%$ respectively), and for complementation by Drt111 of $RecG$ ⁻ DNA-damage sensitivity (50%). Efficiencies were less for higher degrees of damage. These trends may reflect saturation, by high amounts of DNA damage, of activities limited by lack of bacterial translation signals, poor codon usage, RNA instability or protein instability, or insolubility in *E. coli*, or the presence of activity-inhibiting chloroplast processing signals on the Drt111 and Drt112 polypeptides. In fact, in experiments in which proteins were radiolabeled in E. coli 'mini-cells', neither Drt111-encoding nor DrtI 12-encoding plasmids yielded detectable bands of the appropriate molecular weight (Q. Pang, unpublished results). The correction patterns thus suggest that activities able to efficiently resolve a few conjugal-recombination intermediates, despite low intrinsic biochemical proficiency and/or low levels of expression, may not be able to deal with large numbers of intermediates arising during DNA-damage-provoked sister-chromatid exchange.

Drt111 and Drt112 differ in the sequences and length of their polypeptides, their apparent efficiencies for complementation of RuvC- and RuvC- RecG- phenotypes, and in the ability of Drt111, but not Drt112, to efficiently complement a RecG⁻ phenotype. The molecular weight of Drt111, 42 kDa, falls between those of RecG, 76 kD (16) and RuvC and the phage T7 and T4 resolvases, $17-19$ kDa $(8-10)$. Drt112 is highly similar to Arabidopsis plastocyanin (35), a nuclear-encoded chloroplast protein that participates in electron transfer between photosystem I and the cytochrome b/f complex. If Drt112 is processed at the same site at which plastocyanin is thought to be cleaved in the chloroplast (35), the putative nature Drt112 protein (amino acids $69-167$), would have a molecular weight of only 11 kDa, significantly less than those of the RuvC/phage resolvases. The Drtl 12-plastocyanin similarity is reminiscent of the similarity in between respiratory-chain NADH dehydrogenase and ^a protein that binds to the chloroplast DNA replication origin (38).

E. coli appears to process recombination intermediates, such as Holliday structures, in two steps, by at least two pathways. The bacterial RuvA and RuvB proteins together recognize Holliday junctions and catalyze ATP-dependent branch migration (39,40,41); RecG alone efficiently accomplishes both these tasks (42). Both RuvAB and RecG activities resolve Holiday junctions in short linear model substrates, simply via branch migration out to the ends. However, resolution of such intermediates in chromosomes would require that the migrating junctions encounter preexisting DNA strand nicks, or be cleaved by resolving endonucleases. The RuvC protein is one such resolvase $(10,14)$. Since RuvC⁻ (and RuvA⁻ or RuvB⁻ mutants) are only slightly recombination-deficient (19), yet RecG does not appear to cleave Holliday junctions (42), there may another, as yet unidentified, E. coli resolvase that cleaves junctions recognized

by RecG. Preference of the RuvC resolvase for RuvAB-bound Holliday junctions has not been demonstrated (39,40,41). However, the indistinguishability of RuvA⁻, RuvB⁻, and $RuvC^-$ phenotypes (19) suggests that these proteins, which are encoded by three nearly contiguous genes (17), may act cooperatively in vivo. Ruv⁻ mutants are much more sensitive to DNA damage than RecG⁻ mutants, although both classes of mutants are only slightly recombination-deficient (19). This suggests that the Ruv and RecG pathways are nearly, but not perfectly, interchangeable for conjugal recombination, but that the Ruv proteins are designed to play the predominant role in recombinational toleration of DNA damage.

How might the Arabidopsis Drt activities identified here relate to these $E.$ coli functions? Drt111 is slightly more than half the size of RecG. However, both proteins exhibit a nucleotide binding site, recombination frequencies in RuvC-RecG+ and $RuvC^{-}RecG^{-}$ (Drt111) bacteria are similar (Table III), and Drt111 efficiently corrects RecG⁻ phenotypes, despite its apparently low level of expression. Although Drtl 11 may thus serve as a RecG analog, Drt112 does not correct RecG⁻ phenotypes. The pattern of correction, by apparently limited amounts of Drt112, of RuvC⁻ and RuvC⁻RecG⁻ phenotypes is not inconsistent with partial replacement of the RuvC endonuclease, but such an identification requires further study, in vitro as well as in vivo. The lack of extensive similarity among all proteins thus far implicated in resolution of recombination intermediates-Drtlll, Drti12, the E.coli Ruv and RecG proteins, and the phage T4 and T7 resolvases-suggests that various organisms have recruited a wide variety of proteins to mediate this process. The validity of these speculations remains to be tested, by biochemical studies with purified Drt111 and Drt112 proteins.

The two Arabidopsis RecA homologs/analogs described previously (6,7), and the apparent plant resolution proteins described here, incorporate putative chloroplast transit peptides. This suggests that recombinational toleration is an important feature of resistance of chloroplasts to DNA damage.

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