DNA polymerase λ - a novel DNA repair enzyme in higher plant genome

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Abstract

DNA polymerase lambda (Pol λ) is a novel family X DNA polymerase which has been shown to play key role in nuclear DNA repair and meiotic recombination. Recent studies in mammals support function of this enzyme in base excision repair in response to oxidative DNA damage. However, in plants the biological function of Pol λ in oxidative stress response is still largely unknown. This review will survey recent advances in our understanding of role of Pol λ in repair of oxidative DNA damages and its possible potential involvement in plant’s base excision repair (BER) pathway in response to oxidative stress.

Keywords: Arabidopsis thaliana; oxidant stress; plant growth.

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Oxidative stress and reactive oxygen species

Oxidative damage is one of the major stresses in plants (Hodges, Andrews, Johnson, & Hamilton, 1997), which induces the production of reactive oxygen species (ROS) and also other highly reactive free radicals (Wise, & Naylor, 1987). The reactive oxygen species, generated either endogenously by oxidative electron transport in mitochondria and chloroplast, or exogenously in response to abiotic and biotic stresses can affect almost all the cellular components and induces multiple forms of DNA damages, and thus activates genotoxic stress. ROS react with DNA and generate oxidized form of bases, some of which have extremely mutagenic effects. In mammals, 7,8-dihydro-8-oxoguanine (8-oxo-G) and 2,3-dihydro-2-oxoadenine (2-OH-A) are most commonly generated oxidized bases when ROS attack DNA (Collins, 1999). Occurrence of 8-oxo-G in the replicating strand does not significantly affect DNA replication however it promotes mis-incorporation of nucleotides such as high frequency of incorporation of A against 8-oxo-G. In contrast, existence of 2-OH-A on the replicating strand during replication phase strongly block the progression of the replication fork by the replicative DNA polymerases, resulting in single-stranded (ss) DNA due to separation.
between DNA unwinding and synthesis. ssDNA acts as a common DNA-damage intermediate recognized by complexes associated with ATM (ataxia telangiectasia mutated) and ATR (ATM and Rad3-related) protein kinases. ATR kinases, in particular activates cell-cycle check-point function by phosphorylation of the downstream target Chk1 (check point kinase) which subsequently inhibits activation of origin and blocks the S to G2 phase transition, delaying cell cycle progression (Barone et al., 2007).

The Base Excision Repair pathway removes oxidized bases from DNA

In human cells, the oxidized bases in the DNA are mainly repaired via base excision repair (BER) which involves the removal of the damaged base by a specific glycosylase, producing an abasic site, which is subsequently processed by the step-wise action of the APE1 endonuclease (Apurinic/apyrimidinic endonuclease), Pol β and XRCC1-DNA ligase 1 complex to finally seal the nick (Singhal, Prasad, & Wilson, 1995). However, prolonged replication stress due to inefficient BER activity induces the formation of other potentially harmful DNA intermediates including double strand breaks, in addition to single strand breaks. In human cells, a specialized repair pathway called trans lesion synthesis (TLS) has been shown to be activated in response to prolonged replication block via checkpoint activation which recruits specialized DNA Pols, mainly members of family X and Y polymerases which are capable to bypass the lesion to resume replication. On the other hand the double stand breaks (DSBs) generated by stalled replication fork or delayed BER, are repaired via homologous recombination (HR) (during the S phase of cell cycle only) and non-homologous end joining (NHEJ) mechanisms.

DNA polymerase λ - a novel X family DNA polymerase

DNA polymerase λ (Pol λ) is a family X member and is widespread among higher eukaryotes, both in animals and plants. Mammalian Pol λ is an exonuclease-deficient, 575-amino acid single polypeptide DNA polymerase (García-Díaz et al., 2000, 2002; Ramadan, Shevelev, & Hübscher, 2004) which has been found to share high degree of sequence homology with mammalian Pol β (32%
a number of studies have demonstrated the involvement of Pol λ in all the three major repair pathways including BER (Garcia-Ortiz, Ariza, & Roldan-Arjona, 2001), NHEJ (Lee et al., 2004), and TLS (Picher & Blanco, 2007; Maga et al., 2007; Hübscher & Maga, 2011). Earlier studies have established a role of mammalian Pol λ in TLS for error-free bypass of 2-OH-A and 8-oxo-G lesions in response to oxidative DNA damages (Picher & Blanco, 2007; Crespan, Hübscher, & Maga, 2007). More recent studies have indicated that knockdown of Pol λ ceases replication fork progression and stimulates the ATR/Chk1-mediated S phase checkpoint function, resulting in delay of S phase progression in various human cancer cell lines. Interestingly, knockdown of Pol β, the closest homologue of mammalian Pol λ, failed to activate such response. Together, these observations indicate direct functional involvement of Pol λ in cellular response to DNA damage in the context of oxidative DNA damage and replication stress (Zucca et al., 2013).

**Pol λ interacts with key cell cycle regulatory components, repair and replication associated proteins**

Multiple lines of evidence have demonstrated interactions of Pol λ with the key cell cycle regulatory proteins. *In vitro* and *in vivo* studies established interaction of human Pol λ with cyclin-dependent kinase 2 (CDK2), a key component for G1/S transition (CDK2/cyclin E) and S-phase progression (CDK2/cyclin A) of the cell cycle. Phosphorylation at Thr 553 in human Pol λ was found to be crucial for maintaining Pol λ stability during cell cycle progression in the late S and G2 phases, preventing it from being degraded via proteasomal pathway by ubiquitination and allowing the time for repair during and after S phases (Wimmer, Ferrari, Hunziker, & Hübscher, 2008). Recent studies have established the functional and physical interaction of human Pol λ with MutYH (MutY glycosylase homologue) in a mismatch repair pathway for the error-free bypass of 8-oxo-G lesions in the DNA. The interaction was found to be regulated by a delicately balanced phosphorylation and ubiquitination of Pol λ, governing the stabilization of protein in late S phase and its recruitment to chromatin into active 8-oxo-G repair complexes. Interestingly, Pol λ not involved in 8-oxo-G repair, have been shown to be subjected for proteasomal degradation (Markkanen, van Loon, Ferrari, Parsons, Dianov, & Hübscher, 2012). More recent research have demonstrated that Pol λ is functionally associated with S phase DNA damage response machinery in mammalian cancer cell lines in order to protect cells from oxidative DNA damage (Zucca et al., 2013).

In plants, although information is still limited on the post-translational regulation of Pol λ, phospho-proteomic analyses have identified several potential phosphorylation sites in the N terminus Ser/Pro rich domain in *Arabidopsis* Pol λ, indicating the possibility of *in vivo* phosphorylation of Pol λ by cdk/cyclin complexes like cdk5 and cdc2 during cell cycle progression (unpublished data). In addition, *Arabidopsis* and rice Pol λ possess a highly conserved PIP box (PCNA binding domain) QK/RL/GLKY/FF. The PIP box was found to regulate the interaction of Pol λ with PCNA2 (proliferating cell nuclear antigen 2) for enhancing the fidelity and efficiency of translesion synthesis in oxidative DNA damage repair in *Arabidopsis* (Amoroso et al., 2011). Furthermore, *in silico* analyses (http://smart.embl-heidelberg.de/) have predicted possible interactions of *Arabidopsis* Pol λ with additional DNA repair responsive proteins including the BER pathway proteins Pol δ and DNA ligase 1 (Roy et al., 2013a). Interestingly, Pol λ contains a highly conserved N terminus BRCT (BRCA1 C terminus) domain which is predominantly found in proteins involved in cell cycle checkpoint functions in response to DNA damage and known to mediate protein-protein interactions. Our recent results in *Arabidopsis* have shown that Pol λ interacts directly with XRCC4 (X-ray repair cross-complementing protein 4) and DNA ligase4 via its N terminus BRCT domain during repair of high salinity and DNA cross linking agent induced DSBs through NHEJ pathway (Roy et al., 2013b). Together, these observations suggest that Pol λ interacts with various crucial regulatory partners, which appear to govern the stability and activity of Pol λ and its recruitment to the site of DNA damage.

**Is Pol λ involved in plant’s BER?**

In comparison to plants, the understanding of the mechanism of BER, one of the major pathways of repair of oxidative DNA damages, is far more advanced in mammals, yeast and *Escherichia coli*. However, studies in plants mainly involving *Arabidopsis* have identified structural and/or functional homologs of most of the BER proteins in plant genome (Britt, 2002; García-Ortiz et al., 2001; Kimura & Sakaguchi, 2006; Roldan-Arjona & Ariza, 2009). In plants, the general knowledge about Pol λ structure and functions is still limited. Genome-wide sequence analyses have identified Pol λ as the only member of family X DNA Pols in plants and therefore, appear to substitute for the function of Pol β to play a key role in nuclear DNA damage repair and recombination. Similar to human Pol λ, *Arabidopsis* Pol λ (AtPolλ) is comprising of two major domains, the N-terminal domain and the C-terminal highly conserved Pol X motif which is the active site for polymerase activity. The N-terminal part contains a nuclear localization signal (NLS), a BRCT (breast cancer susceptibility C-terminus) module, which acts as the site for protein–protein and protein–DNA interactions, and a Ser–Pro-rich domain which acts as a suppressor of DNA polymerase activity and also serves as target site for post-translational modification (Fig. 2A). AtPolλ showed ~39% identity and ~57% similarity in amino acid sequence with human Pol λ. Amino acid sequence
alignment of human Pol λ and AtPolλ showed a high degree of amino acid residue conservation at the Pol X domain (Roy, Roy Choudhury, Singh, & Das, 2011). In addition, as like mammalian Pol λ, the C-terminal Pol X domain comprises of an N-terminal 8 kDa domain, unique to family X Pols and a polymerase domain.

Fig. 2. Structural similarities between human and Arabidopsis Pol λ. A. Schematic representation of the domain structures of Arabidopsis and human Pol λ (AtPolλ and HSPolλ). The amino acid numbers for the proteins and the related sub-domain features are indicated. The GenBank accession numbers are ADM33939 (AtPolλ) and NP_001167555 (human Pol λ), respectively. B. Alignment of human Pol λ and AtPolλ amino acid sequences using the ClustalW program (Thompson, Higgins, & Gibson, 1994). The image was prepared with Espript (Gouet, Courcelle, Stuart, & Metoz, 1999). Residues that are completely conserved are colored red and highlighted with a blue box. The numbering shown above the alignment represents that of Arabidopsis Pol λ.
organized in fingers, palm and the thumb sub-domains which are common to all polymerases (Roy et al., 2013b).

Previous studies have established key role of Pol λ in diverse repair pathways in plant genome, including nucleotide excision repair (NER) for removal of UV-B induced photoproducts (Roy et al., 2011) and error-free translesion DNA synthesis in response to oxidative DNA damage in Arabidopsis (Amoroso et al., 2011). We have characterized the transcriptional regulation of Pol λ gene in Arabidopsis (AtPolλ, At1G10520) under diurnal conditions (Roy, Roy Choudhury, Singh, & Das, 2012). Our recent studies have demonstrated involvement of Pol λ in the repair of high salinity and DNA cross-linking agents induced DSBs via the NHEJ pathway in Arabidopsis (Roy et al., 2013b). Therefore multiple evidences support that Pol λ acts as key player in various crucial DNA repair pathways in plant genome.

During the lifecycle plants undergo considerable periods of quiescence when the plant embryo is maintained within the dormant seed (Sallon et al., 2008). Dehydration during storage and subsequent water imbibitions of dry seeds during germination was found to be associated with high levels of oxidative stress which cause significant damage to the integrity of the genome of seed embryo. This is commonly found when seeds are stored under unfavourable conditions like high temperatures and moisture contents (Kranner, Minibayeva, Beckett, & Seal, 2010). Loss of seed viability in storage conditions is strongly linked with various forms DNA damage in seed embryo including single and double strand breaks and even chromosomal breakages and aberrant chromosomes (Waterworth, Drury, Bray, & West, 2011). However information on the repair of oxidative DNA lesions in seed embryo via BER activity during germination is very limited. Recent studies have shown that Arabidopsis cell extracts contain the enzymatic machinery required for uracil and abasic (AP) sites repair via short patch BER pathway (Cordoba-Canero, Morales-Ruiz, Roldan-Arjona, & Ariza, 2009). Overexpression of a DNA glycosylase/AP lyase, a BER pathway component, was found to enhance seed longevity and abiotic stress tolerance in Arabidopsis (Chen et al., 2012). In mammal, studies with extracts from mouse embryonic fibroblasts have indicated that mammalian Pol λ possesses dRP lyase (5′-deoxyribose phosphate lyase) activity in vitro, suggesting involvement of Pol λ in BER. This activity has been found to be associated with the 8-kDa domain at the C-terminal Pol X region of the protein (García-Díaz, Bebenek, Kunkel, & Blanco, 2001). Previously, biochemical evidences have indicated that rice Pol λ possesses the dRP lyase activity (a key feature for repair DNA polymerases involved in BER), indicating possible role of Pol λ in the gap-filling step in plant’s BER pathway (Uchiyama, Kimura, Yamamoto, Ishibashi, & Sakaguchi, 2004). However, unlike mammalian its counterpart, role of Pol λ in BER pathway has not been well characterized in plants at the structure-function level. Therefore, unveiling the mechanisms of BER at the structural, molecular and genetic level in the context of role of Pol λ will be further interesting for understanding the biological function of this sole member of family X DNA polymerase in relation to plant growth and productivity.

Outlook

Significant progress has been achieved in the last few years in understanding the mechanisms of plant DNA repair and recombination. The main pathways of repair are becoming well characterized. The mechanisms by which plant cells respond to environmental and genotoxic stress, including the key early steps of damage detection, transfer of signal and activation of cell cycle checkpoint functions, have just begun to unveil. Future challenges will be further expanding knowledge on the fundamental DNA repair functions which are active in plants, to provide the way towards interesting biotechnological applications focused at improving various stress tolerance in crop plants. Significant research in this area in future will provide meaningful insight about how plants survive under stresses and possible strategies to improve crop growth and productivity in the context of understanding the long-term impacts of adverse environmental conditions on plant genomes.

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