Microsatellite instability in Arabidopsis increases with plant development

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Abstract: Plant development consists of the initial phase of intensive cell division followed by continuous genome endoreduplication, cell growth, and elongation. The maintenance of genome stability under these conditions is the main task performed by DNA repair and genome surveillance mechanisms. Our previous work showed that the rate of homologous recombination repair in older plants decreases. We hypothesized that this age-dependent decrease in the recombination rate is paralleled with other changes in DNA repair capacity. Here, we analyzed microsatellite stability using transgenic Arabidopsis (Arabidopsis thaliana) plants that carry the nonfunctional -glucuronidase gene disrupted by microsatellite repeats. We found that microsatellite instability increased dramatically with plant age. We analyzed the contribution of various mechanisms to microsatellite instability, including replication errors and mistakes of DNA repair mechanisms such as mismatch repair, excision repair, and strand break repair. Analysis of total DNA polymerase activity using partially purified protein extracts showed an age-dependent decrease in activity and an increase in fidelity. Analysis of the steady-state RNA level of DNA replicative polymerases α, β, Pol I-like A, and Pol I-like B and the expression of mutS homolog 2 (Msh2) and Msh6 showed an age-dependent decrease. An in vitro repair assay showed lower efficiency of nonhomologous end joining in older plants, paralleled by an increase in Ku70 gene expression. Thus, we assume that the more frequent involvement of nonhomologous end joining in strand break repair and the less efficient end-joining repair together with lower levels of mismatch repair activities may be the main contributors to the observed age-dependent increase in microsatellite instability.

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Microsatellite instability in Arabidopsis thaliana

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Microsatellite instability in *Arabidopsis thaliana* increases with plant development

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Plant development consists of the initial phase of intensive cell division followed by continuous genome endoreduplication, cell growth and elongation. The maintenance of genome stability under these conditions is the main task performed by DNA repair and genome surveillance mechanisms. Our previous work showed that the rate of homologous recombination (HR) repair in older plants decreases. We hypothesized that this age-dependent decrease in the recombination rate is paralleled with other changes in DNA repair capacity. Here, we analyzed microsatellite stability using transgenic plants that carry the non-functional β-glucuronidase gene disrupted by microsatellite repeats. We found that microsatellite instability increased dramatically with plant age. We analyzed the contribution of various mechanisms to microsatellite instability including replication errors, mistakes of DNA repair mechanisms such as mismatch repair, excision repair and strand break repair. Analysis of total DNA polymerase activity using partially purified protein extracts showed an age-dependent decrease in activity and an increase in fidelity. Analysis of the steady-state RNA level of DNA replicative polymerases α, δ, Pol I-likeA and Pol I-like B and the expression of Msh2 and Msh6 showed an age-dependent decrease. An in vitro repair assay showed lower efficiency of non-homologous end joining in older plants, paralleled by an increase in Ku70 gene expression. Thus, we assume that the more frequent involvement of non-homologous end-joining in strand break repair and less efficient end-joining repair together with lower level of mismatch repair activities may be the main contributors to the observed age-dependent increase in microsatellite instability.

**Key words:** microsatellite instability, age-dependent changes, DNA repair, DNA polymerase activity and fidelity, non-homologous end-joining, homologous recombination, excision repair, *Arabidopsis thaliana*
INTRODUCTION

The genome of Arabidopsis thaliana is extensively repetitive, which leads many to believe that Arabidopsis is subject to ancient autoploid events with many subsequent rearrangements and alterations (Meinke et al., 1998; Arabidopsis Genome, 2000; Blanc et al., 2000). Despite the highly reduplicated genome with the potential for a high degree of genetic redundancy, maintaining a consistent level of genome stability is critical. This is especially important when considering that plants do not have a predetermined germ line and gametes are produced from meristematic cells that are products of many somatic cell divisions (Hays, 2002). Furthermore, as plants are sessile organisms, they are continuously exposed to various genotoxic elements such as heavy metals, reactive oxygen species, and ultraviolet irradiation. This constant exposure to harsh environmental conditions imposes a need for precise and efficient genome-maintenance pathways as the persistence of DNA damage and mutagenesis can decrease the fitness of current and future generations (Britt, 1996).

DNA mutagenesis cannot solely be attributed to environmentally induced genotoxic stress as DNA is prone to spontaneous or replication-induced mutagenesis. For example, transitions of 5-methylcytosine to thymine are common spontaneous mutations (Britt, 1996), while DNA replication and repair infidelity can induce numerous errors (Sia et al., 1997; Tuteja et al., 2001). Hundreds of mutations are introduced upon each genome replication due to DNA polymerase infidelity. Repetitive elements are particularly prone to this type of mutation due to replication slippage, which refers to DNA polymerase dissociation during replication of short repetitive sequences followed by the separation and subsequent re-association of the daughter strand in a different but identical repeat (Viguera et al., 2001). Polymerase re-loading and the resumption of DNA synthesis can result in addition or subtraction of the repeated sequence. Microsatellites, the simple tandem repeats of 1-6 nucleotides (Viguera et al., 2001), are highly susceptible to replication slippage.

The frequency at which these and other polymerase-derived errors persist depends largely on the DNA polymerase proofreading activity and the precision and fidelity of core DNA repair enzymes. Since many repair pathways involve DNA polymerase activity, many of them can potentially contribute to an increase in microsatellite instability. Mismatch repair (MMR) is a
repair mechanism involved in the correction of replication errors. It is essential for the maintenance of repeated sequences as mutations in MMR genes are associated with a substantial destabilization of microsatellites (Karran, 1996), and in humans, microsatellite instability increases with aging (Ben Yehuda et al., 2000; Krichevsky et al., 2004; Neri et al., 2005).

The fidelity of different repair pathways can vary largely in the same or similar types of lesions. For example, single- and double-strand breaks (SSBs and DSBs) can be repaired via homologous recombination (HR) or non-homologous end joining (NHEJ) pathways (Britt, 1996; Tuteja et al., 2001; Kovalchuk et al., 2004; Boyko et al., 2006a). Out of these pathways, HR is believed to be precise and largely error-free, while NHEJ can induce numerous mutations ranging from single to thousand nucleotide insertions or deletions (Pelczar et al., 2003; Boyko et al., 2006b). It is unclear how either of these pathways is chosen for repair, but recent evidence from our laboratory suggests that the HR pathway is developmentally regulated, whereby NHEJ is up-regulated and HR is down-regulated with plant development (Boyko et al., 2006b). Currently, there is no information as to whether other DNA repair pathways in plants are developmentally regulated.

Previous publications suggest that aging human cells have a higher frequency of mutations in microsatellites (Ben Yehuda et al., 2000; Krichevsky et al., 2004; Neri et al., 2005). No such data exist for plants. Here, we investigated microsatellite stability during the development of Arabidopsis thaliana using the uidA (GUS) reporter gene inactivated by an artificially incorporated microsatellite (Azaiez et al., 2006). We found a strong increase in instability with plant maturity. We tested the contribution of various repair pathways to age-dependent microsatellite instability and suggest that these changes are primarily due to more frequent involvement of the NHEJ pathway in DNA repair.
RESULTS

Older plants have higher rates of microsatellite instability

We analyzed microsatellite stability using a transgenic *Arabidopsis thaliana* reporter line 121A carrying the GUS gene interrupted by a 31-nucleotide insert comprising a stretch of 16 Gs (Azaiez et al., 2006). The insertion of this microsatellite results in a frameshift generating a premature stop codon (Figure 1). Gene expression and the protein activity can be restored via deletion of one guanine nucleotide or insertion of two guanine nucleotides (Figure 1).

To analyze whether the appearance of blue spots is the result of changes at the microsatellite locus, we have excised plant tissues with blue spots from lightly stained three-week-old plants, prepared the DNA, amplified the region around the satellite using PCR techniques, cloned individual clones and sequenced them. Out of 96 individual clones sent for sequencing, 64 gave readable sequences. Among those, 10 represented the wild-type sequence containing all 16 guanines, 22 had a single guanine deletion, 2 – a single guanine deletion and a single G → A mutation, 14 – a deletion of two guanines, 8 – a deletion of three guanines, 4 – a deletion of four guanines, 1 – a deletion of five guanines, 1 – an insertion of one guanine, and 2 – an insertion of 2 guanines (Figure 1C; Figure 1S). To test whether these sequences are not sequence artifacts, we picked ten random clones containing deletions/insertions of either one or several guanines and performed sequencing of the antiparallel strand; in all cases, we found similar results. Deletions seem to be a predominant type of revertants in this microsatellite region.

Next, we germinated plants from line 121A and harvested them for analysis of mutations at the reporter locus at 3, 5, 7, 10, 13, 16, 19 or 22 days post germination (dpg). We found that the number of blue spots increased dramatically with an increase in plant age; the number of events increased from 0.2 to 343 per plant (Figure 2A; Table 1S). This increase could be simply attributed to the increase in the number of cells as well as the number of genomes carrying the target transgene.

To address this, we analyzed mutation rates, i.e., the number of events per single cell genome. To measure mutation rates in synthetic microsatellites, we first analyzed the number of genomes per cell in plants of different age. This analysis showed that the number of genomes increased from
1.13E+06 in 3-day-old plants to 3.02E+07 in 22-day-old plants (Table 1S). To obtain the mutation rate, we related the mutation frequency to the number of genomes and found that the mutation rate increased 64.2-fold from 1.77E-07 to 1.14E-05 in 3-day-old and 22-day-old plants, respectively (Figure 2B; Table 1S).

Since the mutation rate in microsatellite regions increases with age, it can be suggested that DNA repair in plants is indeed developmentally regulated. As mentioned above, mutations at microsatellite loci could be generated by many mechanisms, most of which can be related to mistakes attributable to replicative or repair DNA polymerases. To test this hypothesis, we measured DNA polymerase activity, DNA polymerase expression as well as repair efficiency and the expression of various DNA repair genes.

Higher microsatellite mutation rates are not due to lower DNA polymerase activity in older plants

To analyze whether older plants have lower levels of DNA polymerase activity, we used a DNA polymerase activity assay. This assay is based on the ability of a freshly prepared protein extract to extend the DNA sequence of a 15-nt primer upon annealing to a complementary 30-nt template (Figure 2S). Incubation of the extract with the template, the primer and a mix of dGTP and dATP results in extension of the primer by 2 nucleotides. Thus, the appearance of a 17-nt extension product is expected, whereas the appearance of products larger than 17 nt, e.g., 18 nt, would indicate misincorporation of nucleotides. The assay allows the analysis of both DNA polymerase activity, the ability to incorporate nucleotides into a growing DNA chain, and DNA polymerase fidelity, the ability to incorporate correct nucleotides. To decrease the number of samples, we used protein extracts from tissue harvested at 5, 12 and 22 dpg. Figure 3A shows a representative image of the assay. The 15-nt product corresponds to the FAM-labeled primer, the 17-nt product represents the extended primer that incorporated precisely two nucleotides, guanine and adenine, whereas the 18-nt product results from incorporation of three nucleotides including guanine, adenine and another misincorporated nucleotide, either adenine or guanine (Figure 3A).

Analysis of the intensity of the 17-nt product showed that DNA polymerase activity inversely correlated with plant age; the older the plants, the lower was the DNA activity (Figure 3A,B,E). Analysis of the intensity of the 18-nt product also showed a substantial age-dependent decrease in product intensity (Figure 3A,C). Relative DNA polymerase fidelity was calculated by relating the intensity of the 18-nt product (due to misincorporation) to the intensity of the 17-nt product.
(precise extension) and then standardizing it to the 5 dpg sample. The analysis showed that 12-day-old plants had comparable DNA polymerase fidelity, whereas 21-day-old plants had a 75% increase in fidelity (Figure 3F).

While analyzing gel images, we noticed the presence of shorter molecules of less than 15 nucleotides. We hypothesized that the appearance of these fragments was due to the exo- and endo-nuclease activity (Figure 3A). The analysis of the intensity of fragments of the 3-nt product showed that the extracts prepared from 5-day-old plants had much higher nuclease activity than those prepared from 12- or 21-day-old plants (Figure 3A,D). Thus, we concluded that younger plants had higher exo-/endonucleolytic activity (Figure 3G).

Steady-state RNA levels of genes coding DNA polymerases are age-dependent

The aforementioned assay gave us a global picture of DNA polymerase activity and fidelity. Unfortunately, it was difficult to decipher the contribution of an individual DNA polymerase in the assay described above. We hypothesized that changes in activity in older plants were in part due to changes in expression of various DNA polymerases. A decrease in DNA polymerase expression in older plants would lead to less protein available for DNA synthesis. Since in our assay it was impossible to find out which DNA polymerases were involved in DNA synthesis, we checked the steady-state RNA level of the following DNA polymerases: α, δ, θ, η, κ, λ, Pol AtREV1, Pol I-like A and Pol I-like B. The analysis showed an age-dependent decrease in the expression of replicative DNA polymerases α, δ, Pol I-like A and Pol I-like B (Table 1). The expression of bypass DNA polymerases η, κ and AtREV1 varied; whereas the expression of Pol η decreased, the expression of Pol κ did not change, and the expression of AtREV1 actually increased with age. The expression of the translesion DNA polymerase AtREV3 increased in 12- and 21- day-old plants, whereas the expression of the translesion Pol θ increased only in 21-day-old plants. Finally, the expression of the DNA repair and recombination DNA Pol λ also increased with age. Replicative DNA polymerases are the most abundant and active in developing organisms. It is thus not surprising to find a correlation between a decrease in the expression of replicative DNA polymerases and a decrease in global DNA polymerase activity. As we mentioned above, microsatellite instability can be the result of mistakes in replication or DNA repair. Such repair mechanisms could include mismatch repair, nucleotide excision repair and
strand break repair. Next, we decided to test the contribution of these repair processes to higher mutation rates at microsatellite loci.

*Mistakes in mismatch repair are unlikely to be the reason of higher microsatellite instability in older plants*

To analyze the possible involvement of MMR, we performed two experiments. First, we crossed the 121A line to an *msh2* mutant, selected homozygous plants and analyzed microsatellite instability in plants of various ages. We hypothesized that if MMR is responsible for the higher rate of mutations at microsatellite loci, complete loss of MMR activity seen in the mutant should result in a much greater increase in the mutation rate in young leaves relative to older leaves. The analysis showed that *msh2* in general had higher microsatellite instability as compared to wild-type plants. However, the difference between old *msh2* plants and young plants was similar to that in wild-type plants; there was approximately a 70-fold increase in the mutation rate in *msh2* plants at 22 dpg vs 3 dpg, as compared to a 64-fold increase in wild-type plants (Table 2S). It has to be noted, however, that since the average number of events in plants increased substantially, it became too difficult to count individual events in each plant (the average number of events in 22-day-old plants was well over 500). We thus cannot completely exclude that older *msh2* plants may have a greater mutation-rate increase at the microsatellite locus of the GUS transgene as compared to wild-type plants.

Next, we tested the expression of genes coding for mismatch repair proteins, MSH2, MSH6 and MSH7. The analysis of steady-state RNA levels showed an age-dependent decrease in the MSH2 and MSH7 genes and a tendency to an increase in the MSH6 gene (Table 2).

*Older plants have less active excision repair*

To test the possible contribution of base excision repair (BER) towards an increase in microsatellite instability, we performed the BER and nucleotide excision repair (NER) assays using extracts prepared from 5-, 12- and 21-day-old plants. The assay is based on the ability of partially purified protein extracts to perform the repair of non-damaged or UV-damaged circular or linear plasmid DNA. Incubation of the plasmid DNA with protein extracts from 5-, 12- and 21-day-old plants showed that repair efficiency of circular DNA was dramatically lower in 21-day-old plants, as compared to 5- or 12-day-old plants (Figure 4). The rate of incorporation of
radionuclides into blunt-cut or non-sticky-ended cut linear DNA was higher than in circular DNA. These types of strand breaks are typically repaired via end-joining repair. In both cases, however, 21-day-old plants showed lower incorporation rates as compared to younger plants (Figure 4).

Repair of a UV-damaged substrate showed yet higher band intensities, which presumably reflects BER/NER repair of UV-damaged DNA. In this case, however, band intensity was the highest in 5-day-old plants; it was the lowest in 21-day-old plants, while in 12-day-old plants, it was intermediate. A decrease in band intensity would suggest less frequent involvement of BER/NER and possibly would reflect the higher number of DNA damages that remain unrepaired. As one of the steps of BER/NER involves the DNA polymerase function and DNA polymerase activity decreases with age, it is possible that the age-dependent decrease in BER/NER efficiency is at least in part due to lower DNA polymerase activity. A higher frequency of mutations at microsatellite loci in older plants could be due to more frequent involvement of BER/NER and thus more frequent mistakes of BER/NER. Since what we observed was the exact opposite, we have concluded that higher microsatellite instability in older plants is not due to increased involvement of BER/NER.

Higher rates of mutations at microsatellite loci could be due to more frequent involvement of NHEJ repair in Arabidopsis

To test whether the involvement of NHEJ has any effect on the frequency of microsatellite stability, we tested the ability of protoplasts prepared from young and old plants to repair double-strand breaks. We used this assay in the past to show the differential efficiency of strand break repair in plant and animal cells (Pelczar et al., 2003; Kovalchuk et al., 2004). For the analysis, protoplasts of 5-day-old, 12-day-old and 21-day-old plants were transfected with 5 μg of linear DNA serving as a template for strand break repair. Our previous experiments showed that the peak of extra-chromosomal strand break repair occurs between 4h and 12h post transfection (Pelczar et al., 2003). Therefore, we incubated DNA with protoplasts for 4h and 12h. Next, the DNA prepared from protoplasts was used for E. coli transfection.

Any difference between the numbers of ampR bacteria obtained from transfection with the DNA recovered from plants of different ages would be a reflection of the efficiency of strand break repair. The analysis showed more ampR bacteria obtained from transfection of older protoplasts; a nearly 50% increase between 5 dpg and 21 dpg was observed (Figure 5A). Next, we
analyzed the percentage of white colonies among all white and blue colonies. The appearance of blue colonies is expected if break repair is faithful, whereas the appearance of white colonies would indicate deletions and insertions of various sizes reflecting error-prone repair. The analysis showed that in 21-day-old plants, the percentage of white colonies was over 1.5-fold higher, as compared to 5-day-old plants (Figure 5B). This experiment suggested that older plants had the increased strand break repair activity but lower fidelity.

Next, we analyzed the steady-state level of *Ku70* and *Rad51* RNAs and found that the expression of *Ku70* increased and the expression of the *Rad51* gene slightly decreased with plant age (Figure 6A). Analysis of protein levels also showed that the amount of KU70 protein increased with age (Figure 6B,C). We attempted to perform Western blot analysis for RAD51, but unfortunately two sets of generated antibodies were too unspecific to draw any conclusion. These experiments suggest that the contribution of NHEJ to strand break repair becomes more frequent and evident with age. Previously, we indirectly showed that HR repair decreased with age (Boyko et al., 2006b). Since these two repair pathways compete for the same substrate (strand breaks), it makes sense that NHEJ repair has to compensate for decreased involvement of HR.

**DISCUSSION**

Here, we report that older plants have an increased frequency of microsatellite instability as compared to younger plants and propose that this increase is associated with a more frequent contribution of the error-prone NHEJ pathway towards strand break repair.

*Deletions are the predominant type of mutations at the artificial microsatellite loci*

Our analysis showed that the majority of mutations observed at the microsatellite locus of the GUS transgene were deletions; among 54 mutated sequences, there were only 3 insertions and 51 deletions. Thus, deletions occurred up to 16 times more frequently than insertions. When starting the experiments, we assumed that GUS protein activity would be predominantly restored via a deletion of a single guanine or an insertion of two guanines. Whereas the former was a frequent case (24 out of 54 mutated sequences had a single G deleted), the latter was a rare event, with only
two cases being an insertion of two guanines. We were surprised to find clones representing deletions of two and three guanines (these events represented 22 out of 54 sequences with mutations). It is hardly possible that these sequences would lead to the restoration of the GUS+ phenotype as they would not restore the coding frame. Since it was difficult to excise blue sectors only, it is possible that some “non-blue” tissues have been PCR-amplified. Since reversions at one allele of 121A reporter plants would be sufficient to activate the transgene, the second copy could also mutate, not necessarily in the same manner; thus, it is possible that the events representing deletions of two or three guanines could have occurred at the second allele.

Curiously, data exist that show that deletions are indeed more common in the microsatellite regions. An in vitro analysis of replication showed that the frequency of mutations at (A)n and (CA)n microsatellites directly correlated with the number of repeats, and it was higher for single rather than dual nucleotide repeats. The frequency of the mutations observed was as high as 1 in 100 replicated repeats, with the contraction rate for (A)n and (CA)n being 5- and 12-fold higher than the expansion rate, respectively (Shinde et al., 2003). Similarly, the analysis of mutation types at microsatellites in wheat showed that deletions at the microsatellites loci were 4-fold more frequent than insertions (Thuillet et al., 2002).

Age-dependent changes in DNA repair efficiency in different organisms

In our experiments, we observed that the rate of mutations at artificial microsatellite loci of a transgene increases with plant age. It is difficult to draw any parallel with the findings reported in literature because similar studies have not been performed. The effect of aging on DNA repair and mutation rates was primarily studied in seeds. Embryo cells within plant seeds do not divide and exist in the G0/G1 stage of the cell cycle for a long period of time (Whittle et al., 2001). Since it is possible to manipulate the duration of the resting stage in seeds, the assessment of the physiological and genetic impact of cell aging on DNA damage and mutation rates is also possible. Several papers studying the impact of seed age on the onset of mutations support our finding (reviewed in (Whittle and Johnston, 2006)). Naturally aged rye (Secale cereale) seeds have increased variations in AFLPs and other genetic markers (Chwedorzewska et al., 2002a; Chwedorzewska et al., 2002b). Plantlets of Zea mays (Peto et al., 2008), Triticum (Melletti and Russo, 1969) and Crepis (Boldyrev et al., 2004) germinated from older seeds contain higher levels of chromosome rearrangements and other mutations.
However, age-dependent changes in the efficiency of DNA repair in plants have not yet been reported. Analysis of mutation rates in germinated plants is more difficult since cell divisions are not synchronized and plant cells have various degrees of endo-reduplication. But older plants contain higher percentage of older cells. The mutation rate is also more difficult to study in living plants due to the rare occurrence of single nucleotide changes. It was estimated that mutations at a given nucleotide of the transgene occur at the rate of $10^{-8} - 10^{-9}$. Given the fact that a three-week-old Arabidopsis plant of an average size contains about $10^6$ cells, it becomes clear that measuring mutation rates is a difficult process. However, cell genomes contain stretches of simple sequence repeats, microsatellites that mutate at much higher rate (Marriage et al., 2009). It was estimated that mutation rates at microsatellites were $10^3$-$10^4$-fold higher than mutation rates at non-repetitive sequences. Thus, the analysis of mutation rates can be easily done via the analysis of mutations at the microsatellite loci. Indeed, we utilized a method for detection of mutations at the artificial microsatellite loci cloned into the GUS transgene and found that older plants had higher rates of mutations in these regions.

The data on the mutation rate in animals are much more complete and suggest that aging animal cells have a lower efficiency of most types of DNA repair, including excision repair, mismatch repair and strand break repair mechanisms (Gorbunova et al., 2007). Mitchell and Hartman (1990) showed that proliferating tissues had a higher repair capacity as compared to terminally differentiated non-dividing cells (Mitchell and Hartman, 1990). More specifically, it was found that senescent cells in G0/G1 do not use HR for strand break repair, and thus they more often rely on the use of the NHEJ pathway (Seluanov et al., 2007). Analysis of excision repair capacity also showed age dependence. An *in vitro* assay measuring repair efficiency of a synthetic DNA substrate containing a single G:U mismatch detected a strong decline in BER activity in brain, liver and germ cell nuclear extracts of old mice (Intano et al., 2003). Also, another large study, using human peripheral blood lymphocytes from over a hundred individuals aged 20 to 60, showed an age-dependent decline of UV damage repair (Wei et al., 1993; Grossman and Wei, 1995). The rate of decline was 0.63% per year, which amounts to a 25% decrease over a 40-year period (Wei et al., 1993; Grossman and Wei, 1995).
Higher rates of microsatellite instability in plants are not due to lower efficiency of replicative DNA polymerases

DNA replication is a precise process, with DNA replicative polymerases incorporating incorrect bases at a rate of $10^{-6}$-$10^{-7}$ per base pair replicated. MMR fixes most of these errors reducing an error rate additionally by $\sim 1,000$-fold. In repetitive tracts that are prone to slippage and mispairing during replication (Kunkel and Bebenek, 2000), potential frameshift mutations including deletions or insertions are more frequent.

Our experiments showed that 21-day-old plants had lower DNA polymerase activity but higher DNA polymerase fidelity. These data were paralleled with a decrease in the expression of all tested replicative polymerases. An age-dependent decrease observed in the expression pattern of replicative DNA polymerases in plants is a novel finding. Similar results have been reported in animals; several reports showed reduced abundance of the replicative DNA polymerase $\beta$ in brain extracts from older mice and rats (Krishna et al., 2005). For example, the expression of DNA polymerase $\beta$ and AP endonuclease was induced by DNA damage in young mice, while older mice showed a lack of inducibility (Cabello et al., 2002).

Unfortunately, no data exist on age-dependent changes in DNA polymerase fidelity in plants. In contrast, several previous reports suggest the existence of lower DNA polymerase activity in older plants (Bottomley, 1970) and aging seeds (Grilli et al., 1995; Reuzeau et al., 1997), thus confirming our findings.

Our finding of the higher fidelity of DNA polymerases in extracts from old plants is puzzling. Since we observed that both RNA levels of major DNA replicative polymerases and the total activity of DNA polymerases dropped with age, it would be logic to expect a drop in the fidelity. It seems so that an increase in fidelity could be some kind of compensatory mechanism for lower abundance and activity. We can only hypothesize that higher mutation rates in microsatellites in older plants are not due to the decreased fidelity of replicative DNA polymerases.

The argument for the role of the replication process in the accumulation of heritable mutations is a long-lasting one. The notion that replication errors are main contributors to heritable mutations in animals seems to compelling (Crow, 2000). In contrast, analysis of evolutionary rates of neutral DNA sequences among various plant taxa suggests that more heritable base substitution mutations occur per unit time during seed ageing than during the lifetime of the plant. This suggests that higher percentage of heritable mutations are the result of accumulated DNA damages in seeds.
rather than the result of replication errors occurring upon multiple cell divisions in meristematic regions of growing plants (Whittle, 2006).

Changes in mismatch repair may be responsible for an age-dependent increase in microsatellite instability

Mismatch repair is one of the first DNA repair mechanisms taking care of replicative errors. We thus hypothesized that mistakes in mismatch repair could be more frequent in older plants than in younger plants. As anticipated, our analysis showed that microsatellite mutations occur more frequently in msh2 plants than in wild-type plants. Alou et al. also reported that knockout of AtPMS1 showed up to a 28-fold increase in microsatellite instability (Alou et al., 2004). The analysis of the number of blue spots showed that an age-dependent increase in mutation rates was similar in msh2 and wild-type plants. We should admit, however, that since the number of events in msh2 plants was so high, it was difficult to make a comprehensive analysis.

The analysis of steady-state RNA levels showed a drastic age-dependent decrease in the expression of MSH2 and MSH6 genes, and a tendency for increased expression of the MSH7 gene. Despite phylogenetic evidence that MSH7 arose from MSH6, the mismatch recognition properties of AtMSH2-MSH6 and AtMSH2-MSH7 appeared to differ sharply. The AtMSH2-MSH7 dimer bound poorly to DNA containing a single extra ‘looped out’ nucleotide, whereas this substrate was effectively recognized by AtMSH2-MSH6 (Wu et al., 2003). In contrast, both pairs recognized the G/T mismatch equally well. A decrease in the expression of MSH2 and MSH6 but not MSH7 could possibly suggest that small loops are less efficiently repaired in older plants as compared to younger ones, whereas the repair of mismatches might have similar efficiency. The replication of microsatellites such as G16 frequently results in polymerase slippage leading to single nucleotide deletions or insertions, thus resulting in the formation of single nucleotide loops. Since AtMSH2-MSH6 is responsible for the repair of such loops, it is possible that an age-dependent decrease in the expression of MSH2 and MSH6 could contribute to an age-dependent increase in microsatellite instability. However, since there is no in vitro DNA repair assay for the analysis of mismatch repair, it is hard to draw a clear conclusion.

An age-dependent increase in microsatellite instability could be due to more frequent involvement of NHEJ

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Our repair assay showed that older plants have more efficient NHEJ repair as compared to younger plants. At the same time, older plants appeared to be sloppier in end-joining repair as compared to younger plants. These changes were paralleled by the higher steady-state level of Ku70 RNA and the amount of KU70 protein. These results suggest that older plants indeed use NHEJ repair more often. Since NHEJ is a more error-prone mechanism of end-joining repair as compared to HR repair (Gorbunova and Levy, 1997; Bleuyard et al., 2006), it can be hypothesized that an increase in the involvement of NHEJ repair could be one of the reasons explaining why microsatellite mutations occur in older plants more often.

Our previous work also suggested that older plants utilize NHEJ more frequently. Previously, we found that homologous recombination rates in plants harvested at 2 dpg decreased by 6-fold compared to plants harvested at 22 dpg (Boyko et al., 2006b). We noticed that whereas the expression of genes involved in recombination repair decreased with age, the expression of genes involved in NHEJ repair increased (Boyko et al., 2006b).

Microsatellites are often subject to frequent unequal crossing-over. Unequal crossing-over between direct repeats involves misaligned pairing of repeats on homologous chromosomes (Lovett, 2004). It can be hypothesized that a decrease in the involvement of HR could be one of the protection mechanisms preventing frequent rearrangements/crossing-over that could contribute to microsatellite instability.

Unfortunately, not much is known about age-dependent changes in strand break repair in plants. In animals, the efficiency of NHEJ was reduced over four-fold in presenescent and senescent cells relative to young cells, indicating that DNA end joining becomes less efficient and more error-prone in senescent cells (Seluanov et al., 2004). The level of Ku70 has been shown to decline markedly in the testes of aging rats (Um et al., 2003) and lymphocytes from aging human donors (Ju et al., 2006). Also, in a recent publication, a 50% reduction in the level of KU70 and KU80 proteins in senescent human fibroblasts was observed (Seluanov et al., 2007). Analysis of NHEJ in the brains of young and old rats using the in vitro plasmid rejoining assay showed a substantial reduction in the efficiency of plasmid rejoining in the brains of old rats (Ren and Pena de Ortiz, 2002; Vyjayanti and Rao, 2006). A similar experiment described in this paper also showed that end-joining repair was more error-prone in older plants. At the same time, we found that older plants use NHEJ more often than younger plants.
CONCLUSION
Our experiments showed that microsatellite instability in plants increased with age. Further analysis showed that there was a correlation between the more frequent involvement of NHEJ and higher microsatellite instability in older plants. Since we observed a decrease in the expression of genes coding for MMR, it cannot be excluded that a lower capacity of MMR contributes to higher mutation rates in older plants. It remains to be shown whether mutants impaired in NHEJ would be impaired in an age-dependent increase in microsatellite mutation rates.
MATERIALS and METHODS

Plant lines and growing conditions
Production of the Arabidopsis thaliana microsatellite instability reporter line 121A has been previously described (Azaiez et al., 2006). Briefly, this line contains a synthetic microsatellite (16 Gs) cloned into the uidA gene’s MscI restriction site, 620 bp downstream of the start codon. This generates a premature stop codon at the position of 8 nucleotides downstream of the cloned fragment. The loss of a single nucleotide or the gain of two nucleotides at the microsatellite locus will provide the correct frameshift to activate the GUS gene (Figure 1). Seeds from line 121A were sowed on MS media (Sigma) plates and grown under controlled light and temperature conditions (16 hours/light at 22 °C and 8 hours/dark at 18 °C with illumination of 200 µmol m⁻² s⁻¹). Plants were harvested on days 3, 5, 7, 10, 13, 16, 19, and 22 post-germination. For the analysis of the role of MMR, line 121A was crossed to the msh2 mutant (Azaiez et al., 2006), homozygous lines were selected, and the frequency of mutations in the microsatellite region was measured at the aforementioned days post-germination.

Histochemical staining procedures
Histochemical staining of plants was performed at the aforementioned developmental stages as described before (Jefferson et al., 1987). Briefly, the plants were vacuum infiltrated for 10 min in a sterile staining buffer containing 100 mg of 5-bromo-4-chloro-3-indolyl glucuronide (X-glu) substrate (Jersey Labs Inc., USA) in 300 mL of 100 mM phosphate buffer (pH 7.0), 0.05% NaN3, 0.05% Tween 80, and 1 mL dimethylformamide. The plants were then incubated at 37°C for 48 hours in the staining buffer and then bleached with ethanol (Figure 1B).

Calculating the number of genomes
Total DNA from respective transgenic lines was isolated from whole plants at different developmental stages using the Nucleon phytopure plant DNA extraction kit (Amersham Life Science, Piscataway, New Jersey, USA). To estimate the number of genomes present, the total DNA yield (µg/plant) was compared with the DNA content of an Arabidopsis cell (0.16 pg) (Swoboda et al., 1993).

To find out whether the DNA extraction method had a significant influence on yield, we prepared DNA using an additional previously published protocol (Boyko et al., 2006a; Boyko et
al., 2006b). Although the DNA yield was increased by 50% as compared to the yield produced by the Nucleon Phytopure Kit, the ratio between the amounts of DNA in different plant organs was the same.

**Preparation of plant crude tissue extracts for the analysis of DNA polymerase activity**

Leaves from 5-, 12- to 21-day-old plants of line 121A were used for preparation of crude extracts. All extraction steps were carried out on ice. Using a pestle and mortar, 10 g of leaves were homogenized with 100 ml of pre-chilled homogenization buffer (100 mM sodium phosphate buffer, pH 7.4, containing 1% v/v protease inhibitor cocktail from Sigma, #P9599-5ML) to obtain a 10% extract. The extracts were filtered through four layers of cheesecloth and centrifuged at 10,000 rpm for 15 min. The supernatant obtained was immediately used for assays. If necessary, crude plant extracts can be partially purified using any suitable method (for instance, the one described by Li et al. (Li et al., 2002)). For the analysis of total protein concentration in crude plant tissue extracts, the Bradford assay with bovine serum albumin as a standard was applied (Bradford, 1976). The experiments were repeated three times.

**Preparation of the primer/template complex**

The primer/template complex for the assay was prepared by annealing the fluorescein amidite (FAM)-labelled 15 bp primer (5’-6-FAM-TCCCAGTCACGACGT-3’, PAGE-purified) to the 30 bp template (5’-TCATCGAGCATGATCACGTCGTGACTGGGA-3’, PAGE-purified). All components were mixed by pipetting in the following order (on ice): \( \Sigma V = 200 \mu l : \) Tris-HCl (1M, pH 8.0) = 10 \( \mu l \), \( \beta \)-Mercaptoethanol (14.3M) = 0.5 \( \mu l \), BSA (10 mg/ml, NEB) = 2 \( \mu l \), Primer (100\( \mu \)M) = 3 \( \mu l \), Template (100\( \mu \)M) = 3 \( \mu l \), \( H_2O \) = 183.5 \( \mu l \). The reaction was incubated for 5 min in boiling water and then allowed to cool to room temperature (20-25°C). The complex was prepared in advance and stored at -20°C.

**Polymerisation reaction**

The crude extract (10 \( \mu g \) of total protein) was mixed with the primer/template complex, 2 mM dNTP and reaction buffer (10x Y+/Tango, Fermentas), 10 mM magnesium acetate, 66 mM
potassium acetate, 0.1 mg/ml BSA: ΣV = 25 µl, dNTP (2 mM) = 2.5 µl, Template/Primer complex = 2.5 µl, Reaction Buffer = 2.5 µl, Plant extract = 10 µg of total protein, H₂O = to 25 µl.

The reaction was carried out at 37 ºC for 15 min in a PCR machine, quenched with 50 µl of loading buffer (95% formamide, 50 mM EDTA, and 0.05% bromphenol blue), heated to 95°C for 3 min and cooled on ice for 2 min. The Klenow enzyme (NEB buffer 2) was used as a positive control.

Denaturing polyacrylamide gel electrophoresis and gel scanning

After quenching, 15 µl of the sample/loading buffer mix was loaded on a 20% polyacrylamide gel (20x20x0.075 cm) containing 8 M Urea. Also, 0.5 µl of the 15 bp 6-FAM-labelled primer was mixed with 2.5 µl of loading buffer and loaded on the polyacrylamide gel to serve as a molecular weight marker. Electrophoresis was carried out in 1x TBE buffer for about 5 h at 500 V. Gels were scanned using Typhoon 9410 at an excitation wavelength of 88 nm using a 520 BP 40 emission filter, the PMT voltage of 685 V, at a resolution of 100 µm. Images of scanned gels were analyzed using ImageQuant 5.2 software (Molecular Dynamics). For the adjustment of total values of the object outline, the Local Median background correction method was applied.

Real time PCR analysis

Approximately 80 mg of plant tissue was ground in liquid nitrogen, transferred to a chilled 1.5mL eppendorf tube, and 160 µL TRIzol reagent (Invitrogen) was added. The remainder of the extraction was performed as per the manufacturer’s protocol. RNA quantity and quality were measured using a spectrophotometer (Ultraspec 1100 pro) in 20 mM Tris pH 7.5 in RNase-free double distilled water. cDNA was then prepared from total RNA using the RevertAID H First Strand cDNA synthesis kit (Fermentas).

Real time quantitative PCR (RT-qPCR) was performed using SsoFast EvanGreen Supermix (Bio-Rad). cDNAs were amplified under the following conditions: (i) 98°C for 3 min for one cycle; (ii) 98°C for 5 s, 54.3°C for 5 s, 65-95°C for 5 s, with a 0.5°C increment for 40 cycles. Primers (see Table 3S) for RT-qPCR were designed using the Primer3 program. For every set of primers, annealing temperature optimization, melt curve analysis and gel analysis of amplicon were performed. To evaluate PCR efficiency, the standard curve was established using series of cDNA
dilutions. The expression of polymerases was related to the expression of *RCE1*. Conditions of *RCE1* amplification were as follows: (i) 98°C for 3 min for one cycle; (ii) 98°C for 5 s, 45.0°C for 5 s, 65-95°C for 5 s, with a 0.5°C increment for 46 cycles. The expression of *RCE1* is commonly used for the analysis of expression of developmentally or metabolically regulated genes (Seki et al., 2001). The average of 4 reactions (2 dilutions per each of two RNA preparations) was obtained, and the fold induction was calculated. The statistical significance of the experiment was confirmed by performing a Student’s t-test (two-tailed paired or non-paired).

Partial plant extract purification for the in vitro DNA repair assay
All steps were performed on ice. Frozen plant tissues from 5-, 12- and 21-day-old plants were grounded with liquid nitrogen and resuspended in 7 volumes (w/v) of ice-cold homogenization buffer (25 mM Hpes-KOH, pH 7.8, 100 mM KCl, 5 mM MgCl₂, 250 mM sucrose, 10% glycerol, 1 mM DTT, and the protease inhibitor cocktail Complete Mini (Roche, Mannheim, Germany)). The homogenates were filtered through 45-µm filters and placed on a magnetic stirrer. 2M KCl was added slowly to the stirring homogenates to a final concentration of 450 mM. After 30 minutes of extraction, the homogenates were centrifuged at 40,000 g for 1 h to remove cellular and nuclear debris. The supernatants were transferred to glass beakers, and solid (NH₄)₂SO₄ was added slowly to a final concentration of 70%. Ten µl of 1 M NaOH were added for each 1 g of (NH₄)₂SO₄ to neutralize the supernatant. After 1 h incubation on ice, the precipitated proteins were collected by centrifugation at 20,000 g for 1 h. The supernatants were discarded, and the pellets were dissolved in a minimum volume (3 ml) of a dialysis buffer (25 mM Hpes-KOH, pH 7.8, 100 mM KCl, 12 mM MgCl₂, 1 mM EDTA, 17% glycerol, 2 mM DTT) and dialyzed overnight against the same buffer (in 4 L) using 7,000 MWCO Slide-A-Lyser dialysis cassettes (Thermo Scientific). After dialysis, the samples were split into aliquots, frozen in liquid nitrogen and kept at -80°C.

In vitro repair assay
All reactions were done in triplicates. LITMUS29 (NEB, USA) was used in the experiments. Circular, linearized blunt-ended (*Stu*I, Fermentas) non-damaged and damaged (UV light, *λ*ₘₐₓ = 254 nm, 450 J/m²) as well as linearized non-sticky-ended (BamHI/KpnI, Fermentas) non-damaged and damaged plasmids (UV light, *λ*ₘₐₓ = 254 nm, 450 J/m²) were used for the assay. Each reaction (25 µl total volume) contained one form of plasmid DNA (300 ng/reaction), 2.5 µl 10x reaction
buffer (Roche, Mannheim, Germany), 2.5 µl 10x dNTP/DIG-11-dUTP mix (Roche, Mannheim, Germany), and 10 µg of a partially purified plant extract (Li et al., 2002). Plant extracts were prepared from 5-, 12- and 21-day-old plants (10 plants per each sample). Reaction mixes were incubated in the dark for 2 hours at 25°C. The reactions were stopped with 20 mM EDTA (the final concentration). DNA (120 ng) was separated by electrophoresis in 0.8% agarose/1x TAE, visualized with ethidium bromide, photographed and transferred to a nylon membrane (Roche, Mannheim, Germany) in the 10x SSC solution using a vacuum blotter (Appligene, UK) (see Figure 3S). After transfer, DNA was bound to a membrane by UV light using a Spectrolinker XL-1000 (Spectronics Corp.) and detected with the DIG Nucleic Acid Detection Kit (Roche, Mannheim, Germany) using anti-DIG-alkaline phosphatase conjugate and NBT/BCIP as a substrate according to the manufacturer’s instructions (Roche, Mannheim, Germany). Membranes were scanned and quantified with ImageJ.

Analysis of strand break repair using plant protoplasts

Protoplasts were prepared from leaves of 5-, 12- and 21-day old Arabidopsis thaliana plants and transfected essentially as described (Pelczar et al., 2003). For the analysis, the pGEM (Promega) plasmid carrying the ampicillin resistance gene and the LacZ gene was digested with KpnI to generate 3’ protruding ends. Digested DNA was purified from the gel and dephosphorylated; the absence of a circular plasmid molecule was tested by PCR across the restriction site region (data not shown). Protoplasts of 5-day-old, 12-day-old and 21-day-old plants were transfected with 5 µg of linear DNA. To normalize for equal transfection efficiency, each protoplast sample was also transfected with 0.5 µg of circular pM1-Luc (Roche) plasmid carrying a kanamycin resistance gene.

Materials were added to 15 mL Falcon tubes in the following order: DNA 5.5 µg, 0.3 mL of a suspension containing 1.0 x 10^6 protoplasts/mL and 0.3 mL 40% PEG 6000. The mix was incubated for 5 min at room temperature, and then 4 mL of K3 medium was added. After mixing, the tubes were incubated in a horizontal position at 28°C in the dark. Incubation was performed for 4h and 12h. Harvesting of transfected protoplasts was done using 10 mL of W5 buffer (150mM NaCl, 125mM CaCl₂, 5mM KCl, 6mM glucose) and by centrifugation for 10 min at 1000 rpm. The Qiagen miniprep kit was used to purify the circular form of DNA. This DNA was used for E. coli transformation.
Following transformation, 90% of the bacteria were plated on ampicillin-containing media and the remainder – on kanamycin-containing media supplemented with X-gal/IPTG. The number of amp\textsuperscript{R} and kan\textsuperscript{R} bacterial colonies was counted and compared between different age groups. The number of kan\textsuperscript{R} bacterial colonies was equal in all samples prepared from protoplasts of different ages, suggesting that protoplast transfection steps, DNA recovery steps and bacteria transfection steps were similarly efficient in all three protoplast groups.

*Western immunoblotting*
Western blot analysis of the amount of KU70 was performed from ~100 mg of plant tissue using anti-KU70 primary antibodies (1:500, Santa Cruz Biotechnology) and anti-goat secondary antibodies (1:5000, Santa Cruz Biotechnology), as described before (Boyko et al., 2006b).

*Statistical analysis*
The experiments were repeated at least three times, and the mean values ± SD or SE were calculated. The statistical significance of the results was confirmed by performing Student’s t-test. Statistical analyses were performed using Microcal Origin 6.0.

*Acknowledgements*
We thank Valentina Titova for proof-reading the manuscript. IK acknowledges the financial support of NSERC and Human Frontiers Science Foundation.
Table 1. The steady-state RNA level of genes coding for various DNA polymerases

<table>
<thead>
<tr>
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<th>5d</th>
<th>12d</th>
<th>21d</th>
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<tbody>
<tr>
<td><strong>Pol α</strong></td>
<td>1±0.064</td>
<td>0.24±0.006***</td>
<td>0.21±0.010***</td>
</tr>
<tr>
<td><strong>Pol δ</strong></td>
<td>1±0.062</td>
<td>0.64±0.034**</td>
<td>0.69±0.070*</td>
</tr>
<tr>
<td>PolI like-A</td>
<td>1±0.073</td>
<td>1.04±0.062</td>
<td>0.56±0.182*</td>
</tr>
<tr>
<td>PolI like-B</td>
<td>1±0.080</td>
<td>0.66±0.038**</td>
<td>0.69±0.029**</td>
</tr>
<tr>
<td>AtREV1</td>
<td>1±0.033</td>
<td>2.34±0.066***</td>
<td>2.47±0.058***</td>
</tr>
<tr>
<td>AtREV3</td>
<td>1±0.052</td>
<td>0.77±0.161</td>
<td>1.20±0.062*</td>
</tr>
<tr>
<td><strong>Pol κ</strong></td>
<td>1±0.081</td>
<td>0.69±0.089*</td>
<td>0.66±0.074*</td>
</tr>
<tr>
<td><strong>Pol η</strong></td>
<td>1±0.111</td>
<td>0.95±0.097</td>
<td>0.94±0.045</td>
</tr>
<tr>
<td><strong>Pol θ</strong></td>
<td>1±0.078</td>
<td>2.84±0.182***</td>
<td>2.02±0.080***</td>
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<tr>
<td><strong>Pol λ</strong></td>
<td>1±0.102</td>
<td>0.44±0.290*</td>
<td>0.18±0.034***</td>
</tr>
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</table>

The real-time PCR data for each DNA polymerase were related to the RCE1 data and then standardized to the data from 5-day-old plants (taken as 1.0). The data represent the average of 3 independent experiments with standard deviation. The asterisks show a significant difference as compared to the data from 5-day-old plants. One asterisk, $P<0.05$; two asterisks, $P<0.01$; three asterisks, $P<0.001$. 
Table 2. The steady-state RNA level of genes coding for mismatch repair proteins

<table>
<thead>
<tr>
<th></th>
<th>5d</th>
<th>12d</th>
<th>21d</th>
</tr>
</thead>
<tbody>
<tr>
<td>MSH2</td>
<td>1±0.082</td>
<td>0.43±0.023***</td>
<td>0.41±0.017***</td>
</tr>
<tr>
<td>MSH6</td>
<td>1±0.069</td>
<td>1.10±0.015</td>
<td>1.16±0.057*</td>
</tr>
<tr>
<td>MSH7</td>
<td>1±0.147</td>
<td>0.23±0.092**</td>
<td>0.33±0.023***</td>
</tr>
</tbody>
</table>

The real-time PCR data for each gene were related to the RCE1 data and then standardized to the data from 5-day-old plants (taken as 1.0). The data represent the average of 3 independent experiments with standard deviation. The asterisks show a significant difference as compared to the data from 5-day-old plants. One asterisk, $P<0.05$; two asterisks, $P<0.01$; three asterisks, $P<0.001$. 
Literature Cited


Viguera E, Canceill D, Ehrlich SD (2001) Replication slippage involves DNA polymerase pausing and dissociation. EMBO J 20: 2587-2595


Figure captures

Figure 1. Schematic presentation of the construct for the analysis of microsatellite instability
A. Insertion of the sequence (in red) carrying the G16 microsatellite results in the frameshift leading to in-frame stop codon couple nucleotides downstream. Deletion of one guanine or insertion of two guanines would restore the open reading frame of the GUS gene and thus activate the protein.
B. Cells and their progeny in which reading-frame restoration occurred can be visualized after histochemical staining as blue sectors. The picture shows leaves of 21-day-old plants with a number of blue sectors. The insert shows the entire plant.
C. Analysis of the types of reversions occurring at the microsatellite region of the GUS transgene. The numbers show the frequency of occurrence of a particular sequence. A template represents the sequence of the plasmid used to produce line 121A. “WT” identifies a wild-type sequence, whereas G\( \rightarrow \)A/1G-, 1G-, 2G-, 3G-, 4G-, 5G-, 1G+ and 2G+ represent mutations. The bottom of the figure shows representative chromatograms for the wild-type sequence (G16) and deletion of 4 guanines (G12).

Figure 2. Mutation frequency and mutation rates at microsatellite loci increase with age
Seeds of transgenic Arabidopsis thaliana plants were germinated and grown on MS media. Plants were harvested for histochemical staining at 3, 5, 7, 10, 13, 16, 19 and 22 days post germination.
A. The graph shows the average mutation frequency (as calculated from three independent experiments), and error bars indicate a standard error. Mutation frequency was calculated as the average number of mutation events (blue spots) per the number of plants used for the analysis. The values that are not connected by the same letter are significantly different (Student’s test, \( \alpha = 0.05 \)).
B. The graph shows the average mutation rate (as calculated from three independent experiments), and error bars indicate a standard error. The mutation rate was calculated by relating mutation frequency to the total number of genomes present in plants of different ages. The values that are not connected by the same letter are significantly different (Student’s test, \( \alpha = 0.05 \)).
Figure 3. Analysis of DNA polymerase activity and fidelity in plants of different ages

A. Extracts from 5-, 12- and 21-day-old *Arabidopsis thaliana* plants (ecotype Columbia) were incubated in triplicates with 2 mM dGTP+dATP. The 3-, 15-, 17- and 18-nt bands are indicated by arrows. K – the Klenow fragment reaction products, P – the 15 nt 6-FAM labelled primer only. 5 dpg, 12 dpg and 21 dpg indicate samples prepared from plants collected at 5, 12 and 21 days post germination, respectively.

B. The average intensity (in arbitrary units ± SD) of the 17-nt band. The numbers on the top of bars show the average intensity. Asterisks show significant differences in 12- and 21-day-old plants as compared to 5-day-old plants (two asterisks, *P*<0.01; three asterisks, *P*<0.001).

C. The average intensity (in arbitrary units ± SD) of the 18-nt band. The numbers on the top of bars show the average intensity. Asterisks show significant differences in 12- and 21-day-old plants as compared to 5-day-old plants (one asterisk, *P*<0.05; three asterisks, *P*<0.001).

D. The average intensity (in arbitrary units ± SD) of the 3-nt band. The numbers on the top of bars show the average intensity. Asterisks show significant (*P*<0.001 in both cases) differences in 12- and 21-day-old plants as compared to 5-day-old plants.

E. DNA polymerase activity (fold) is calculated by relating the intensity of the 17-nt band in 12- and 21-day-old plants to the intensity of this band in 5-day-old plants. Bars show the average of three experiments with SD. Asterisks show significant differences in 12- and 21-day-old plants as compared to 5-day-old plants (two asterisks, *P*<0.01; three asterisks, *P*<0.001).

F. DNA polymerase fidelity (fold) is calculated by relating the intensity of the 18-nt band (as shown in Figure 3C) to the intensity of the 17-nt band (as shown in Figure 3B). The data for 12- and 21-day-old plants are shown as a fold change to 5-day-old plants. Bars show the average of three experiments with SD. Asterisks show significant differences in 21-day-old plants as compared to 5-day-old plants (*P*<0.001).

G. Exo- (endo) nuclease activity (fold) is calculated by relating the intensity of 3-nt band in 12- and 21-day-old plants to the intensity of this band in 5-day-old plants. Bars show the
average of three experiments with SD. Asterisks show significant ($P<0.001$ in both cases) differences in 12- and 21-day-old plants as compared to 5-day-old plants.

**Figure 4. The *in vitro* DNA repair assay shows a lower excision repair capacity of older plants**

The DNA repair assay was performed by incubation of circular and linear non-exposed and UV-irradiated DNA with partially purified protein extracts prepared from 5-, 12- and 21-day-old plants.

A. The figure shows the representative gel. “Circular” shows the band stemming from incorporation of dig-labeled dNTPs into circular DNA incubated with protein extracts. “Blunt”, “blunt-UV”, “non-sticky” and “non-sticky-UV” show the bands stemming from incorporation of dNTPs into blunt-cut DNA, blunt-cut-UV irradiated DNA, non-sticky-end-cut DNA, and non-sticky DNA irradiated with UV.

B. The figure shows units of the average band intensity as seen in Figure 4A prorated to the loading controls (Figure 3S). The data are shown as the average (calculated from three independent experiments with SD).

**Figure 5. The fidelity of strand break repair is lower in protoplasts prepared from older plants**

Protoplasts prepared from leaves of 5-, 12- and 21-day-old plants were transfected with linear plasmid DNA. 4 and 12 hours post transfection, bacterial cells were transformed with repaired circular DNA, and the number of white and blue colonies was calculated.
A. The figure shows the average number of colonies (blue and white) as counted from three independent experiments (with SD). The asterisk shows significant differences ($P<0.05$) in 12- and 21-day-old plants as compared to 5-day-old plants.

B. The average (with SD) percentage of white colonies out of all colonies was calculated from three independent experiments. The asterisk shows significant differences ($P<0.05$) in 12- and 21-day-old plants as compared to 5-day-old plants.

**Figure 6. Older plants have higher levels of Ku70 expression and lower levels of Rad51 expression**

A. cDNA prepared from 3-, 5-, 7-, 10-, 13-, 16-, 19- and 22-day-old plants was used for amplification of $KU70$ and $RAD51$. Amplification of tubulin was used as a control. Three independent PCR reactions using cDNA prepared from three biological repeats were performed. The figure shows the average intensity (in arbitrary units ± SD) of amplified fragments as related to the intensity of tubulin calculated from three independent experiments. Lines show trend lines for $KU70$ (dashed) and $RAD51$ (solid).

B. Western blot analysis of the amount of KU70 was performed from protein extracts prepared from plant of different ages. The picture shows the representative blot. Silver staining of the gel shows equal loading.

C. The figure shows the average expression (in arbitrary units with SD) as quantified from three independent Western blots and prorated to the loading controls. Asterisks show significant differences in 5- to 22-day-old plants as compared to 3-day-old plants (one asterisk, $P<0.05$; two asterisks, $P<0.01$; three asterisks, $P<0.001$).
A. Insertion of the sequence (in red) carrying the G16 microsatellite results in the frameshift leading to in-frame stop codon couple nucleotides downstream. Deletion of one guanine or insertion of two guanines would restore the open reading frame of the GUS gene and thus activate the protein.

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B. The average intensity (in arbitrary units ± SD) of the 17-nt band. The numbers on the top of bars show the average intensity. Asterisks show significant differences in 12- and 21-day-old plants as compared to 5-day-old plants (two asterisks, $P<0.01$; three asterisks, $P<0.001$).

C. The average intensity (in arbitrary units ± SD) of the 18-nt band. The numbers on the top of bars show the average intensity. Asterisks show significant differences in 12- and 21-day-old plants as compared to 5-day-old plants (one asterisk, $P<0.05$; three asterisks, $P<0.001$).

D. The average intensity (in arbitrary units ± SD) of the 3-nt band. The numbers on the top of bars show the average intensity. Asterisks show significant ($P<0.001$ in both cases) differences in 12- and 21-day-old plants as compared to 5-day-old plants.

E. Polymerase activity (fold) is calculated by relating the intensity of the 17-nt band in 12- and 21-day-old plants to the intensity of this band in 5-day-old plants. Bars show the average of three experiments with SD. Asterisks show significant differences in 12- and 21-day-old plants as compared to 5-day-old plants (two asterisks, $P<0.01$; three asterisks, $P<0.001$).

F. Polymerase fidelity (fold) is calculated by relating the intensity of the 18-nt band (as shown in Figure 3C) to the intensity of the 17-nt band (as shown in Figure 3B). The data for 12- and 21-day-old plants are shown as a fold change to 5-day-old plants. Bars show the average of three experiments with SD. Asterisks show significant differences in 21-day-old plants as compared to 5-day-old plants ($P<0.001$).

G. Exo- (endo) nuclease activity (fold) is calculated by relating the intensity of the 3-nt band in 12- and 21-day-old plants to the intensity of this band in 5-day-old plants. Bars show the average of three experiments with SD. Asterisks show significant ($P<0.001$ in both cases) differences in 12- and 21-day-old plants as compared to 5-day-old plants.
Figure 4. The in vitro DNA repair assay shows a lower excision repair capacity of older plants
The DNA repair assay was performed by incubation of circular and linear non-exposed and UV-irradiated DNA with partially purified protein extracts prepared from 5-, 12- and 21-day-old plants.

A. The figure shows the representative gel. “Circular” shows the band stemming from incorporation of dig-labeled dNTPs into circular DNA incubated with protein extracts. “Blunt”, “blunt-UV”, “non-sticky” and “non-sticky-UV” show the bands stemming from incorporation of dNTPs into blunt-cut DNA, blunt-cut-UV irradiated DNA, non-sticky-end-cut DNA, and non-sticky DNA irradiated with UV.

B. The figure shows units of the average band intensity in Figure 7A. The data are shown as the average (calculated from three independent experiments with SD).
Figure 5. The fidelity of strand break repair is lower in protoplasts prepared from older plants

Protoplasts prepared from leaves of 5-, 12- and 21-day-old plants were transfected with linear plasmid DNA. 4 and 12 hours post transfection, bacterial cells were transformed with repaired circular DNA, and the number of white and blue colonies was calculated.

A. The figure shows the average number of colonies (blue and white) as counted from three independent experiments (with SD). The asterisk shows significant differences ($P<0.05$) in 12- and 21-day-old plants as compared to 5-day-old plants.

B. The average (with SD) percentage of white colonies out of all colonies was calculated from three independent experiments. The asterisk shows significant differences ($P<0.05$) in 12- and 21-day-old plants as compared to 5-day-old plants.
Figure 6. Older plants have higher levels of Ku70 expression and lower levels of Rad51 expression

A. cDNA prepared from 3-, 5-, 7-, 10-, 13-, 16-, 19- and 22-day-old plants was used for amplification of Ku70 and Rad51. Amplification of tubulin was used as a control. Three independent PCR reactions using cDNA prepared from three biological repeats were performed. The figure shows the average intensity (in arbitrary units ± SD) of amplified fragments as related to the intensity of tubulin calculated from three independent experiments. Lines show trend lines for Ku70 (dashed) and Rad51 (solid).

B. Western blot analysis the amount of Ku70 was performed from protein extracts prepared from plant of different ages. The picture shows the representative blot. Silver staining of the gel shows equal loading.

C. The figure shows the average expression (in arbitrary units with SD) as quantified from three independent Western blots. Asterisks show significant differences in 5- to 22-day-old plants as compared to 3-day-old plants (one asterisk, $P<0.05$; two asterisks, $P<0.01$; three asterisks, $P<0.001$).