Review Article

Early Steps in the DNA Base Excision Repair Pathway of a Fission Yeast *Schizosaccharomyces pombe*

**Kyoichiro Kanamitsu and Shogo Ikeda**

*Department of Biochemistry, Faculty of Science, Okayama University of Science, 1-1 Ridai-cho, Kita-ku, Okayama 700-0005, Japan*

Correspondence should be addressed to Shogo Ikeda, ikeda@dbc.ous.ac.jp

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DNA base excision repair (BER) accounts for maintaining genomic integrity by removing damaged bases that are generated endogenously or induced by genotoxic agents. In this paper, we describe the roles of enzymes functioning in the early steps of BER in fission yeast. Although BER is an evolutionarily conserved process, some unique features of the yeast repair pathway were revealed by genetic and biochemical approaches. AP sites generated by monofunctional DNA glycosylases are incised mainly by AP lyase activity of Nth1p, a sole bifunctional glycosylase in yeast, to leave a blocked 3′ end. The major AP endonuclease Apn2p functions predominantly in removing the 3′ block. Finally, a DNA polymerase fills the gap, and a DNA ligase seals the nick (Nth1p-dependent or short patch BER). Apn1p backs up Apn2p. In long patch BER, Rad2p endonuclease removes flap DNA containing a lesion after DNA synthesis. A UV-specific endonuclease Uve1p engages in an alternative pathway by nicking DNA on the 5′ side of oxidative damage. Nucleotide excision repair and homologous recombination are involved in repair of BER intermediates including the AP site and single-strand break with the 3′ block. Other enzymes working in 3′ end processing are also discussed.

1. Introduction

DNA molecules in cells always suffer from chemical decay due to exposure to endogenous and environmental agents [1–3]. Cells die when the damage to DNA obstructs replication and transcription. Moreover, base damage causes mutations, which are responsible for cancer, aging, and the hereditary diseases [3–5]. Base excision repair (BER) is a DNA repair pathway directed mainly at nonbulky lesions, such as, alkylated and oxidized bases, and at some types of mismatched bases that are produced during replication or by deamination [3, 5–7]. The BER pathway is usually initiated by DNA glycosylase that removes damaged bases to leave apurinic/apyrimidinic (AP) sites. The AP sites are further processed by an AP endonuclease that cleaves phosphodiether bonds 5′ to the AP site to leave a 3′ OH and 5′-deoxyribose phosphate (5′-blocked end). Bifunctional DNA glycosylase associated with AP lyase removes damaged bases and cleaves 3′ to the AP site, leaving a 3′-α,β-unsaturated aldehyde (3′-blocked end) and a 5′ phosphate. These blocked ends are subsequently converted into 3′ OH and 5′ phosphate by appropriate enzymes. Finally, a repair DNA polymerase fills the gap, and a DNA ligase seals the nick.

The basic mechanism of BER was first elucidated in *Escherichia coli* [3]. Subsequent studies showed that the process is conserved in eukaryotes including the budding yeast *Saccharomyces cerevisiae* and mammals. Fission yeast (*Schizosaccharomyces pombe*) is an ascomycetous yeast that shares many fundamental cellular properties with higher multicellular eukaryotes [8]. Although *S. pombe* has been used as a prominent model organism, research of the BER pathway in yeast has started late. Completion of the *S. pombe* genome project in 2002 provided a list of the yeast BER machineries including DNA glycosylases and AP endonucleases, which are evolutionarily conserved from bacteria to man [9]. Over the past decade, the BER pathway of *S. pombe* has been fairly well characterized by genetic approaches using many BER-defective mutants constructed by gene targeting. Biochemical properties of BER enzymes were also examined using purified recombinant proteins. In this paper, we describe the roles of the enzymes involved in the early steps of BER in fission yeast with an emphasis on...
2. DNA Glycosylases in S. pombe

DNA glycosylases identified in E. coli, S. cerevisiae, S. pombe, and the human are classified in Table 1 on the basis of their substrate specificities and structural features [3, 7, 10]. Monofunctional DNA glycosylase simply hydrolyzes the N-glycosidic bond to release damaged bases and leaves the AP site. Nth1 and Ogg1 DNA glycosylases are associated with AP lyase activity, which cleaves the AP site by β elimination, leaving a 3′-α,β-unsaturated aldehyde. E. coli Fpg and Nei and human NEILs are other types of bifunctional glycosylases which cleave the AP site by β,δ elimination and generate a 3′-phosphate end. S. pombe has five monofunctional DNA glycosylases (Ung1p, Thp1p, Mag1p, Mag2p, and Myh1p) [11]. Unlike other organisms, Nth1p is a sole DNA glycosylase with AP lyase activity in S. pombe cells. Nth1 DNA glycosylase is β lyase and does not possess δ lyase activity [2, 5, 7].

2.1. Uracil DNA Glycosylases Ung1p and Thp1p Function in Avoidance of Spontaneous Mutation. Deamination of cytosine residues in DNA occurs spontaneously in pH- and temperature-dependent reactions, and results in conversion to uracil [1, 3]. Cytosine deamination is estimated to produce 400 uracil residues per mammalian genome per day [12]. Promutagenic U:G mispair causes C:G to T:A transition mutations, if not repaired prior to DNA replication. Uracil also occurs in DNA through incorporation of dUMP instead of dTMP by DNA polymerase during replication. Deamination of 5-methylcytosine in DNA results in the formation of thymine, and hence T:G mispairs. Uracil residues can be removed through the activity of uracil DNA glycosylase (UDG), which consists of four distinct families in mammalian cells: the UNG family (the major activity of UDG in cells), the MUG (mismatch-specific uracil DNA glycosylase)/TDG (thymine DNA glycosylase) family, the SMUG (single strand-specific monofunctional uracil-DNA glycosylase) family, and the MBD4 (methyl-binding domain 4) family [3, 5, 13]. The UDGs, other than MBD4, form a single protein superfamily with a common α/β-fold structure and must have been evolved from a common ancestor [14].

S. pombe has two UDG genes, ung1 (systematic name SPCC1183.06) and thp1 (SPCC965.05c). The Ung1p protein is highly conserved (51% identity to human UNG) and localizes predominantly to the nucleus [15]. The bacterially expressed protein showed an apparent enzyme activity for uracil-containing DNA, which was inhibited by an inhibitor protein (UGI). Overexpression of ung1 induces a DNA checkpoint-dependent cell cycle delay and causes cell death [15]. Human MUG/TDG was first isolated as an enzyme that excises thymine from T:G mispairs as well as uracil from U:G mispairs [16, 17]. S. pombe Thp1p is a member of the MUG family, and maintains a high level of glycosylase activity towards substrates containing U, 5-fluorouracil, 3,N4-ethenocytosine, and 5-hydroxyuracil [18]. Unlike human enzyme, Thp1p cannot excise thymine residues from the T:G mispair substrate. Since cytosine methylation is undetectable in fission yeast cells, the potential to process this substrate appears to be correlated with the degree of cytosine methylation in the genome of organisms [19]. Thp1p can also remove deaminated purine bases: xanthine and oxanine from guanine and hypoxanthine from adenine [20]. Both ung1 and thp1 mutants showed a moderate mutator phenotype [21]. Double mutation of the genes additively increased the mutation frequency. Moreover, expression of Ung1p and Thp1p suppressed spontaneous mutagenesis in UDG-deficient cells. These results indicate that both proteins play important roles in the prevention of spontaneous mutagenesis of S. pombe cells.

2.2. Two AlkA Homologs Mag1p and Mag2p Function in Removal of Alkylated Base. Alkylating agents such as methyl methanesulfonate (MMS) generate covalent modifications at nitrogen residues of DNA bases, in particular N3-methyladenine (3-meA) and N7-methylguanine (7-meG), which account for 11% and 83% among the alkylation products such as 3-meA and 7-meG spontaneously generate 600 and 4,000 residues a day, respectively [12]. In E. coli, alkylation products such as 3-meA and 7-meG are mainly removed by two structurally different
monofunctional DNA glycosylases, AlkA and TagA [3]. The *S. pombe* genome encodes two *alkA* paralogs *mag1* (SPAPB24D3.04c) and *mag2* (SPBC23G7.11). The amino acid sequences of Mag1p and Mag2p share 44.8% similarity. The *S. pombe* *mag1* gene was cloned by its ability to reverse the MMS-sensitive phenotype of an *E. coli* *alkA*/*tagA* double mutant [22]. The substrate range of Mag1p overproduced in *E. coli* is limited to the main alkylation products, such as, 3-meA, 3-meG, and 7-meG, whereas no significant activity was found toward deamination products, ethenoadducts, or oxidation products [23]. The efficiency of 3-meA and 3-meG removal was 5–10 times slower for Mag1p than for *E. coli* AlkA, whereas the two enzymes remove 7-meG at a similar rate. On the other hand, biochemical analysis of Mag2p has not been performed yet because the recombinant protein expressed in *E. coli* showed no glycosylase activity [23].

Mutant strains *mag1Δ* and *mag2Δ* hardly showed MMS sensitivity [23–25]. The BER mutants *nth1Δ*, *apn2A*, and *rad2A* were sensitive to MMS, while the double mutants *nth1Δ/mag1Δ*, *apn2Δ/mag1Δ*, and *rad2Δ/mag1Δ* restored resistance to MMS [23, 25]. This showed that Mag1p is involved in the initial step of MMS-damaged repair both by the Nth1p-dependent short patch BER pathway and by the Rad2p (flap endonuclease)-dependent long patch BER pathway (Figure 1). Moreover, double mutant *nth1Δ/mag2Δ* was more resistant to MMS than *nth1Δ*, indicating that Mag2p also functions in the removal of alkylated bases [25]. *rad16* encodes an XPF homolog that functions as a DNA repair endonuclease in nucleotide excision repair (NER). The double mutant of *rad16* with *mag1* or *mag2* increased sensitivity to MMS [25]. In addition, the *rad16Δ/mag1Δ/mag2Δ* triple mutant exhibited the highest MMS sensitivity. Expression of Mag1p or Mag2p in the triple mutant restored tolerance to MMS. These results showed that BER and NER could independently repair the alkylation DNA damage. Comparison of the substrate specificity and kinetic parameters of Mag1p and Mag2p will be needed to dissect the precise roles of these redundant enzymes in MMS resistance. *rhp51* encodes a RecA-like protein that functions in homologous recombination (HR). The double mutant of *rhp51* with *mag1* decreased the sensitivity to chronic MMS exposure compared with *rhp51Δ* single mutant [24]. In addition, spontaneous intrachromosomal recombination frequencies increased 3-fold in the *mag1* mutant [23]. These results show that both Mag1p and HR contribute to repair of the alkylation base. Deletion of *mag1* from *nth1*, *rad16*, or *rad2* decreased the recombination frequency, indicating that AP sites generated by Mag1p removal of spontaneous base lesions are substrates of short patch BER, long patch BER, NER, and HR [23].

2.3. *Nth1p Is the Major Contributor for Incision of the AP Site.* An 8-oxoguanine (8-oxoG) produced by the oxidation of the guanine residue in G:C could pair not only with cytosine but also with adenine, and the G:C to T:A mutation occurs during the next stage of replication [3, 5, 7]. In a human cell, about 1,000 residues of 8-oxoG are generated per genome in a day [12]. Bifunctional DNA glycosylases Fpg (MutM) in *E. coli* and OGG1 in budding yeast and mammals can excise 8-oxoG paired with cytosine. Thymine glycol (Tg), which is the oxidation product of thymine, interrupts DNA replication and transcription. In human cells, production of Tg and similar oxidized pyrimidines is estimated to be 500 residues per genome per day [12]. In *E. coli* endonuclease III (Nth) and endonuclease VIII (Nei) excise Tg and oxidized pyrimidines from DNA and subsequently cleave the strand by AP lyase activity [3]. *S. pombe* has no homolog of Fpg, OGG1, or Nei [11].

The *S. pombe* *nth1* gene (SPAC30D11.07) was cloned on the basis of homology to *E. coli* *nth* [26]. Nth1 family proteins have a helix-hairpin-helix motif in the vicinity of the active lysine residue and an iron-sulfur cluster [4Fe-4S] at the C terminal. In addition, a eukaryote’s enzyme has a nuclear localization signal. Indeed, fusion protein of *S. pombe* Nth1p with green fluorescent protein was predominantly localized in the nucleus [27]. Unlike budding yeast Ntg1p and the enzyme in mammals, *S. pombe* Nth1p was not observed in mitochondria.

A recombinant *S. pombe* Nth1p expressed in *E. coli* shows a broad substrate specificity. Mass spectrometry of released bases from oxidized DNA by treatment of Nth1p revealed that the enzyme efficiently excised five pyrimidine-derived lesions: 5-hydroxycytosine, Tg, 5-hydroxy-6-hydrothymine, 5, 6-dihydroxycytosine, and 5-hydroxyuracil [28]. *S. pombe* Nth1p could remove 5-formyluracil and 5-hydroxymethyluracil as efficiently as *E. coli* Nth [29]. Moreover, 8-oxoG in 8-oxoG:G and 8-oxoG:a mismap is also the substrate for *S. pombe* Nth1p. The expression of *S. pombe* Nth1p reduced hydrogen peroxide (H2O2) toxicity and the frequency of spontaneous mutations in the *E. coli* nth1/nei double mutant [29]. Although the *S. pombe* *nth1Δ* is sensitive to MMS, Nth1p did not remove 3-meA and 7-meG [30]. However, methyl-formamidopyrimidine, a cytotoxic lesion generated from 7-meG by opening the imidazole ring, was excised efficiently by Nth1p. Nicking activity to oligonucleotides containing Tg and the AP site virtually disappeared in the extract from *nth1Δ* cells, indicating that these lesions are mainly incised by Nth1p in the yeast cell [27].

An *nth1Δ* strain was tolerant to oxidative damage (H2O2 and menadione) and UV irradiation, but exhibited moderate sensitivity to MMS [27, 30, 31]. In addition, the mutant strain exhibited a more than 6-fold increase in the frequency of interchromosomal recombination [30]. Epistasis analysis of the *nth1* gene versus *rad2*, *rad16*, and *rhp51* showed that MMS damage could be repaired through NER and HR other than the BER pathway [30]. Although a mutant of the major AP endonuclease gene *apn2* showed hypersensitivity to MMS, double mutation of *nth1* and *apn2* became tolerant to MMS [31]. This shows that Nth1p functions upstream of Apn2p in the same pathway; that is, the 3′-blocked end generated by Nth1p is converted to 3′ OH by phosphodiesterase activity of Apn2p (Figure 1). In *S. pombe* cells, Nth1p not only removes the oxidized base as DNA glycosylase, but also incises a large portion of the AP site generated by the action of Mag1p and Mag2p DNA glycosylases. Thus, Nth1p plays a central role during early steps of the BER pathway in the fission yeast [27].
Figure 1: A schematic illustration of the BER pathway in *S. pombe*. In the Nth1p-dependent (or short patch) pathway, repair of the AP site is initiated by incision of Nth1p leaving the 3’-block. Apn2p functions primarily in the removal of the 3’-block, and poorly in the incision of AP sites. Apn1p is a back-up enzyme for 3’-phosphodiesterase activity of Apn2p. Finally, a DNA polymerase (Pol.) fills the gap, and a DNA ligase seals the nick. NER and HR could repair a part of the AP sites and single-strand breaks with 3’-blocks. AP endonuclease incises the AP site leaving 5’-dRP end, which will be removed by 5’-dRP lyase activity of Pol4p to feed into short-patch BER. In the long patch pathway, Rad2p incises the flap DNA containing the lesion after DNA synthesis. Uve1p and Apn1p are possibly involved in the NIR pathway. Details are discussed in the text.

2.4. DNA Glycosylase for Mismatch. An 8-oxoG : A mispair in DNA, which is generated by incorporation of an adenine residue opposite to 8-oxoG, leads to a G : C to T : A mutation [3, 5]. The repair of this mismatch is initiated by excision of an adenine residue by adenine-specific mismatch DNA glycosylase (MutY in *E. coli*). The *mutY* homolog of *S. pombe*, namely, *myh1* (SPAC26A3.02), has been cloned [32]. Expression of Myh1p in the *mutY* mutant of *E. coli* could reduce the spontaneous mutation frequency of the cells. Purified Myh1p recognized A : G and A : 8-oxoG, as well as 2-aminopurine : G and A : 2-aminopurine [32]. In addition, Myh1p probably prevents a C : G to G : C transversion mutation because it can remove the guanine from G : 8-oxoG [33]. The *myh1Δ* strain displays a 36-times higher frequency of spontaneous mutation than the wild strain [34]. Moreover, *myh1Δ* showed sensitivity to H2O2. Myh1p binds to proliferative cell nuclear antigen (PCNA) and PCNA-like heterotrimer Rad9p/Rad1p/Hus1p [35–37]. All subunits of the latter complex are required for proper functioning of the checkpoint and DNA replication under stress [38]. Myh1p may act as an adaptor to recruit checkpoint proteins to the DNA lesion [36].
3. AP Endonucleases in S. pombe

AP sites are generated by removal of damaged bases by monofunctional DNA glycosylase and more frequently by nonenzymatic hydrolysis of the N-glycosidic bond of DNA [1, 3]. In a human cell, about 9,000 residues of the AP endonuclease could also convert 3′-OH and 5′-deoxyribose phosphate (dRP). E. coli has two types of AP endonuclease, exonuclease III (Xth), and endonuclease IV (Nfo) [3, 39]. AP endonucleases from various organisms are classified on the basis of structural similarity to Xth and Nfo (Table 2). APE1 in mammals is an Xth-type homolog, which accounts for more than 90% of the AP endonuclease activity in cells [3, 39]. A second Xth-type homolog APE2 was found [40, 41], but no Nfo-type enzymes were present in mammalian cells. In the budding yeast, Apn1p is the major contributor of AP endonuclease activity [30]. However, the apn2Δ single mutant showed the same MMS sensitivity as the nth1Δ single mutant [30]. However, the apn2Δ/nth1Δ double mutant was more sensitive than the apn2Δ single mutant [44]. Moreover, when Apn1p was overexpressed in apn2Δ cells, MMS sensitivity was partially restored [49]. Therefore, Apn1p removes the 3′-blocked end as a back-up function of the major enzyme Apn2p (Figure 1). Since Apn1p was observed in both the nucleus and cytoplasm, it possibly functions in more than just nuclear DNA repair [49]. S. cerevisiae Apn1p has been shown to move into the mitochondria to maintain genomic stability [51].

3.2. Other AP Endonucleases of S. pombe. apn1Δ (SPCC622.17) encodes an Nfo-type AP endonuclease [50]. Although the APN1 mutant of S. cerevisiae is hypersensitive to MMS, the S. pombe apn1Δ mutant exhibited no sensitivity to MMS and oxidative stress. In addition, the nth1Δ/apn1Δ double mutant showed the same MMS sensitivity as the nth1Δ single mutant [30]. Although the apn2Δ/apn1Δ double mutant was more sensitive than the apn2Δ single mutant [44]. Moreover, when Apn1p was overexpressed in apn2Δ cells, MMS sensitivity was partially restored [49]. Therefore, Apn1p removes the 3′-blocked end as a back-up function of the major enzyme Apn2p (Figure 1). Since Apn1p was observed in both the nucleus and cytoplasm, it possibly functions in more than just nuclear DNA repair [49]. S. cerevisiae Apn1p has been shown to move into the mitochondria to maintain genomic stability [51].

UvE1p (SPBC19C7.09c), which is primarily a UV-photoprodct specific endonuclease of S. pombe, could recognize non-UV-induced DNA damage (e.g., the AP site and dihydrouracil) and mismatched bases in vitro, and hydrolyze immediately 5′ to the damage [52, 53]. The protein has a TIM barrel fold that is very similar to the structure of Nfo [54]. S. cerevisiae has no reported UvE1p homolog. Deletion of the uve1 gene in apn2Δ significantly increased MMS sensitivity [44]. Neurospora crassa UvE1p could complement E. coli xthΔ/nfoΔ with respect to sensitivity to MMS and t-butyl hydroperoxide [55]. The mutation frequency of uve1Δ increased when treated with H2O2 [56]. This effect was additively elevated by the deletion of apn2Δ, indicating that both UvE1p and Apn2p contribute to the avoidance of H2O2-induced mutagenesis, but these enzymes define a distinct oxidative damage repair pathway, E. coli Nfo and S. cerevisiae Apn1p can nick DNA on the 5′ side of several oxidized base lesions, generating a 3′ OH and a 5′ dangling damaged nucleotide, which provide an alternative nucleotide incision repair (NIR) pathway to the classic BER [57, 58]. Therefore, S. pombe UvE1p and Apn1p possibly engage in the NIR.

### Table 2: AP endonucleases in bacteria, yeasts, and human cells.

| Organism | Xth | Nfo |
|----------|-----|-----|
| E. coli  | Apn2p | Apn1p |
| S. cerevisiae | Apn2p | Apn1p |
| S. pombe | (Uve1p) | |

AP endonucleases showed in boldface contribute to the major activities in each organism.
pathway by nicking oxidative damage on DNA in an Nth1p-independent manner (Figure 1). In such a case, damaged bases might be removed by a flap endonuclease, Rad2p, after long-patch repair DNA synthesis is primed by the actions of Uve1p and Apn1p [59, 60]. Uve1p works efficiently in mitochondria as well as in the nucleus [61].

3.3. Short- and Long-Patch Repair Subpathways. In the Nth1p-dependent repair pathway, 3′-phosphodiesterase activity of AP endonuclease removes 3′-blocked end resulting in a 1-nt gap, which will be refilled via short-patch BER (Figure 1). However, it is not known whether the BER synthesis extends beyond a single nucleotide, with incorporation of two or more nucleotides (long-patch BER). When AP endonuclease incises the AP site, single-strand break with 3′-OH and 5′-dRP ends is generated. The dRP moiety is released by combined action of a DNA polymerase and Rad2p endonuclease (long-patch BER). An X-family DNA polymerase of S. pombe, Pol4p, has been shown to contain an intrinsic 5′-dRP lyase activity like mammalian DNA polymerases β and λ, suggesting another AP endonuclease pathway to feed into short-patch BER (Figure 1) [62]. S. cerevisiae Trf4 protein plays a similar role in short-patch BER of the yeast [63]. In mammalian cells, the choice of short-patch or long-patch BER may depend on the state of the 5′ terminal moiety and protein-protein interaction of BER components [64]. In vitro analyses of repair synthesis with whole cell extracts or purified enzymes are needed to clarify the relative contribution of short-patch and long-patch BER in S. pombe cells.

4. Enzymes for Removing 3′ Blocked Ends in S. pombe Cells

Single-strand breaks with 3′ phosphate ends in DNA are induced by chemical attacks of endogenous ROS and environmental oxidants [3]. Fpg/Nei-type DNA glycosylases cleave the AP site by β,δ elimination and generate DNA strand breaks with a 3′-phosphate end. In human cells, removal of 3′ phosphate is dependent on 3′-DNA phosphatase (hPNK), and not the major AP endonuclease hAPE1 [65]. Unlike bacteria and humans, S. pombe does not possess Fpg/Nei-type DNA glycosylase, and the physiological source of 3′ phosphate in the yeast is probably due to the action of tyrosyl-DNA phosphodiesterase I (TDP1). TDP1 catalyzes the hydrolysis of 3′-phosphotyrosyl bonds of the irreversible topoisomerase I (Top1)-DNA covalent complex to generate a 3′-phosphate end [66, 67]. A 3′-DNA phosphatase (Pnk1p; SPAC23C11.04c) has been purified from S. pombe on the basis of its ability to process H2O2-damaged DNA to allow DNA synthesis by DNA polymerase [68]. Like human PNK, Pnk1p localizes in nuclei and has both 5′-DNA kinase and 3′-DNA phosphatase activities [69]. The pnk1 mutant showed hypersensitivity to ionizing radiation and camptothecin, an inhibitor Top1, but not to MMS. It remains to be elucidated whether Pnk1p is the major 3′-DNA phosphatase in S. pombe cells.

TDP1 has been implicated in the repair of the irreversible Top1-DNA covalent complex, which can be generated by either exogenous or endogenous factors [67]. Additionally, TDP1 hydrolyses a variety of 3′ lesions, including 3′ phosphoglycolate [70, 71]. A recessive mutation in the human (TDP1) gene is responsible for the inherited disorder, spinocerebellar ataxia with axonal neuropathy (SCAN1) [72]. A tdp1 (SPCP31B10.05) mutant of S. pombe was completely sensitive to camptothecin [73]. Moreover, the mutant cells progressively accumulate DNA damage and rapidly lose viability in nondividing conditions [73]. Low cellular respiration levels protected the tdplΔ cells, indicating that the production of endogenous ROS is a major cause for the accumulation of DNA lesions in the absence of Tdp1p [73]. Therefore, Tdp1p with Pnk1p processes the same naturally occurring 3′ ends, produced from oxidative DNA damage. Rapid and extensive death of the tdplΔ/rhp51Δ double mutant strain suggested the pivotal role of the HR process in DNA repair in tdplΔ cells [73].

5. BER in a Catalase-Deficient Mutant of S. pombe

The BER mutants of S. pombe are substantially resistant to H2O2. A catalase-deficient mutant (ct1Δ), in which ROS scavenging activity is extensively reduced, became sensitive to H2O2 [74, 75]. Deletion of the BER gene (nth1, apn1, apn2, or uve1) from ct1Δ further increased the sensitivity to H2O2, indicating that catalase activity obscures the functions of BER enzymes in vivo [75]. Double mutants in both ct1 and BER genes showed extremely high spontaneous mutation rates, especially in the ct1/nth1 mutant. Vitamin C relieved the mutator phenotype of the ct1/nth1 mutant [75]. The results provide evidence that BER enzymes as well as catalase and an antioxidant contribute in vivo to avoidance of ROS-induced mutagenesis and cell death.

6. Conclusion

In this paper, we described the early steps of BER in the fission yeast highlighting the key roles of DNA glycosylases, AP endonucleases, and other end-cleaning activities in maintenance of genomic integrity. Although the yeast BER pathway consists of evolutionarily conserved enzymes, the major activity for processing the AP site is different from these of the budding yeast and mammals. AP sites generated by monofunctional glycosylase are mainly repaired via the Nth1p-dependent BER pathway [23, 27, 30, 31]. Unlike other model organisms, the major AP endonuclease of this yeast is assigned to Apn2p, which works primarily in the removal of the 3′ block, and poorly in the incision of AP sites [23, 44, 49]. Uve1p engages in an Nth1p-independent pathway by nicking DNA on the 5′ side of oxidative damage [56, 75]. Genetic interactions of BER genes with NER and HR genes suggest synergism among the different DNA repair pathways in the protection of alkylation and oxidative damage [23–25, 30]. NER and HR probably repair the intermediates of BER such as, the AP site and single-strand breaks with 3′ blocks. BER operates most efficiently when specific protein-protein coordination occurs [76]. A study of physical interaction
of the BER proteins will facilitate an understanding of the regulation of BER protein activities and crosstalk between BER and other DNA transaction pathways in yeast cells. Upon nitrogen-starvation, mostly G2 vegetative cells promote two rounds of division and enter the G0 state with 1C DNA [77]. The S. pombe G0 state will provide an excellent model to reconsider the roles of BER and other DNA pathways in resting and nondividing physiological conditions [73, 78].

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