RESEARCH PAPER

RecA maintains the integrity of chloroplast DNA molecules in Arabidopsis

Beth A. Rowan†, Delene J. Oldenburg and Arnold J. Bendich*

Department of Biology, University of Washington, Seattle, WA 98195 USA

† Present address: Department of Molecular Biology, Max Planck Institute for Developmental Biology, D-72076 Tübingen, Germany.
* To whom correspondence should be addressed. E-mail: bendich@u.washington.edu

Received 28 January 2010; Revised 15 March 2010; Accepted 16 March 2010

Abstract

Although our understanding of mechanisms of DNA repair in bacteria and eukaryotic nuclei continues to improve, almost nothing is known about the DNA repair process in plant organelles, especially chloroplasts. Since the RecA protein functions in DNA repair for bacteria, an analogous function may exist for chloroplasts. The effects on chloroplast DNA (cpDNA) structure of two nuclear-encoded, chloroplast-targeted homologues of RecA in Arabidopsis were examined. A homozygous T-DNA insertion mutation in one of these genes (cpRecA) resulted in altered structural forms of cpDNA molecules and a reduced amount of cpDNA, while a similar mutation in the other gene (DRT100) had no effect. Double mutants exhibited a similar phenotype to cprecA single mutants. The cprecA mutants also exhibited an increased amount of single-stranded cpDNA, consistent with impaired RecA function. After four generations, the cprecA mutant plants showed signs of reduced chloroplast function: variegation and necrosis. Double-stranded breaks in cpDNA of wild-type plants caused by ciprofloxacin (an inhibitor of Escherichia coli gyrase, a type II topoisomerase) led to an alteration of cpDNA structure that was similar to that seen in cprecA mutants. It is concluded that the process by which damaged DNA is repaired in bacteria has been retained in their endosymbiotic descendent, the chloroplast.

Key words: DNA structure, fluorescence microscopy, homologous recombination, leaf variegation, pulsed-field gel electrophoresis.

Introduction

All organisms experience DNA damage, which must be repaired in order to avoid deleterious phenotypic consequences (Thompson and Schild, 2001; Tuteja et al., 2009). Many processes can be used to maintain the integrity of the genome: photoreactivation, base excision repair, nucleotide excision repair, mismatch repair, homologous recombination, and cross-link repair (Sancar and Sancar, 1988; Niederhofer, 2008). Mechanistic details are becoming increasingly clear in bacteria, eukaryotic nuclei, and the mitochondria of yeast and mammals (Bohr and Anson, 1999; Nowosielska, 2007; Aguilera and Gomez-Gonzalez, 2008). By comparison, however, DNA repair processes in the mitochondria and chloroplasts of plants remain poorly understood (Kimura and Sakaguchi, 2006; Vlek et al., 2008).

Homologous recombination is important for the repair of double-strand DNA breaks (DSB), DNA gaps, and interstrand cross-links, and occurs by invasion of a homologous DNA strand followed by strand exchange (Li and Heyer, 2008). The Escherichia coli RecA/RecBCD pathway of strand invasion and exchange is perhaps the best characterized system for DNA repair by homologous recombination (Cox, 2007; Michel et al., 2007). The RecA protein is highly conserved among diverse bacterial species, and the RecBCD complex has different functional analogues in different bacterial lineages (Rocha et al., 2005).

Abbreviations: DAPI, 4′,6-diamidino-2-phenylindole; cpDNA, chloroplast DNA; PFGE, pulsed-field gel electrophoresis; DSB, double-strand DNA breaks.
© 2010 The Author(s).
This is an Open Access article distributed under the terms of the Creative Commons Attribution Non-Commercial License (http://creativecommons.org/licenses/by-nc/2.5), which permits unrestricted non-commercial use, distribution, and reproduction in any medium, provided the original work is properly cited.
Eukaryotic homologues of RecA (RAD51) and organelle-targeted prokaryotic RecA homologues are encoded by the nuclear genomes of many organisms, including plants (Lin et al., 2006). Repair and recombination of chloroplast DNA (cpDNA) in *Chlamydomonas reinhardtii* is suppressed when a dominant-negative version of *E. coli* RecA is targeted to chloroplasts (Cerutti et al., 1995). The prevalence of chloroplast-targeted RecA mRNA increases in response to DNA damage in *C. reinhardtii* (Nakazato et al., 2003), *Physcomitrella patens* (Inouye et al., 2008), pea (Cerutti et al., 1993), and *Arabidopsis* (Cao et al., 1997). Mitochondrial-targeted RecA has been shown to be important for mitochondrial DNA (mtDNA) repair in *P. patens* (Odahara et al., 2007) and for mtDNA recombination and repair in *Arabidopsis* (Khazi et al., 2003; Shedge et al., 2007).

The nuclear genome of *Arabidopsis* encodes five putative homologues of RecA that are predicted to be localized in mitochondria and chloroplasts (Table 1). Locus AT1G79050 encodes a protein (RECA1; cpRecA) that has been shown experimentally to be localized in chloroplasts (Cao et al., 1997; Shedge et al., 2007) and has a 61% identity with cyanobacterial RecA (Cerutti et al., 1992). Locus AT3G12610 encodes a protein (DRT100) that has only weak homology to *E. coli* RecA and has a putative chloroplast-targeting signal peptide (Pang et al., 1992). DRT100 cDNA can, however, partially restore the growth phenotypes of recA mutants of *E. coli* (Pang et al., 1992, 1993). Locus AT2G19490 encodes a protein (RECA2) that localizes to both mitochondria and chloroplasts, and locus AT3G10140 (RECA3) encodes a protein that localizes to mitochondria only (Shedge et al., 2007). Locus AT3G32920 has been annotated as encoding a mitochondrial-targeted RecA protein, but the gene is probably a pseudogene (Shedge et al., 2007).

Previous reports describing the consequences of mutation in a chloroplast-targeted RecA homologue for any plant could not be found [other than lethality previously reported for RECA1(cprecA) (Shedge et al., 2007)]. In this study, the function of two of the three chloroplast-targeted RecA proteins in *Arabidopsis* is addressed using T-DNA insertion mutants and it is shown that the cpRecA gene encodes a protein that functions similarly to bacterial RecA. Pulsed-field gel electrophoresis (PFGE) and fluorescence microscopy of individual ethidium bromide-stained cpDNA molecules were used to analyse cpDNA structure. In wild-type (*Arabidopsis*), as in other plants, in-gel-prepared cpDNA consists of multigenomic, complex forms that do not migrate into the gel, discrete bands of monomeric and oligomeric linear molecules, and a smear of linear molecules. Circular forms are rarely found. Reduction of *cpRecA* mRNA leads to a decrease in the prominence of the bands, an increase in the smear, and leaf abnormalities that probably result from a lack of DNA damage repair in chloroplasts.

### Materials and methods

#### Plant growth conditions

Seeds of *Arabidopsis thaliana* (Columbia or SALK T-DNA insertion lines obtained from the Arabidopsis Biological Resource Center, http://www.biosci.ohio-state.edu/~plantbio/Facilities/abrc/abrchome.htm) were sown on soil (for the experiments shown in Figs 1–4) or on sterile agar plates containing MS salts (Murashige and Skoog, 1962), 1% sucrose, and 0, 0.5, 1, or 2 μM ciprofloxacin (for the experiments shown in Figs 5 and 6). Seeds were held at 4 °C for 3 d to promote uniform germination before transferring them to a greenhouse (soil) or growth room at 20 °C with constant light at 30 μEinsteins m⁻² s⁻¹.

#### Obtaining homozygous T-DNA insertion mutants

Wt plants and plants homozygous for a T-DNA insertion near the 3′ end (drt100-1) or in the middle (drt100-2) of the single exon in the *DRT100* gene were selected from SALK lines 021479 and 102492, respectively, using PCR-based genotyping. Primers to distinguish wt alleles from those containing the T-DNA insertion were designed using the T-DNA express tool at http://signal.salk.edu/cgi-bin/dnaexpress (see Table 2). The wt plants and plants homozygous for a T-DNA insertion in the third to last exon of the *cpRecA* gene (*cprecA*) were selected similarly from SALK line 072979. Seeds from single individual homozygous plants (wt or *cprecA*; generation 1) were collected and raised to maturity (generation 2). Seeds from two individuals were then collected (generation 3) and used to produce the plants used for chloroplast isolation and PFGE experiments. Two individuals were selected at each generation starting at generation 3 to give rise to generations 4–7.

#### Measurement of mRNA levels for DRT100 and cpRecA using reverse transcriptase real-time quantitative PCR

RNA from shoots of 13-day-old seedlings was prepared using a Sigma Spectrum™ Plant Total RNA Kit, treated with DNase (DNA-free™ Kit, Applied Biosystems, http://www3.appliedbiosystems.com/AB_Home/index.htm) and reverse transcribed using the BioRad iScript™ cDNA synthesis kit (www.biorad.com). Amplification of 2 μl of cDNA was carried out using the BioRad iQ™ SYBR Green Supermix (www.biorad.com) with the primer sets described in Table 2. Following an initial denaturation at 94 °C for 3 min 15 s, 45 cycles of 20 s denaturation at 94 °C, 20 s annealing at 57 °C, and 30 s extension at 72 °C were

### Table 1. Organelle-targeted homologues of bacterial RecA encoded by the *Arabidopsis* nuclear genome

<table>
<thead>
<tr>
<th>Locus</th>
<th>Name</th>
<th>Compartment of gene product</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>AT1G79050</td>
<td>RECA1; cpRecA</td>
<td>Chloroplast</td>
<td>Cerutti et al. (1992); Cao et al. (1997); Shedge et al. (2007)</td>
</tr>
<tr>
<td>AT3G12610</td>
<td>DRT100</td>
<td>Chloroplast</td>
<td>Pang et al. (1992, 1993)</td>
</tr>
<tr>
<td>AT2G19490</td>
<td>RECA2</td>
<td>Chloroplast; mitochondria</td>
<td>Shedge et al. (2007)</td>
</tr>
<tr>
<td>AT3G10140</td>
<td>RECA3</td>
<td>Mitochondria</td>
<td>Khazi et al. (2003); Shedge et al. (2007)</td>
</tr>
<tr>
<td>AT3G32920</td>
<td>None</td>
<td>Mitochondria</td>
<td>Shedge et al. (2007)</td>
</tr>
</tbody>
</table>

* Mitochondrial targeting based on NBCI GenBank annotation.
performed, and amplification of the reactions was monitored using the Chromo 4 real-time detection system (Bio-Rad Laboratories, www.biorad.com). A melting curve from 65 °C to 95 °C was used to confirm the presence of single products. Data were analysed using the Opticon Monitor 3 software (Bio-Rad Laboratories, www.biorad.com) and the amount of each transcript was determined relative to the actin gene ACT2 using the 2–DDCT method (Livak and Schmittgen, 2001). The highest amount was assigned a value of 1 and all other values are expressed relative to 1.

Chloroplast isolation and preparation for PFGE and fluorescence microscopy

Tissue collected from plants grown in a greenhouse was washed for 3–5 min in 0.5% sarkosyl and rinsed exhaustively four times in tap water and four times in distilled water before isolating chloroplasts using the high-salt method that avoids using DNase (Shaver et al., 2006; Rowan et al., 2007). For PFGE and analysis of cpDNA molecules from individual chloroplasts, the isolated chloroplasts were embedded in agarose and lysed overnight at 48 °C in 1 M NaCl, 5 mM EDTA, 1% Sarkosyl, and 200 µg ml⁻¹ proteinase K. Agarose plugs were washed extensively in many plug volumes of 10 mM TRIS, 1 mM EDTA (TE) before soaking in 30 mM sodium acetate, pH 7.5, 50 mM NaCl, 1 mM ZnCl₂, 5% glycerol for 1 h on ice and treating with or without 2.5 U of mung bean nuclease for 15 min or 30 min at 37 °C in a total volume (liquid plus gel) of 120 µl. Digestion was stopped by adding 0.1% Sarkosyl, 10 mM EDTA, pH 9. For analysis of cpDNA from individual chloroplasts, agarose-embedded cpDNA was prepared at a concentration 100–500-fold less than was used for PFGE, stained with 0.1 µg ml⁻¹ ethidium bromide, and visualized as described (Oldenburg and Bendich, 2004). For estimation of genome equivalents per plastid, isolated chloroplasts were stained with 4′,6-diamidino-2-phenylindole (DAPI) and the relative fluorescence intensity (Rfl) was measured as described (Rowan et al., 2004, 2007). Rfl was determined similarly for glutaraldehyde-fixed, DAPI-stained vaccinia virus particles. The number of chloroplast genome equivalents per plastid was calculated using the equation: chloroplast genome equivalents = 1.33V (where V = the DAPI–DNA Rfl of the plastid divided by the mean Rfl of vaccinia virus particles). The value 1.33 is a constant that accounts for the differences between the size and base composition between the Arabidopsis chloroplast genome and the vaccinia virus genome, and was determined as (%AT content of vaccinia virus genome/ %AT content of Arabidopsis chloroplast genome) × (number of bp of vaccinia virus

![Fig. 1. The effect of a T-DNA insertion in DRT100 on cpDNA amount and structure. (A) PFGE of cpDNA obtained from an equal volume of pelleted chloroplasts from wt and drt100-1 mutant plants after staining with ethidium bromide. (B) Blot hybridization of the gel in (A) with an 854 bp cpDNA-specific probe that contains a portion of the petA gene. Similar results were obtained with a different T-DNA insertion allele (drt100-2; data not shown). Immature, entire shoots of plants grown for 16 d post germination; mature, third rosette leaves of plants grown for 30 d post-germination. The ratio of the hybridization signals for each of the lanes is 2.5:2.2:1:1.2 for wt immature:drt100-1 immature:wt mature:drt100-1 mature, respectively. Linear DNA sizes (in kb) are indicated. cz, compression zone.](https://academic.oup.com/jxb/article-abstract/61/10/2575/429840?redirectedFrom=fulltext&searchPhrase=RecA%20maintains%20cpDNA%20in%20Arabidopsis)
Fig. 2. The effect of a T-DNA insertion in cpRecA on cpDNA amount and structure. (A) PFGE of cpDNA obtained from an equal volume of pelleted chloroplasts from wt and cprecA mutant plants after staining with ethidium bromide. (B) Blot hybridization of the gel in (A) with a petA probe. Immature, entire shoots of plants grown for 14 d post-germination; mature, first and second rosette leaves from plants grown for 23 d post-germination. The ratio of the hybridization signals for each of the lanes is 6:3.8:1:1.3 for wt immature:cprecA immature:wt mature:cprecA mature, respectively. Linear DNA sizes (in kb) are indicated. cz, compression zone. (C and D) Size and DNA content of plastids isolated from wt and cprecA mutant plants. Genome equivalents per plastid were determined by DAPI staining, using vaccinia virus particles as a standard [results using this method are similar to those using real-time qPCR (Rowan et al., 2009)]. D14 shoots, entire shoots of plants grown for 14 d post-germination. D23 L1,2, first and second rosette leaves from plants grown for 23 d post-germination. Numbers in the upper right corners of (C) and (D) represent the mean of genome equivalents per plastid. The mean μm² per plastid (and number of plastids analysed) at D14 are 42 (69) for wt and 45 (68) for cprecA. The corresponding values at D23 are 35 (51) for wt and 36 (44) for cprecA. The relative cpDNA amounts among samples are similar when assessed by quantification of the blot hybridization signal (B) and genome copy number (C and D).
genome/number of bp of Arabidopsis chloroplast genome), where %AT for vaccinia virus (Copenhagen strain) is 66.6, %AT for Arabidopsis cpDNA is 64%, number of bp for vaccinia virus DNA is 197361 and number of bp for Arabidopsis cpDNA is 154361. Brightfield images of the chloroplasts were recorded and used to measure plastid area.

Blot hybridization of cpDNA from PFGE
PFGE-separated cpDNA was alkali-denatured, transferred onto a nylon membrane, and neutralized. Procedural details were described previously (Oldenburg and Bendich, 2004b, 2006). An 854 bp fragment of the Arabidopsis petA gene was labelled with alkaline phosphatase using AlkPhos Direct Labeling Reagents, and hybridization was detected using the CDP-Star Detection Reagent (GE Healthcare). The hybridization signals were quantified using NIH Image J software (http://rsb.info.nih.gov/ij/). Lanes on the image of the blot were selected and the software plotted the intensity of the signal down the lane. The area under the curves generated by the software was calculated to determine the strength of the signal from the corresponding regions of the lanes.

Protein alignment, phylogenetic analysis, and targeting sequence prediction
Protein sequences were obtained from NCBI GenBank (http://www.ncbi.nlm.nih.gov/sites/entrez?db=nuccore), aligned using MacVector™ 8.0 (http://www.macvector.com/), and analysed by the Predotar v. 1.03 (http://urgi.versailles.inra.fr/predotar.html) and TargetP 1.1 (http://www.cbs.dtu.dk/services/TargetP) targeting sequence prediction algorithms. The coding sequences, also obtained from NCBI GenBank, were aligned using MacClade 4.0.8. (http://macclade.org/macclade.html), and a Neighbor–Joining tree was created with these aligned sequences using MEGA 4 (http://www.megasoftware.net/) with the default settings.

Results
Organelle-targeted homologues of RecA encoded by the Arabidopsis nuclear genome
For two of the three genes encoding chloroplast-targeted RecA proteins of Arabidopsis, homozygous T-DNA insertion mutants were obtained by PCR-based genotyping of seeds from SALK T-DNA insertion lines (see Materials and methods). The seeds produced by an individual plant of each genotype (homozygous mutants and their wt counterparts) were collected and the resulting plants were raised to maturity in order to obtain enough seeds to generate sufficient plant tissue for the analysis of cpDNA by PFGE and blot hybridization.
Amount and form of cpDNA in wt and drt100 mutant plants

At 16 d post-germination, wt and drt100-1 mutant seedlings had produced 6–7 leaves. For both genotypes, the leaves of the immature plants were <70% of the length reached when fully expanded at maturity. The drt100-1 mutants appeared morphologically indistinguishable from wt plants. DNA from an equal volume of packed chloroplasts isolated from entire shoots was analysed by PFGE and blot hybridization. For both genotypes, some of the cpDNA remained in the well (Fig. 1, immature). The cpDNA that migrated into the gel was found as linear molecules, including bands at the monomeric, dimeric, and trimeric length of the genome, as well as subgenomic fragments. No band migrating between the well and the compression zone (corresponding to supercoiled circular cpDNA) was observed. The relative amounts and forms of cpDNA did not differ between wt and drt100 mutant plants, and these results were similar to those for wt plants of several other species (Shaver et al., 2008). The cpDNA from fully expanded third leaves of mature 30-day-old wt and mutant plants (Fig. 1, mature) showed a similar pattern with no evident difference between the two genotypes. However, the amount of cpDNA was reduced ~2-fold in mature plants for both genotypes. Similar results were obtained for plants homozygous for a T-DNA insertion in the middle of the single exon of the DRT100 gene (drt100-2, data not shown). The amount of DRT100 transcripts was reduced 24-fold in drt100-1 mutants compared with their wt counterparts as measured by real-time quantitative PCR (RT-qPCR), and transcripts were essentially undetectable in drt100-2 mutants (Table 3).

It is concluded that severely reducing the RNA level for the DRT100 gene has no effect on the structure or amount of cpDNA in Arabidopsis plants.

Amount and form of cpDNA in wt and cprecA mutant plants

The cprecA mutants appeared morphologically indistinguishable from wt plants. At 14 d post-germination, the immature seedlings of both wt and cprecA mutants had produced six leaves, and all were <70% of the maximum expanded length at maturity. Chloroplasts were isolated from entire shoots and their DNA was analysed by PFGE, blot hybridization, and fluorescence microscopy. Well-bound cpDNA and the genomic monomer and dimer were detected with wt plants, but the cpDNA of the cprecA mutants did not show prominent bands at the size of the monomer or dimer. Only the well-bound cpDNA and a smear of linear DNA molecules from ~40 kb to 500 kb were evident, and the hybridization signal was only 60% as intense as the signal from DNA obtained from an equal volume of wt chloroplasts (Fig. 2A, B, immature). Linear forms of cpDNA from fully-expanded first and second leaves of mature 23-day-old wt and cprecA mutant plants were not detected, even after a longer exposure (not shown), probably because the amount of DNA from chloroplasts of mature plants was 3- to 6-fold lower than that from an equal volume of chloroplasts from immature plants (Fig. 2A, B,
mature). The \(cpr\text{Rec}A\) transcript level was only 15% of the wt level as determined by RT-qPCR (Table 3). Individual chloroplasts from immature \(cpr\text{Rec}A\) mutant plants contained ~2-fold fewer genome equivalents on average compared with their wt counterparts (Fig. 2C). For mature leaves, the amount of DNA per chloroplast did not differ between wt and \(cpr\text{Rec}A\) mutants (Fig. 2D). It is concluded that the \(cpr\text{Rec}A\) gene has a role in maintaining the structure of cpDNA molecules and the amount of DNA retained in immature chloroplasts.

**Amount and form of cpDNA in \(cpr\text{Rec}A\) \(drt100\text{-}1\) mutant plants**

It is possible that \(cpr\text{Rec}A\) exhibits functional redundancy with \(DRT100\), allowing \(cpr\text{Rec}A\) to compensate for the loss of \(DRT100\) function in \(drt100\) mutant plants. To examine this possibility, the cpDNA of \(cpr\text{Rec}A\) \(drt100\text{-}1\) double mutants was analysed. If these genes have partially overlapping functions, the phenotype of the double mutants is expected to differ from that of the single mutants. \(F_1\) seeds from the cross \(cpr\text{Rec}A\times\text{drt}100\text{-}1\) were allowed to self, generating a segregating \(F_2\) population. Two \(cpr\text{Rec}A\) \(drt100\text{-}1\) individuals and one wt plant isolated from this population were allowed to self. Progeny seeds from these plants and from an independent wt strain (Col) were grown for 13 d. All of these immature plants appeared visually indistinguishable and had produced six leaves; each was <70% of its final expanded length. Chloroplasts were isolated from entire shoots and the cpDNA was analysed by PFGE. The forms of cpDNA for both wt siblings of the double mutants and the independent wt strain consisted of both well-bound cpDNA and migrating linear forms, including bands at the positions of the monomer and dimer (Fig. 3). In contrast, both of the \(drt100\text{-}1\) \(cpr\text{Rec}A\) lines showed a phenotype that was similar to that of the \(cpr\text{Rec}A\) single mutant: linear forms of DNA migrating as a smear of fragments ranging in size from ~40 kb to 500 kb with no prominent bands. Double mutants generated using a reciprocal cross (\(\text{drt}100\text{-}2\times\text{cpr}\text{Rec}A\)) gave the same result (data not shown). If \(DRT100\) and \(cpr\text{Rec}A\) acted redundantly, the cpDNA phenotype of double mutants is expected to differ.
from the phenotype of either single mutant. The double mutant phenotype did not differ from that of the cprecA single mutant. Thus, these genes do not act redundantly.

Single-stranded cpDNA in wt and cprecA mutants

In *E. coli*, RecA is involved in the repair of single-stranded gaps and DSBs (Michel et al., 2007). During the repair of DSBs, single-stranded DNA (ssDNA) is produced through the exonucleolytic action of the RecBCD protein (Kowalczykowski, 2000). The RecA protein binds along the single-stranded regions and facilitates repair by homologous recombination, while preventing more extensive exonuclease activity (Skarstad and Boye, 1993). Thus, it might be expected that a reduction in RecA activity would result in more ssDNA. Wt and cprecA cpDNA (from 14-day-old seedling shoots) were treated with mung bean nuclease [an enzyme with endonuclease activity on single-strand gaps in double-stranded DNA (dsDNA) and 5′ to 3′ exonuclease activity on ssDNA (Kroeker and Kowalski, 1978)] for 15 min or 30 min or left the DNA untreated. Linear genomic oligomers of wt cpDNA were unaffected by treatment with this nuclease (Fig. 4). However, the amount of well-bound cpDNA decreased after treatment with mung bean nuclease. This result indicates that the well-bound form of cpDNA contains single-stranded regions, whereas the linear genomic oligomers consist primarily of dsDNA. An increase in the smear of subgenomic fragments smaller than 50 kb is also apparent after digestion with mung bean nuclease. These fragments may arise from cpDNA that has been liberated from the well. Thus, the replicating forms of cpDNA from young wt plants may consist of single-stranded regions interspersed with dsDNA at intervals of <50 kb.

After treatment with mung bean nuclease, cprecA mutants exhibited a decrease in both the hybridization signal from the well-bound cpDNA and the range of sizes of the linear forms of cpDNA. Thus, all of the forms of cpDNA in the cprecA mutants contain single-stranded regions, but only the well-bound cpDNA in wt contains ssDNA detectable with this enzyme, suggesting that cpRecA and *E. coli* RecA may act similarly.

![Fig. 6.](https://academic.oup.com/jxb/article-abstract/61/10/2575/429840)

Comparison of the structural forms of individual cpDNA molecules in wt and cprecA plants after ciprofloxacin treatment. Wt and cprecA plants after 11 d of growth followed by 3 d of treatment with water or 30 μM ciprofloxacin. (A) Deproteinized cpDNA was stained with ethidium bromide, visualized by fluorescence microscopy, and characterized by structural class. Class I structures, complex forms consisting of a network of connected fibres or fibres connected to a large, dense core; class II structures, complex forms with a greater number of unconnected fibres than connected fibres; class III structures, unconnected fibres without a complex form. Representative images of class I, II, and III molecules are shown in the insets. The small dots in the images are shorter fibres of DNA that remained condensed during preparation for fluorescence microscopy. The scale bar in the class I image is 10 μm and applies to all images in (A). The number of molecules examined for wt, wt+cipro, cprecA, and cprecA+cipro was 62, 48, 56, and 57, respectively. No circular forms were observed among these 223 molecules. A chi-squares test of independence showed that the distributions of molecules among classes I–III for wt+cipro, cprecA, and cprecA+cipro differed from the wt (*P* <0.001). (B) Representative images of wt and cprecA plants after treatment with ciprofloxacin. Treatment with ciprofloxacin caused bleaching along the vasculature of older leaves of both wt and cprecA mutant plants. Arrows indicate bleaching along the vasculature. The small gradations are millimetres.
Double-strand cpDNA breaks in wt and cprecA mutants

It is possible that the extensive smear of linear cpDNA fragments in cprecA mutants is due to a lack of DNA damage repair. Thus, cprecA mutant plants would be especially vulnerable to agents that damage DNA. Also, some types of DNA damage, such as DSB, in wt plants would be expected to produce a similar cpDNA phenotype to that in cprecA mutants. Ciprofloxacin is a type II topoisomerase inhibitor that induces DSBs during DNA replication in prokaryotes, but not in the eukaryotic nucleus (Jacoby, 2005), and has been shown to affect chloroplasts in Arabidopsis (Wall et al., 2004). The nuclear genome of Arabidopsis encodes organelle-targeted type II topoisomerases homologous to bacterial gyrase: a GyrA subunit that is targeted to mitochondria only, and one GyrB subunit that is targeted to mitochondria only, and one GyrB subunit that is targeted to chloroplasts only (Wall et al., 2004). Wt and cprecA plants grown on agar plates containing 0, 0.5, 1, or 2 μM ciprofloxacin were investigated.

The immature, 14-day-old wt seedlings grown without ciprofloxacin had produced 6–7 rosette leaves and all were <70% of their final size (Fig. 5C). After 14 d of growth in 0.5 μM ciprofloxacin, wt seedlings had a more compact rosette and produced 5–6 leaves, but appeared otherwise similar to the control plants grown without ciprofloxacin. Plants grown in 1 μM ciprofloxacin appeared slightly chlorotic and produced only two rosette leaves; and those in 2 μM ciprofloxacin appeared extremely chlorotic and produced no rosette leaves. The cprecA plants appeared similar to wild-type plants for all growth conditions except 1 μM ciprofloxacin. Under this condition, cprecA mutants appeared more chlorotic and had smaller rosette leaves than the wt. Thus, a dose-dependent effect of ciprofloxacin on seedling morphology is found for both the wt and cprecA. The effect, however, is more pronounced for the cprecA mutant.

For wt plants, growth in 0.5 μM ciprofloxacin resulted in a less distinct monomer band and no DNA band at the dimer position (Fig. 5B). The monomer band was completely absent in wt plants grown in 1 μM or 2 μM ciprofloxacin. The size range of the smear of cpDNA fragments for cprecA plants became smaller with increasing concentrations of ciprofloxacin. The decrease in prevalence of the oligomeric bands (and decline in the amount of well-bound DNA) with increasing concentrations of ciprofloxacin for the wt plants probably reflects an increase in DSBs caused by ciprofloxacin-mediated inhibition of type II topoisomerase activity (Jacoby, 2005). Similarly, the decrease in the monomeric and dimeric bands and increase in the smear of linear DNA for the cprecA mutants would be due to an increase in DSBs without repair.

In order to characterize further the effect of the cprecA mutation and cpDNA damage on cpDNA structure, individual cpDNA molecules were examined using fluorescence microscopy of deproteinized cpDNA stained with ethidium bromide. After 11 d of growth, wt and cprecA plants were either mock treated (with water) or treated with 30 μM ciprofloxacin and allowed to grow for a further 3 d. Plants that were treated with ciprofloxacin showed bleaching along the vasculature of leaves that were produced before the treatment with the drug (Fig. 6B, white arrows). The basal halves of the leaves that were produced after the treatment were entirely bleached. Figure 6A shows that most (71%) of the cpDNA images from mock-treated wt plants were found as class I forms (complex, multigenomic, branched linear structures with more DNA fibres connected to a core than unconnected fibres). Class II forms (complex structures with more unconnected DNA fibres than fibres connected to a core) made up 16% and class III (no complex structures; only unconnected DNA fibres) made

Table 2. Primers used for PCR genotyping and RT-qPCR

<table>
<thead>
<tr>
<th>Product</th>
<th>Forward primer</th>
<th>Reverse primer</th>
</tr>
</thead>
<tbody>
<tr>
<td>Genotyping</td>
<td></td>
<td></td>
</tr>
<tr>
<td>drt100-1</td>
<td>CTGATTTTCTTCCCGGGAG (wt) or LBb1 (drt100-1)</td>
<td>TACCTGTCCGTGTCGTAAC</td>
</tr>
<tr>
<td>drt100-2</td>
<td>GATACCATGCACATGCAAATG (wt) or LBb1 (drt100-2)</td>
<td>GGGATAATGCTCTTCGAAAGGGCGC</td>
</tr>
<tr>
<td>cprecA</td>
<td>TAGGGTGAGATTGGAATGCAAG (wt) or LBb1 (cprecA)</td>
<td>AAGAGCTGCTGCTCATCTAAAG</td>
</tr>
<tr>
<td>LBb1</td>
<td>GCGTGGACCGCTTGCAACT</td>
<td></td>
</tr>
<tr>
<td>RT-qPCR</td>
<td></td>
<td></td>
</tr>
<tr>
<td>DRT100</td>
<td>GCACTCACTTCCCTCGTTCT</td>
<td>TTTCCGGAGGGTTTGGCGATT</td>
</tr>
<tr>
<td>cpRecA</td>
<td>CAGAGGTTACTAGCGGAGGAA</td>
<td>ACTCTTAGCAGGCGGCGAA</td>
</tr>
<tr>
<td>ACT2</td>
<td>GCCATCCAAGCTGTCTCTCT</td>
<td>GCTCGTAGCAGCAAGCACAA</td>
</tr>
</tbody>
</table>

Table 3. mRNA for DRT100 and cpRecA in wt and T-DNA mutants

<table>
<thead>
<tr>
<th>Genotype</th>
<th>DRT100 transcripts</th>
<th>cpRecA transcripts</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wt</td>
<td>0.81±0.08</td>
<td>0.00±0.00</td>
</tr>
<tr>
<td>Wt</td>
<td>1.00±0.07</td>
<td>0.03±0.00</td>
</tr>
<tr>
<td>Wt</td>
<td>1.00±0.10</td>
<td>0.15±0.03</td>
</tr>
<tr>
<td>Wt</td>
<td>0.03±0.00</td>
<td>0.00±0.00</td>
</tr>
<tr>
<td>Wt</td>
<td>0.15±0.03</td>
<td></td>
</tr>
<tr>
<td>Wt</td>
<td>0.07±0.00</td>
<td>0.00±0.00</td>
</tr>
<tr>
<td>Wt</td>
<td>0.08±0.00</td>
<td>0.00±0.00</td>
</tr>
<tr>
<td>Wt</td>
<td>0.00±0.00</td>
<td>0.00±0.00</td>
</tr>
<tr>
<td>Wt</td>
<td>0.00±0.00</td>
<td>0.00±0.00</td>
</tr>
<tr>
<td>Wt</td>
<td>0.00±0.00</td>
<td>0.00±0.00</td>
</tr>
</tbody>
</table>

a Wt plants from the T-DNA insertion line for the drt100-1 allele.
b Wt plants from the T-DNA insertion line for the drt100-2 allele.
c Wt plants from the T-DNA insertion line for the cprecA allele.
up only 13% of the forms of cpDNA. After treatment with ciprofloxacin, the proportion of class I forms decreased to 35%, and there was a corresponding increase in the proportion of class II forms (52%) but no increase in class III forms. Class I represented only 18% of all forms in mock-treated cprecA mutants, with most of the cpDNA found in class II (55%) and class III (27%). After treatment with ciprofloxacin, there was only a minor decrease in the proportion of class I (13%) and a minor increase in the proportion of class III (33%). It is surmised that ciprofloxacin-induced DSBs cause a conversion from class I to class II forms in wt plants. The unconnected fibres seen in classes II and III appear as simple linear molecules that would migrate in the smear of cpDNA during PFGE, whereas the complex branched forms remain immobile in the well of the gel (Oldenburg and Bendich, 2004b). Both assay procedures used to assess molecular integrity (Figs. 2–6) show that the molecules of cpDNA from cprecA mutant plants are more fragmented than those of wt plants.

**Phenotypic effects after growth of cprecA mutants for multiple generations**

None of the cprecA mutants from the first three generations of selfing showed any abnormal visible phenotype. During the next four generations, however, cprecA mutants exhibited a range of visible phenotypes (Fig. 7). The most common was colour variegation in leaf sectors. Tissue necrosis in mature leaves was also observed. The percentage of plants exhibiting a phenotype varied from 1.1% to 4.2% and did not increase with each successive generation (Table 4). Additional unlinked inserts in the genomes of these mutants might be responsible for the phenotypes observed. This possibility, however, is unlikely because none of the wt plants isolated from the same T-DNA insertion line exhibited an abnormal phenotype in seven generations, suggesting that the phenotypes observed in the cprecA mutants are caused by disruption of the cprecA gene. It is possible, however, that additional T-DNA insertions very tightly linked to the insertion in cpRecA may be present in this line. PCR was used to determine whether any of the 13 genes in the region from 25 kb upstream to 25 kb downstream of cprecA contained an insertion (Supplementary Table S1 available at JXB online). For each gene, two individual plants were examined from different generations that either contained or did not contain the insert in cpRecA. No inserts were found in the genes in the 50 kb region surrounding cpRecA. It is concluded that the T-DNA disruption of the cpRecA gene is responsible for the phenotypes observed.

### Discussion

There is a dearth of information about repair of DNA in chloroplasts. Two Arabidopsis nuclear genes predicted to encode chloroplast-targeted proteins that are similar to bacterial RecA were investigated. Impairing the function of one of these genes, cpRecA, led to abnormalities in both cpDNA and plant morphology. Although chloroplast-targeted E. coli RecA homologues have been implicated in cpDNA maintenance and repair (Cerutti et al., 1992, 1993, 1995; Pang et al., 1992; Cao et al., 1997; Inouye et al., 2008), the results provide the first evidence that homologous recombination is used to maintain the structural integrity of DNA molecules in the chloroplasts of vascular plants.
A mutation in the other gene, DRT100, had no apparent effect and did not act redundantly with \textit{cpRecA}. The amino acid sequence of the DRT100 protein differs considerably from that of bacterial RecA and other nucleo-encoded, organelle-targeted RecA proteins (Supplementary Fig. S1 at JXB online). Pang \textit{et al.} (1992) reported that this protein had a chloroplast localization signal. Two current signal peptide prediction algorithms, TargetP and Predotar, both predict that this protein is targeted to the secretory pathway (see Materials and methods). Although \textit{DRT100} can complement \textit{recA} mutants of \textit{Escherichia coli} (Pang \textit{et al.}, 1992), it may not function as a RecA protein in chloroplasts.

\textbf{Abnormal cpDNA molecules in \textit{cprecA} mutants}

In PFGE, the cpDNA from wt plants and \textit{drt100} mutants was found as well-bound and as a smear of migrating linear forms, including discrete bands representing the genomic monomer, dimer, and trimer. The well-bound cpDNA is partially single stranded, consistent with an earlier proposal that the DNA that remains in the well is a branched form of replicating DNA that serves as the segregating genetic unit, the chromosome, in both chloroplasts (Oldenburg and Bendich, 2004\textit{a}, \textit{b}) and mitochondria (Oldenburg and Bendich, 1996). The molecular integrity of the branched cpDNA was greatly reduced in the \textit{cprecA} mutants, as it was in wt plants treated with the prokaryotic type II topoisomerase inhibitor ciprofloxacin (Fig. 6A). Fragmentation of the cpDNA molecules in \textit{cprecA} mutants increased only slightly after treatment with ciprofloxacin. If cpDNA replication occurs through a recombination-dependent mechanism (Oldenburg and Bendich, 2004\textit{a}, \textit{b}), cpDNA replication might be compromised in \textit{cprecA} mutants. If so, fewer topoisomerase-DNA complexes would be expected for the cpDNA of \textit{cprecA} mutants, leading to fewer instances of DSBs generated by ciprofloxacin-mediated suppression of topoisomerase activity.

During PFGE, the migrating cpDNA of \textit{cprecA} mutants appeared as a smear of linear fragments without any distinct bands (Fig. 2). Prominent linear genomic monomers and oligomers of cpDNA have been observed for many wt plant species, including maize (\textit{Zea mays}; Oldenburg and Bendich, 2004\textit{a}, \textit{b}; Oldenburg \textit{et al.}, 2006), \textit{Medicago truncatula} (Shaver \textit{et al.}, 2006, 2008), wheat (\textit{Triticum aestivum}; Shaver \textit{et al.}, 2006), \textit{Chenopodium album} (Backert \textit{et al.}, 1995), tobacco (\textit{Nicotiana tabacum}; Backert \textit{et al.}, 1995; Lilly \textit{et al.}, 2001; Scharff and Koop, 2006), pea (\textit{Pisum sativum}; Bendich and Smith, 1990; Lilly \textit{et al.}, 2001), watermelon (\textit{Citrullus vulgaris}; Bendich and Smith, 1990), spinach (\textit{Spinacia oleracea}; Deng \textit{et al.}, 1989), and \textit{Marchantia polymorpha} (Oldenburg and Bendich, 1998). Thus, it is likely that the cpDNA repair process has been conserved among plants and maintains linear monomeric and oligomeric forms of cpDNA. The lack of such monomers and oligomers in \textit{cprecA} mutants may result from the failure either to repair DSBs or to produce telomeric ends that prevent degradation of such linear DNA molecules within the plastid. Homologous recombination has been suggested as a telomerase-independent mechanism for maintenance of eukaryotic telomeres (Tarsounas and West, 2005; Raji and Hartsuiker, 2006) and may similarly protect the ends of linear cpDNA molecules even though the structures of linear cpDNA ends and eukaryotic telomeres differ.

It might seem surprising that ciprofloxacin did not have a more substantial impact on the growth or morphology of \textit{cprecA} mutants than wt plants. On the other hand, cpDNA may be present in excess, so that morphological effects may go undetected until most of the cpDNA is damaged. Additionally, DSBs generated by ciprofloxacin remain in a complex with topoisomerase (Chen \textit{et al.}, 1996) and may not act as substrates for repair by RecA. Furthermore, the additional chloroplast-targeted RecA protein (RECA2; AT2G19490) may partially complement the \textit{cprecA} mutation.

\textbf{Delayed phenotype of \textit{cprecA} mutants}

It is demonstrated that a recombinational repair system is required for the stability of the chloroplast chromosome. After several generations of selfing, the \textit{cprecA} mutants exhibited visible phenotypes, including tissue necrosis and colour variegation among leaf sectors. Why was the morphological consequence of cpDNA damage apparent only after several generations? One reason follows from the non-Mendelian nature of organelar inheritance and another from analogy to recombination-dependent DNA repair in \textit{E. coli}.

It may have taken several generations to generate sectors of cells containing chloroplasts in which most of the DNA is damaged or degraded, given the stochastic partitioning of damaged and wt chloroplast chromosomes during chloroplast division and the stochastic partitioning of chloroplasts during cell division (Birky, 2001). Many animals experience a lifetime accumulation of mtDNA damage, which varies among cells in different tissues and organs (Trifunovic and Larsson, 2008; Kukat and Trifunovic, 2009). Variation in mtDNA damage among cells may reflect differential efficiency of mtDNA repair among tissues (Gredilla \textit{et al.}, 2008). Impairment of mitochondrial function due to accumulated mtDNA mutations is proposed to contribute to ageing. Decline of respiratory function may result from random partitioning of normal and damaged mtDNA molecules during cell division, leading to clonal expansion of cells with damaged mtDNA (Nagley \textit{et al.}, 1992; Trifunovic and Larsson, 2008). As it takes many cell generations to observe the phenotypic consequences of mtDNA damage during ageing in animals, it may similarly require many cell generations for the consequences of cpDNA damage in \textit{cprecA} mutants to become evident. Another possibility is that the degradation of cpDNA with damaged ends occurs rather slowly. In \textit{E. coli}, RecA-dependent recombination involves the RecBCD exonuclease, which is maintained at a low concentration (~10 enzyme complexes per cell) in order to prevent unintentional degradation of the \textit{E. coli} chromosome (Dillingham and Kowalczykowski, 2008). The functional analogue of the RecBCD complex for \textit{Arabidopsis} chloroplasts (which is not yet known) may similarly be maintained.
at a low concentration, leading to a gradual degradation of cpDNA molecules when \(cpRecA\) activity is impaired.

Visible phenotypes due to chloroplast chromosome instability might be expected to increase in frequency with each successive generation. Instability of the nuclear genome due to telomerase deficiency results in increasingly deleterious phenotypes with successive generations in Arabidopsis and mouse (Lee et al., 1998; Riha et al., 2001). However, the percentage of cprecA mutant plants with observable phenotypes of cprecA mutants varied from 1.1% to 4.2% but with no generational increase. One explanation for this surprising observation is that additional repair processes might be activated after several generations without functional cprecA. For example, non-homologous end-joining (a mechanism not yet known for cpDNA repair) is used as an alternative to homologous recombination for DSB repair in both prokaryotes and eukaryotes (Pitcher et al., 2007). RECA2 is targeted to both mitochondria and chloroplasts, might be partially functionally redundant with cprecA, and might compensate for the lack of cprecA activity. Another possibility is that the chloroplast chromosomes that are transmitted to the next generation might sustain less damage than cpDNA in green cells because gametes originate from meristematic cells where photosynthesis [a major source of DNA-damaging reactive oxygen species (Roldan-Arjona and Ariza, 2009)] does not occur.

The chloroplast mutator of barley leads to cytoplasically inherited chlorophyll deficiencies (Prina, 1992), one of which is due to point mutations in the chloroplast infA gene (Landau et al., 2007), suggesting that the chloroplast mutator gene may be responsible for cpDNA repair in barley. The chloroplast mutator (CHM) of Arabidopsis was originally identified as a nuclear mutation affecting maintenance of cpDNA (Redei and Plurad, 1973) and is now recognized as a homologue of the E. coli mismatch repair gene, MutS. Although CHM (renamed MSFL) is targeted to both mitochondria and chloroplasts, only the mitochondrial genome is affected in chm mutants (Martinez-Zapater et al., 1992; Sakamoto et al., 1996; Shedeg et al., 2007). A recent study identifies a function for the Whirly proteins (ssDNA-binding proteins) in stabilizing the chloroplast genomes of A. thaliana and Z. mays (Marechal et al., 2009). For Arabidopsis, the authors report variegated phenotypes arising in 4.6% of AtWhy1 and AtWhy3 double knockout plants due to illegitimate recombination. Many Arabidopsis variegation mutants have not yet been characterized (Yu et al., 2007). Future research on these proteins and RECA2 will shed light on the spectrum of repair processes for cpDNA.

In conclusion, the \(cpRecA\) gene maintains the integrity of cpDNA, possibly through repair of DSBs or maintenance of telomere-like ends of linear molecules. The phenotypic abnormalities that arise after multiple generations are reminiscent of the ageing phenotypes attributed to declining mitochondrial function in yeast and animals (Mandavalli et al., 2002), an interesting parallel considering that the mitochondrion-targeted RecA homologue was lost in the common ancestor of yeast and animals (Lin et al., 2006).

**Accession numbers**

Arabidopsis Genome Initiative (AGI) locus identifiers for the RecA homologues described in this article are listed in Table 1. The Arabidopsis Information Resource (TAIR) accession numbers are 2091260 for DRT100 and 2207445 for \(cpRecA\).

**Supplementary data**

Supplementary data are available at JXB online.

**Supplementary Table S1.** PCR-based detection of insertions in 13 genes in the 50 kb region surrounding \(cpRecA\). The primers were used to amplify DNA from at least two individuals from two different generations with or without the insertion in \(cpRecA\). If the gene was larger than 1 kb, then overlapping sets of primers were used to span the entire length of the gene (indicated in the table after the name of the gene as “s1, s2, etc.”). N indicates that no inserts were detected. The primers were used to amplify DNA from at least two individuals from two different generations with or without the insertion in \(cpRecA\). If the gene was larger than 1 kb, then overlapping sets of primers were used to span the entire length of the gene (indicated in the table after the name of the gene as “s1, s2, etc.”). N indicates that no inserts were detected.

**Fig. S1.** Protein alignment and phylogeny of coding regions of putative RecA homologues. (A) Protein alignment for cpRecA and three other RecA homologues (DRT100, AT2G19490, and AT3G10140) encoded in the Arabidopsis nuclear genome, Chlamydomonas reinhardtii (C. reinhardtii, nuclear-encoded, chloroplast-targeted homologue), Synechococcus elongatus (S. elongatus, cyanobacterial homologue), and E. coli RecA. The last line shows the consensus sequence. (B) Neighbor–Joining tree of coding sequences (excluding the region coding for the signal peptide). Values at nodes are bootstrap values from 20 000 replicates.

**Acknowledgements**

We thank Amanda Rychel for assistance with the phylogenetic analysis of RecA sequences, and Jennifer Nemhauser and the Comparative Genomics Center at the University of Washington for material support. This investigation was supported in part by Public Health Service National Research Award T32 GM07270 from the National Institute of General Medical Sciences and by the Frye-Hotson-Rigg fellowship from the University of Washington Department of Biology.

**References**


Niedernhofer LJ. 2008. DNA repair is crucial for maintaining hematopoietic stem cell function. DNA Repair 7, 523–529.


