REVIEW ARTICLE

Genetic instability in budding and fission yeast—sources and mechanisms

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One sentence summary: The stability of budding and fission yeast genomes is influenced by two contradictory factors: (1) the need to be fully functional, which is ensured through the replication fidelity pathways of nuclear and mitochondrial genomes through sensing and repairing DNA damage, through precise chromosome segregation during cell division; and (2) the need to acquire changes for adaptation to environmental challenges.

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ABSTRACT

Cells are constantly confronted with endogenous and exogenous factors that affect their genomes. Eons of evolution have allowed the cellular mechanisms responsible for preserving the genome to adjust for achieving contradictory objectives: to maintain the genome unchanged and to acquire mutations that allow adaptation to environmental changes. One evolutionary mechanism that has been refined for survival is genetic variation. In this review, we describe the mechanisms responsible for two biological processes: genome maintenance and mutation tolerance involved in generations of genetic variations in mitotic cells of both Saccharomyces cerevisiae and Schizosaccharomyces pombe. These processes encompass mechanisms that ensure the fidelity of replication, DNA lesion sensing and DNA damage response pathways, as well as mechanisms that ensure precision in chromosome segregation during cell division. We discuss various factors that may influence genome stability, such as cellular ploidy, the phase of the cell cycle, transcriptional activity of a particular region of DNA, the proficiency of DNA quality control systems, the metabolic stage of the cell and its respiratory potential, and finally potential exposure to endogenous or environmental stress.

Keywords: fidelity of replication; dNTP pool; homologous recombination; transcription-associated genome instability; aneuploidy; mitochondrial genome maintenance

INTRODUCTION

The efficiency of the processes that maintain stable DNA is important for cell viability. However, due to the unpredictability of their rapidly changing environment, cells that are unable to adapt might not persist. With respect to evolution, populations must secure a margin of genome variability that allows for the adjustment to new environmental conditions. This state is achieved through mutations. Mutations contribute to genome variations that provide the population with a reservoir of variability that enables it to survive in a changing environment; in this manner, mutations certainly drive evolution. Some mutations appear spontaneously as a consequence of replication errors, error-prone repair of DNA lesions provoked by endogenous factors (e.g. metabolites or reactive oxygen species) and unequal segregation of chromosomes during mitosis (even in cells not exposed to any genotoxic stress conditions). The types of emerging mutations and their rate depend on the availability and efficiency of DNA quality control systems, cell age and ploidy, cell-cycle phase and the metabolic stage of the cell.
Types of mutations also rely on specific features of the DNA sequence itself. For example, repetitive DNA sequences, such as mononucleotide tracts, are difficult templates for DNA polymerases, causing frequent slippage (Fortune et al. 2005). Moreover, trinucleotide repeats are susceptible to the formation of DNA secondary structures and are likely to undergo expansions or contractions (Razidlo and Lahue 2008). In diploid cells in which the second copy of the genome sequence is present, loss of heterozygosity (LOH) occurs at a high rate, predominantly through allelic crossover but also via gene conversion and chromosome loss (Ohnishi et al. 2004). The frequency of rearrangements between two homologous loci in yeast genomes strongly depends on their chromosomal localization; therefore, nuclear architecture also influences genome maintenance (Agmon et al. 2013).

Both simple unicellular organisms and metazoan cells have similar mechanisms for genome maintenance; the same is true for the mechanisms that cause mutations. Because these mechanisms are evolutionarily conserved and their efficiency translates into similar error rates in the cells of all mesophilic organisms (Drake et al. 1998; Drake 2009), the rules governing the preservation of the intact genome established for yeast should be easily applicable to other cell types. Because yeast cells have served as the model for genome stability research for years, the wealth of knowledge in this field has been gathered for this organism. In this review, we present information collected in this field, not only for the budding yeast Saccharomyces cerevisiae that has served as the most frequent model organism but also for the fission yeast Schizosaccharomyces pombe. We hope that these data might be useful for work on all types of cells.

REPLICATION ERRORS

Replication is one of the main sources of spontaneous mutations despite the evolution of mechanisms to avoid mutations and maintain unchanged DNA sequences. Spontaneous mutations result from imperfections in any of three processes: (1) accurate nucleotide selection by DNA polymerases; (2) proofreading 3′ → 5′ exonuclease activity of replicative DNA polymerases; and (3) correction by DNA repair systems of the mistakes made by all DNA polymerases and filling the gaps left after replication.

Nobody is perfect: the DNA polymerases–enzymes imperfecta

Eukaryotic cells usually possess more than 10 DNA polymerases. For example, S. cerevisiae contains 11 polymerases: α, β, γ, δ, ε, σ, θ, ζ, η and Rev1 (Saccharomyces Genome Database. http://www.yeastgenome.org/, 12 November 2014 date last accessed), S. pombe contains 12 (all homologs of S. cerevisiae polymerases plus Pol ε) (PomBase database. http://www.pombase.org/, 12 November 2014 date last accessed) and human cells contain up to 18 polymerases: α, β, γ, δ, ε, σ, θ, τ, υ, κ, λ, μ, ν, Rev1, PRIMPOL and DNTT (GeneCards. http://www.genecards.org/, 12 November 2014 date last accessed) (Table 1). These DNA polymerases belong to several polymerase families including A, B, X and Y. The role they play in cells is determined by their fidelity and processivity (Table 1). The enzymes that are the most precise in DNA synthesis belong to the B and A families of polymerases and are involved in replication. The less accurate enzymes belong mostly to the Y and X families of polymerases and are involved in DNA repair (e.g. in translesion synthesis, TLS). Because the functional mechanisms and roles of DNA polymerases in various processes were extensively studied in yeast S. cerevisiae cells, we will focus on data obtained from this model organism.

Budding yeast possesses three replicative DNA polymerases: polymerase alpha-prime, Pol α (Lucchini et al. 1987; Brooke and Dumas 1991; Brooke et al. 1991); polymerase delta, Pol δ (Chang 1977); and polymerase epsilon, Pol ε (Morrison et al. 1990). Pol α makes one mistake per 10⁴–10⁵ bases replicated (Kunkel et al. 1989), whereas Pol ε (operating on the leading strand) and Pol δ (operating on the lagging strand) (Karthikeyan et al. 2000; Larrea et al. 2010) exhibit error frequencies of 10⁻⁵–10⁻⁷ due to their proofreading 3′ → 5′ exonuclease activity (Shcherbakova et al. 2003; Fortune et al. 2005; McCulloch and Kunkel 2008; Burgers 2009). Polymerase gamma (Pol γ) has a similar error rate (10⁻⁵–10⁻⁶) and is responsible for replication of the mitochondrial genome (McCulloch and Kunkel 2008). The fallibility of DNA synthesis increases when the polymerase does not have proofreading activity. Under these conditions, the error frequency rises to 10⁻³–10⁻⁴, as was shown for Pol ρ and Pol β, or even to 10⁻¹–10⁻², as was calculated for Pol η (Zhong et al. 2006; McCulloch et al. 2007). These polymerases perform poorly on undamaged templates, and their contribution to spontaneous mutagenesis is limited due to their low processivity and strictly regulated access to the DNA template during normal undisturbed DNA synthesis. Instead, they have a unique ability to synthesize DNA from a damaged template, often allowing a proper reconstruction of the sequence in the nascent DNA strand (McCulloch et al. 2004b; Gibbs et al. 2005). These DNA polymerases are known by several different names in the literature. Sometimes they are called alternative polymerases because they are alternatives to replicative polymerases, while other times they are called TLS polymerases because they synthesize across lesions in DNA templates. They have even been called specialized polymerases because they are able to insert the right nucleotide across from the damaged one. Each alternative polymerase has its own cognate substrate; we will describe this phenomenon in more detail in a later section. Only Pol ρ differs from this scheme because it synthesizes longer DNA stretches. Due to its low fidelity, it is responsible for the majority of mutations (Stone et al. 2009; Northam et al. 2010; Stone, Lujan, Kunkel 2012). In fact, Pol ρ is the most mutagenic among all DNA polymerases found in yeast.

DNA synthesis errors can be introduced during two stages of the process: during selection of a correct dNTP for synthesis and at the extension step. dNTP selection occurs mainly by Watson–Crick base pair binding and discrimination against each of the possible mismatched bases prior to covalent incorporation (Echols and Goodman 1991). Misincorporation might take place as a result of the balance between synthesis and excision that compromises the rate of DNA extension, both of which can be influenced by template sequence, abundance of dNTP pools and their bias (Bebenek and Kunkel 1990, 2000). Thus, insertion of an improper dNTP during replication may lead to the creation of a mismatch, as well as the occurrence of a frameshift mutation, according to the following factors: (1) base to base hydrogen bonding between nucleotides in the template and nascent strands; (2) base pair geometry and substrate-induced conformational changes by the DNA polymerase; (3) strength of contact between the polymerase and the DNA minor groove at and upstream of the active site of the polymerase, which influences nucleotide selectivity; and (4) the efficiency of 3′ → 5′ proofreading and strand misalignment (Kunkel and Bebenek 2000) (Fig. 1). DNA polymerases are influenced differently by the factors listed above; therefore, each polymerase generates its unique pattern of errors by which it can be
Table 1. DNA polymerases and their functions in budding and fission yeast.

| Family | DNA polymerase | Fidelity of replication | Subset or orthologs not found | Molecular function | Cell cycle process | Interaction important for DNA synthesis |
|--------|----------------|-------------------------|-------------------------------|--------------------|-------------------|----------------------------------------|
| E      | Replicative DNA polymerase | Pol1, Pol2, Pol3, Pol4, Pol5 | Pol1, Pol2, Pol3 | Pol1, Pol2, Pol3 | Pol1, Pol2, Pol3 | Pol1, Pol2, Pol3 |
| B      | TLS polymerase | Pol1, Pol2, Pol3 | Pol1, Pol2, Pol3 | Pol1, Pol2, Pol3 | Pol1, Pol2, Pol3 | Pol1, Pol2, Pol3 |
| B      | Pol β | Pol1, Pol2, Pol3 | Pol1, Pol2, Pol3 | Pol1, Pol2, Pol3 | Pol1, Pol2, Pol3 | Pol1, Pol2, Pol3 |
| B      | Pol α | Pol1, Pol2, Pol3 | Pol1, Pol2, Pol3 | Pol1, Pol2, Pol3 | Pol1, Pol2, Pol3 | Pol1, Pol2, Pol3 |

Additional information: All information listed in this Table 1 is available at [Saccharomyces Genome Database](http://www.yeastgenome.org/) and [PomBase](http://www.pombase.org). Additional data have been published in [Skoneczna et al., 2008; McCulloch and Kunkel, 2008; Burgers, 2009].

**Gene Ontology annotations for S. cerevisiae protein. Orthologs in other fungi mostly play similar role in the cell. In some cases more information is available for gene product from other fungi than for its S. cerevisiae ortholog.**
recognized. Each polymerase also has its own function in the cell, such as acting on the leading or lagging strand, gap filling and synthesizing mitochondrial DNA (mtDNA). The same factors influence both replicative and non-replicative DNA polymerases, with the exception that the latter group lacks 3′→5′ exonuclease activity. Additionally, Rev1 does not use canonical Watson–Crick base pairing during synthesis. Probably for this reason, this enzyme has the lowest DNA fidelity among all known yeast DNA polymerases (Nair et al. 2005; Prakash, Johnson and Prakash 2005; Waters et al. 2009). On the opposite side of the DNA synthesis fidelity scale are the major replicative polymerases Pol δ and Pol ε. Pol δ was shown to have high selectivity for the correct nucleotide based on kinetic studies of the insertion of individual dNTPs opposite a known nucleotide in a DNA template. This ability together with its proofreading 3′→5′ exonuclease activity establishes Pol δ as the most accurate polymerase in yeast. Nonetheless, Pol δ often generates single nucleotide deletions in homopolymeric runs and wastefully proofreads these types of mismatches with a much higher error rate (approximately three single nucleotide deletions per 10⁴ nucleotides polymerized). Additionally, Pol δ may also slide over templates containing direct repeats spaced by three or more base pairs, resulting in deletions of the nucleotides between these direct repeats. Pol δ strand slippage on repetitive sequences during replication is believed to be the major source of insertions and deletions in yeast genomes (Fortune et al. 2005). As mentioned above, DNA polymerases can be distinguished by the pattern of mistakes they make. Although both Pol δ and Pol ε incorporate nucleotides with high fidelity, Pol δ has a higher probability of sliding on repetitive sequences. In contrast, the high fidelity of Pol ε results from the high effectiveness of its 3′→5′ exonuclease proofreading activity rather than from base selection correctness; this finding is based on the in vitro spectrum of errors (mutation spectra) observed for a proofreading-deficient form of Pol ε that showed a unique error signature with a high proportion of transversions resulting from T-T, T-C and C-T mispairs (Scherbakova et al. 2003). In turn, the disruption of S. cerevisiae Pol γ exonuclease activity increased the mtDNA deletion rate 160-fold, indicating that exonuclease activity is crucial for avoiding deletions during mtDNA replication (Stumpf and Copeland 2013). This result also suggested a possible source of mtDNA deletions of the progeroid phenotype in exonuclease-deficient DNA polymerase γ in mice (Stumpf and Copeland 2013). Pol γ proofreading 3′→5′ exonuclease activity minimizes the frequency of point mutations and prevents deletions, thereby contributing to the stabilization of mtDNA in yeast cells (Vanderstraeten et al. 1998). However, apart from its proofreading activity, the exonuclease domain of Pol γ contributes to the coordination of its polymerase and exonuclease functions. This was documented by the analysis of phenotypes of mip1 (Pol γ) alleles, in which mutations were localized to the DNA-binding channel of the exonuclease domain in close vicinity to the polymerase domain. In these mutants, the imbalance between DNA synthesis and degradation caused poor mtDNA replication (Szczebanowska and Foury 2010). However, increased mutagenesis was also detected in strains encoding mutant mip1 variants that were unable to maintain mtDNA, although they were not affected by polymerase fidelity or exonuclease proofreading activity. Increased mutagenesis was in this case caused by slowing down the replication fork, thereby predisposing the template DNA to irreparable damage that was bypassed with a poor fidelity (Stumpf and Copeland 2014).

During normal replication, three DNA polymerases (Pol α, Pol ε and Pol δ) work together at the replication fork to duplicate the DNA. The replication fork polymerases are programmed to replicate opposite DNA strands; Pol ε synthesizes the leading strand, while primases Pol ρδ and Pol δ polymerize the Okazaki fragments on the lagging strand (Karthikeyan et al. 2000; Larrea et al. 2010). This spatial arrangement causes replication fork asymmetry, which is observed in both budding and fission yeast and influences DNA synthesis fidelity (Kunkel 1994; Miyabe, Kunkel and Carr 2011). Nonetheless, the replicative polymerases might operate at the very same cytological foci (Hiraga et al. 2005), correcting one another’s mistakes (Morrison and Sugino 1994; Pavlov et al. 2004; Burgers 2009).

Yeast replicative DNA polymerases work in complexes composed of several subunits that require proper assembly (Table 1). Formation of the holoenzyme depends on the Mms19 (Met18) protein belonging to the cytosolic iron–sulfur protein assembly machinery and encompasses not only assembly of the active enzyme from several subunits (which most likely requires chaperone assistance) but also installation of the necessary
cofactors (Stehling et al. 2012). Mms19 is involved in assembly of the [4Fe-4S] cluster in the CysB motif of several DNA polymerases, including Pol α, Pol ε, Pol δ and Pol γ. This prosthetic group is needed for enzyme stabilization. Installation of another cofactor (the Zn$^{2+}$ ion in the CysA motif of Pol δ) is required for its PCNA-mediated processivity (Netz et al. 2011). The major subunit in the DNA polymerase holoenzyme is responsible for DNA polymerization and optionally for its 3′→5′ exonuclease proofreading activity. In S. cerevisiae cells, the accessory proteins contribute to the activity of the enzyme and influence its fidelity and processivity. The accessory subunits play an additional role in maintaining contact between the holoenzyme and other cellular components via various interactions. These interactions permit both access to the DNA template and the transmission of important cellular signals to the polymerase, allowing for a proper response. Thus, the accessory subunits may modulate polymerase activity. For example, the interaction between Pol32 (one of the non-catalytic subunit of Pol δ) and Pol30 determines Pol δ processivity. The homotrimer of Pol30 forms a circular structure called PCNA (proliferating cell nuclear antigen) that serves as the DNA polymerase processivity factor. The PCNA works as a sliding clamp encircling the DNA strand and tethers the polymerase to the template, thereby preventing its dissociation (Fukuda et al. 1995; Johansson, Garg and Burgers 2004). Pol32 interacts with the Pol1 subunit of Pol α. This interaction enables cooperation between Pol α and Pol δ that is critical for lagging-strand synthesis (Johansson, Garg and Burgers 2004). The accessory subunits of both replicases Pol δ and Pol ε permit their interaction with different proteins engaged in the DNA repair pathways. They compete for the same interaction partners (e.g. PCNA) with specialized polymerases, influencing their access to the DNA template. The Pol31 and Pol32 subunits of Pol δ and the Dpb2, Dpb3 and Dpb4 subunits of Pol ε contribute to the fidelity of replication by influencing dNTP selection and/or the proofreading activity of catalytic subunits of the respective holoenzyme and by stabilizing its interaction with DNA (Giot et al. 1997; Huang et al. 2002; Aksenova et al. 2010). Accordingly, mutant versions of the Dpb2 gene significantly influence spontaneous mutagenesis (Jaszczur et al. 2009). The polymerase subunits also have an impact on regulation of the replication process. Disrupted interactions between Pol ε subunits and the GINS complex result in increased mutagenesis (Grabowska et al. 2014). The Pol1 interaction with chromatin-binding protein Ctf4 associated with the GINS complex couples the heterohexameric Mcm2-7 helicase to Pol α. This interaction is important for the formation of the correct replisome progression complex on every Okazaki fragment during lagging-strand synthesis (Gambus et al. 2009). In contrast, the Pol2 Pol ε catalytic subunit binds to Mrc1 when this S-phase checkpoint protein is associated with the Mcm2-7 helicase. This binding permits matching of the polymerization with the unwinding rate on the leading DNA strand of the replication fork during normal replication (Lou et al. 2008). Another possible source of replication errors is continued DNA synthesis when the DNA is subjected to damage. To avoid this scenario during DNA damage stress, the checkpoint protein Mrc1 is phosphorylated by a Mecl-dependent manner. Then, phosphorylated Mrc1 interacts with Tof1 to form a pausing complex that is required for Pol ε stabilization at stalled replication forks (Osborn and Elledge 2003). Thus, transient interactions of replicative polymerases with the Ctf4 and Mrc1 proteins protect cells against chronic activation of the DNA damage checkpoint during chromosome replication and permit the finishing of DNA synthesis and subsequent finalization of the cell cycle. Replication blocks can also be overcome by activating the checkpoint response to regulate template switching or origin firing, fork restart and cell-cycle progression, all of which will engage DNA repair (Friedel, Pike and Gasser 2009). Presumably, almost all components of the replication fork machinery contribute to replication accuracy. Indeed, improper functioning of proteins engaged in initiation, elongation or replication control manifest as a mutator phenotype. Among other abnormalities, genome instability phenotypes are present in strains defective in origin recognition complex, various replication factor C (RFC) complexes, and Mcm2-7 helicase complex, and these phenotypes have been reported for cdc45, csm3, ctf18, dbp11, pol30 and sl4d mutant cells (Stone et al. 2008; Alabrudzinska, Skoneczny and Skoneczna 2011; Li and Tye 2011; Cheng et al. 2012). Significantly, mutations in genes encoding proteins functioning in the replication fork that frequently cause replication stalling often lead to gross chromosomal rearrangements (GCR), whereas mutations arising as a consequence of polymerase defects are mainly base substitutions and frameshifts.

The availability of DNA synthesis precursor dNTPs affects the quality of the nascent DNA

DNA synthesis requires a constant supply of dNTPs to proceed. The size of the dNTP pools and their bias (i.e. relative concentrations) must be tightly regulated to ensure optimized DNA metabolism. dNTP pool concentrations oscillate during the cell cycle, reaching a maximum during transition through the G1/S boundary. This feature helps cells set up for the next round of replication and allows progression of the cell cycle (Koç et al. 2003; Chabes and Stillman 2007). dNTP pools also rise in response to DNA damage and stalling of the replication fork, facilitating replication through DNA lesions (Chabes et al. 2003; Koç et al. 2004; Chabes and Stillman 2007; Lis et al. 2008; Sabouri et al. 2008; Davidson et al. 2012; Poli et al. 2012). However, according to data obtained for various eukaryotic cells, this bypass is often mutagenic (Echols and Goodman 1991; Lis et al. 2008; Sabouri et al. 2008). As was shown for human cells, low dNTP concentrations favor high-fidelity replication, whereas high dNTP concentrations stimulate polymerization reactions and inhibit the proofreading activity of replicases, thereby enhancing the dNTP misincorporation rate (Kunkel, Silber and Loeb 1982; Echols and Goodman 1991). However, the amount of dNTPs is not the only factor that contributes to this phenomenon. As was shown for yeast cells, increased levels of misinsertions, strand misalignments and mismatch extensions at the expense of proofreading can result from an imbalance in the dNTP pools (Kumar et al. 2011). Interestingly, changes in the dNTP pool size or bias can induce not only point mutations but also DNA breaks, which can lead to GCR (Ouspenski, Elledge and Brinkley 1999; Chabes et al. 2003; Basullo et al. 2010; Davidson et al. 2012).

The maintenance of proper dNTP levels is essential for cellular viability and genome stability, therefore, the level and activity of the ribonucleotide reductase complex (RNR), the enzyme responsible for the rate-limiting step in dNTP synthesis, is one of the most tightly regulated in the cell (Fig. 2) (Sanvisens, de Llanos and Puig 2013). The RNR complex is composed of small and large subunits: R2 and R1, respectively. In S. cerevisiae, the small subunits (R2) are encoded by the RNR2 and RNR4 genes, and the large subunits (R1) are encoded by the RNR1 and RNR3 genes (Elledge and Davis 1987, 1990). At the transcriptional level, RNR genes are regulated by the cell cycle and DNA damage response networks. Cell-cycle-regulated expression of RNR genes in S. cerevisiae depends on the MBF (Mbp1-Swi6) transcription factor and leads to gradual accumulation of their transcripts during the G1 phase.
Figure 2. The RNR regulation in *S. cerevisiae*. The transcription of RNR genes in the cell cycle is MBF dependent, and occurs in G1 reaching the maximum near G1/S transition. When Stb1 is unphosphorylated, it binds the Swi6 subunit of the MBF complex causing repression of RNR genes. In early G1 phase, Stb1 is phosphorylated by CDK which turns on RNR transcription. In late G1, Nrm1 promotes the MBF-dependent repression of RNR genes. In response to DNA damage in the S phase, the Mec1/Rad53/Dun1 kinase cascade activates RNR transcription. Hyperphosphorylation of Crt1 repressor turns on the expression of RNR2,3,4 genes. Moreover, phosphorylation by Rad53 of Nrm1 causes its release from RNR promoters; and phosphorylation of Ixr1 activates the transcription of RNR1 gene. Additionally, the Mec1/Rad53/Dun1 pathway activates RNR activity by promoting the phosphorylation and degradation of the RNR inhibitor, Sml1 and enhancing the relocation of small RNR subunits to the cytoplasm via modifications of Wtm1 (nuclear anchor protein for RNR small subunits) and Dif1 (importin). Protein names in white refer to transcription factors, in black to proteins with other functions.
allows DNA synthesis to resume and is managed by type 2A-like protein phosphatase activity of the Pph3-Psy2 complex. It was shown that Rad53 independently regulates restart at the replication forks and firing of late origins and that regulation of these processes is mediated by specific Rad53 phosphatases (O’Neill et al. 2007). In addition, rad53 mutants treated with hydroxurea accumulate unusual DNA structures at the replication forks, resulting in genome rearrangements. This finding indicates that Rad53 prevents the collapse of the fork and avoids genome destabilization when replication pauses (Lopes et al. 2001). Notably, impaired function of other DNA damage or replication block checkpoint proteins, such as Mec1, Tel1, Rad9, Ctf18 or Mrcl, also lead to elevated genome instability (Lustig and Petes 1986; Yuen et al. 2007; Lou et al. 2008; Razidlo and Lahue 2008; Alabrudzinska, Skoneczny and Skoneczna 2011).

An optimal dNTP pool supply is also ensured by the post-transcriptional regulation of RNR complex biogenesis and activity (Ouspenski, Elledge and Brinkley 1999; Chabes et al. 2003; Fusulio et al. 2010; Kumar et al. 2010, 2011; Davidson et al. 2012). In response to DNA damage, RNR activity inhibitor Sml1 (Spd1 in S. pombe) is phosphorylated and degraded in a Mec1/Rad53/Dun1-dependent manner; thus, the production of dNTPs necessary for DNA repair and replication effectively increases (Zhao and Rothstein 2002; Nestoras et al. 2010). In S. pombe, Spd1 impedes RNR holoenzyme formation by sequestering the R2 subunit in the nucleus from the cytoplasmic R1 subunit (Fleck et al. 2013). Additionally, RNR activity is regulated by the compartmentalization of its subunits. It was shown that the small subunits of S. cerevisiae RNR (Rnr2 and Rnr4) are imported to the nucleus in a Dif1-dependent manner and then anchored by Wtm1 (Lee and Elledge 2006). Similarly to Sml1 and Spd1, Dif1 is phosphorylated and degraded following DNA damage (Wu and Huang 2008), permitting the redistribution of RNR subunits to the cytoplasm, where they can assemble into an active complex (Yao et al. 2003). Proper assembly of RNR also requires a number of cofactors, including an atypical iron prosthetic group called diferroc-tirosyl radical cofactor [FeIIIY•], monothiol glutaredoxin heme protein complex Grx3-Grx4 and iron–sulfur assembly complex Dre2-Tah18 (Zhang et al. 2008; Netz et al. 2010; Stehling et al. 2012; Zhang 2014). The active holoenzyme also requires thioredoxin and glutaredoxin as electron donors during DNA precursor synthesis; thus, dNTP production can also be influenced by cytoplasmic thioredoxins (Trx1 and Trx2), glutathione reductase (Glr1) and gamma glutamylcysteine synthetase (Gsh1). In fact, trx1Δ trx2Δ cells are unable to accumulate dNTPs and display elevated levels of glutathione reductase (Koc et al. 2006). It has also been shown that thiorexin metabolism is employed to buffer deficiencies in RNR by enabling a compensatory increase in de novo purine biosynthesis that provides additional rate-limiting substrates for dNTP production (Hartman 2007).

The next level of RNR regulation involves allosteric control of its enzymatic activity, permitting an adequate and balanced supply of all four dNTPs and fast adaptation to perturbation of cellular dNTP pools. The R2 subunit of RNR generates a tyrosyl radical that is continuously shuttled to a cysteine residue in the active site of the R1 subunit during catalysis (Kolberg et al. 2004). In addition to the active site, the R1 subunit also contains two separate allosteric sites (the S site and A site) to control activity and substrate specificity (Jordan and Reichard 1998; Eklund et al. 2001). The allosteric sites bind specific NTPs and dNTPs as effectors. Through binding ATP, dATP, dTTP or dGTP, the S site controls the specificity for each of the four substrates. Through binding ATP and dATP, the A site adjusts enzyme activity.

Binding of ATP to the A site stimulates enzyme activity, while binding of dATP inhibits it (Cooperman and Kashlan 2003; Zhang, Liu and Huang 2014).

Interestingly, the interaction between Spd1 (an allosteric regulator of RNR) and DNA-associated PCNA has recently been reported in S. pombe. This interaction is required for Spd1 degradation following ubiquitination by ubiquitin ligase CRL4Cdc2 (Salguero et al. 2012). This finding places RNR directly at the replication fork. Moreover, it has been demonstrated that a mutation in the R1 subunit of RNR (cdc22-D57N) that alleviates allosteric feedback inhibition resulted in highly elevated dNTP pools that were further increased by deletion of the SPD1 gene. The Δspd1 cdc22-D57N double mutant showed increased mutation rates and was more sensitive to damaging agents, causing double-strand breaks (DSBs) stress compared to single mutants. Thus, Spd1 can protect the genome when dNTP pools are high. In contrast, overexpression of Spd1 generates replication stress and provokes genome instability (Fleck et al. 2013).

Correction systems of replication errors

Cleaning up after replication-mismatch repair and ribonucleotide excision repair

When the proofreading activity of replicases fails to remove the deoxynucleotides misinserted during DNA synthesis, the last chance to correct the mistake is employment of the mismatch repair (MMR) system. The functioning of the MMR system was nicely reviewed by Li (2008), Spampinato et al. (2009) and Kolodner and Marsischky (1999). The first step in this pathway depends on specialized protein complexes Msh2-Msh3 or Msh2-Msh6 that recognize structural abnormalities in the DNA helix. The structural anomalies may be caused by mispaired bases or the insertion/deletion of one or more nucleotides. To properly correct the spatial anomaly detected in the DNA, the nascent and template DNA strands have to be distinguished. In yeast, the nascent strand is discriminated based on the discontinuity associated with DNA replication. Consistent with this model, nucleosome Exo1 was found to preferentially repair errors made by Pol α on the lagging strand (Liberti, Larrea and Kunkel 2013). It has been postulated that MMR initiates at Okazaki fragment termini in the lagging strand and at nicks generated in the leading strand by the mismatch-activated Mlh1-Pms2 endonuclease. Because cells lacking RNase H2 display a partial MMR defect combined with an increase in mutagenesis, it was recently proposed that a single ribonucleotide in the vicinity of a mismatch incised by RNase H2 can also act as an initiation site for MMR. Therefore, ribonucleotides misincorporated during DNA replication may serve as physiological markers of the nascent DNA strand (Ghodgaonkar et al. 2013). Following recognition of the template and nascent strands, they are marked via asymmetric binding of the Mlh1-Pms1, Mlh1-Mlh2 or Mlh1-Mlh3 complexes. Then, the fragment of nascent DNA containing the distortion can be removed by flap endonucleases, mainly Exo1, but also Rad27 (Fen1) or Sgs1-Top3 (Kao et al. 2002; Tran et al. 2002; Fricke and Brill 2003). The recent studies established that also Mlh1-Pms1 endonuclease is required for MMR in Exo1-independent MMR subpathway (Smith et al. 2013; Goellner et al. 2014). Excision of the DNA fragment containing the replication error creates an ssDNA gap, which not only provides Pol α with an opportunity to fill it using the original DNA as template but also contributes to checkpoint activation that is needed to maintain genome integrity because ssDNA is believed to be the checkpoint activation signal (Mojas, Lopes and Jiricny 2007; Reha-Krantz et al. 2011). The final step of MMR is ligation. Other components of this
pathway include the ssDNA-binding complex RPA (Rfa1-Rfa2-Rfa3), the PCNA-loading clamp (RFC complex) and PCNA. PCNA serves various DNA repair processes including those involved in MMR as a handy platform facilitating access to DNA (Clark et al. 2000; Shell, Putnam and Kolodner 2007; Stone et al. 2008).

Bowen et al. (2013) performed a series of amazing experiments reconstituting MMR reactions using purified S. cerevisiae proteins engaged in MMR and a set of DNA substrates containing diverse defects. It was found that a mixture of Msh2-Msh6, Exo1, RPA, RFC-A1N, PCNA and Pol δ was sufficient to repair substrates containing various mispairs, frameshifts and a 3′- or 5′-strand break with a range of efficiencies. Furthermore, Bowen et al. showed that the Msh2-Msh3 complex could substitute for the Msh2-Msh6 complex; however, analysis of heterodimers showed a different specificity towards repair of the various mispairs and the addition of the Mlh1-Pms1 complex had no effect on MMR. The Msh2-Msh6 complexes and all three Mlh1-heterocomplexes (Mlh1-Pms1, Mlh1-Mlh3 and Mlh1-Mlh2) also function in the resolution of recombination intermediates and bind with high affinity to a cross-shaped four-stranded structure called a Holliday junction (HJ) that contributes to reciprocal meiotic recombination (Marsischky et al. 1999; Argueso et al. 2003; Rogacheva et al. 2014).

As mentioned above, the MMR system can effectively correct not only mispaired bases but also various frameshifts that are particularly difficult for polymerases to correct using their proofreading activities (Romanova and Crouse 2013). Frameshifts appear most often when the polymerase reaches a polynucleotide tract, which is a hotspot for mutagenesis during DNA synthesis. These tracts result from spontaneous, replication-associated strand slippage (Kunkel 1990) or one of two possible template misalignment mechanisms (primer-template misalignment or dNTP-stabilized misalignment). The insertion of an incorrect nucleotide creates a mispaired primer terminus that is difficult for the DNA polymerase to extend. Subsequent primer-template misalignment can restore proper base pairing, thereby promoting efficient primer extension by relocation of the terminus. This results in a +1 frameshift if the misinserted nucleotide is complementary to the next base of the template strand or a –1 frameshift if it is complementary to the previous base (Bebenek and Kunkel 1990). In the dNTP-stabilized misalignment mechanism, a dNTP substrate is paired correctly, although not with the next available template base. If the next base does not match, the mismatched base is paired with a downstream base on a ‘looped out’ template strand instead. This mechanism generates a –1 frameshift due to the skipping of nucleotides on the template strand by the polymerase (Efrati et al. 1997; Kobayashi et al. 2002). Recently, it was also suggested that the flanking sequence of mononucleotide tracts can play a role in initiating primer-template misalignment; additionally, non-homologous end-joining pathway (NHEJ) defects and MMR system deficiencies can generate frameshifts in a replication-independent manner. Moreover, it was shown that NHEJ is uniquely required for the de novo creation of tandem duplications from non-iterated sequences (Lehner et al. 2012).

The results obtained recently for both S. cerevisiae and Sc. pombe cells suggest an additional class of replicative errors. These errors arise from ribonucleotide monophosphate (rNMP) incorporation into nascent DNA (Watt et al. 2011). These are very common because they happen on average once per 2 kb of newly synthesized DNA; their high frequency is thought to be due to high cellular levels of ribonucleotide triphosphates, although their incorporation is mostly limited by the selectivity of DNA repliases. However, if these misinsertions persist after the DNA synthesis step, they cannot be corrected by MMR; instead, they require the special enzyme RNase H2 for repair. RNase H2 is normally used for the removal of RNA primers from Okazaki fragments, but this enzyme is also responsible for incision of the ribonucleotide, which is the initiation step of ribonucleotide excision repair (NER). During the NER pathway, the incision step is followed by excision by the flap endonuclease Fen1 (or with lower efficiency by Exo1). Next, strand displacement synthesis performed by DNA Pol δ or Pol ε and the process is finally completed by DNA ligase I (Miyabe, Kunkel and Carr 2011; Sparks et al. 2012). The NER pathway also employs PCNA, which is loaded onto DNA by the RFC clamp loader complex. Due to the interaction with PCNA, all successive components of the pathway gain access to the DNA (Sparks et al. 2012). In RNase H2-deficient strains, rNMP incorporated into the leading strand can be removed by processing outside the context of replication in a process that requires Top1 and repetitive sequences and gives rise to misaligned intermediates, resulting in short deletions (Clark et al. 2011; Kunkel 2011; Miyabe, Kunkel and Carr 2011). Another way to overcome rNMP incorporation into DNA in RNase H2-deficient strains is to use post-replication repair (PRR) pathways. In these strains, PCNA is constantly mono- or polyubiquitinated, so PRR is constitutively activated, and misincorporated ribonucleotides can be efficiently bypassed by Pol ζ or omitted in a Mms2-dependent template switch manner (Lazzaro et al. 2012).

Recent studies have demonstrated the importance of yet another protein for the proper functioning of the RER pathway. DNA ligases can generate DNA damage through abortive ligation that produces chemically adducted, toxic 5′-adenylated DNA lesions, (e.g. adenylated 5′ ends containing ribose), which is characteristic of RNase H2 incision. Fortunately, both Sc. pombe and S. cerevisiae also harbor aprataxin (Aptx and Hnt3, respectively), an enzyme with 5′-adenylated RNA–DNA deadenylase activity. Aprataxin efficiently repairs 5′-adenylated RNA–DNA and acts in the RNA–DNA damage response to promote cellular survival and prevent S-phase checkpoint activation in budding yeast undergoing RER (Tumbale et al. 2014).

**PRR: a specialized pathway to repair replication stall-borne problems**

The processes described so far are designed to make replication error free. They generally perform flawlessly while all components remain functional. However, when any of them fails, mutations arise. Another error-prone situation is the presence of DNA lesions in the template. DNA lesions appear frequently in response to endogenous or exogenous stresses, but as long as they are recognized and corrected by appropriate repair pathways, they do not cause any problems. DNA lesions become mutagenic if they persist until the next DNA replication round. Unrepaired damage in the template strand causes difficulties during DNA synthesis. The type of lesion emerging in DNA determines selection of the appropriate response pathway. Possible responses include (1) TLS performed either by the replicative polymerase or specialized DNA polymerase(s), which is facilitated by an increase in the dNTP pool; and (2) the DNA damage avoidance pathway, in which lesions may be passed by the DNA template change. However, in some cases, the lesions block replication completely and cause a permanent stall in the replication fork, followed by cell-cycle arrest and, eventually, death. The repair pathway responsible for overcoming DNA lesion problems during replication is the PRR pathway. Recently, a controversy arose regarding the actual time window during which PRR occurs. The previous belief was that PRR operates during replication. Now, the scientific community has split into
two groups. Some believe that PRR functions during replication, while others favor the view that PRR operates on single-stranded gaps left behind the replication fork or formed during the repair of lesions arising at the G2/M phase (Daiagaku, Davies and Ulrich 2010; Karras and Jentsch 2010; Putnam, Hayes and Kolodner 2010). Nevertheless, PRR is the cellular pathway designed to cope with replication of the problematic template.

Two PRR subpathways exist in \textit{S. cerasvisea}: one that is error prone and one that is error free. It has been shown that PCNA trimer stability is required for TLS by Pol \( \delta \) and by Pol \( \eta \) (Dieckman and Washington 2013). Moreover, PCNA functions as a molecular switch between the error-prone and error-free PRR subpathways. A change in the operating pathway occurs in response to a DNA damage signal followed by checkpoint activation and is realized by alternative covalent modifications of PCNA (Hoege et al. 2002; Watts 2006; Andersen, Xu and Xiao 2008; Zhang et al. 2011) and the different affinity of DNA polymerases for its modified forms (Garg et al. 2005; Parker et al. 2007; Acharya et al. 2011). Access to the perturbed replication fork is thought to be regulated by competition between polymerases, leading to the conclusion that it can also be influenced by a change in the relative abundance of polymerases in the nucleus. In turn, the relative abundance of polymerases in the nucleus is regulated by their varied expression levels during the cell cycle (e.g. expression of the replicases peaks at the G1/S phase (Verma et al. 1991), while the expression of Pol \( \zeta \) peaks at G2/M (Waters and Walker 2006)), by their transport and assembly (Netz et al. 2011), and finally by their stability (e.g. Pol \( \eta \) and Rev1 levels are regulated via ubiquitin-dependent degradation; Podlaska et al. 2003; Skoneczna et al. 2007; Wiltrout and Walker 2011). The fluctuations in polymerase delivery to the replication fork may influence the probability of their interaction with PCNA because various polymerases have similar affinities for the same modified form of PCNA, as do Pol \( \eta \), Pol \( \zeta \) and Rev1 in budding yeast. It was reported that in the cells of higher eukaryotes, DNA polymerases are also subject to modifications that control their PCNA-binding abilities or even exclude them from the chromatin (Lehmann 2011). Recently, the first report was published, concerning ubiquitin-dependent degradation of yeast Pol3 (catalytic subunit of Pol \( \delta \)), which excludes it from the replication fork after DNA damage, allowing access of Rev1 (Dara and et al. 2014). This finding is in line with transient stabilization of Pol \( \eta \) observed after UV irradiation (Skoneczna et al. 2007). Together, these data suggest that polymerase switch (specialized polymerase in place of replicative one in case of DNA damage) can be the ubiquitin-mediated feature.

During undisturbed replication, PCNA is sumoylated by the Ubc9–Siz1 complex (Stelter and Ulrich 2003; Huang et al. 2007), which permits interaction with replicative Pols \( \delta \) and \( \epsilon \) and fast replisome progression at the replication forks. Detection of bulky lesions on the DNA template impedes fork progression and induces Rad6–Rad18-mediated ubiquitination of PCNA, therefore promoting Ubi–PCNA-dependent translesion DNA synthesis (TLS); further polyubiquitination of Ubi–PCNA performed by the Ubc13–Mms2–Rad5 ubiquitin ligase complex in turn favors the damage avoidance pathway (Fig. 3). When synthesis over a lesion is finished, Ubp10 deubiquinates PCNA, allowing replisome remodeling to switch back to the replicase and the efficient resumption of replication (Gallego-Sánchez et al. 2012). Desumoylation of PCNA is performed by the SUMO proteases Ulp1 that removes SUMO and Ulp2 that removes poly-SUMO chains (Stelter and Ulrich 2003; Huang et al. 2007; Drag and Salvesen 2008).

Information on PRR in \textit{Sc. pombe} is limited. It is known that \textit{Sc. pombe} and \textit{S. cerasvisea} share homologous sets of PRR genes, but they are used in different ways. The major distinctions of \textit{Sc. pombe} cells in the control of lesion tolerance pathways are the existence of pathways that involve various TLS polymerases, the TLS requirement for enzymes needed for mono- and polyubiquitination of PCNA and, surprisingly, the mostly error-free bypass of various DNA lesions regardless of the polymerase or polymerases synthesizing through them (Coulon et al. 2010). Although Siz1 and Ubc9 homologs exist in \textit{Sc. pombe}, PCNA sumoylation has not been documented. The data obtained for cells carrying mutations in various DNA polymerase genes showed that even though all four \textit{Sc. pombe} TLS polymerases operate on DNA templates containing lesions (i.e. the substrate that is recognized by the receptors as a DNA damage signal), only cells lacking Pol \( \kappa \) or Pol \( \eta \) (but not Pol \( \zeta \) or Rev1) activated the checkpoint after UV irradiation. Interestingly, the enhanced checkpoint response in cells lacking Pol \( \kappa \) and Pol \( \eta \) was not due to stalled replication forks but was instead caused by post-replication DNA gaps with unrepaired UV lesions in the template that acted as both substrates for TLS polymerases and as signals for checkpoint activation (Callegari et al. 2010). Previous results demonstrated that checkpoint activation in fission yeast contributes to Pol \( \kappa \) function in the DNA damage-tolerant pathway differently than in budding yeast. The Rad17 checkpoint protein provides both activation of \textit{dinB} (encoding Pol \( \kappa \)) expression and supports its physical interaction with the checkpoint-clamp components Hus1 and Rad1 (Kai and Wang 2003). Thus, it could be expected that TLS is a part of the checkpoint response.

Pol \( \eta \) is the only polymerase for which transcriptional activation of the encoding gene following DNA damage has been demonstrated in budding yeast. After UV induction, RAD30 transcript levels increase about 2-fold (Pavlov, Nguyen and Kunkel 2001). However, because the overexpression of Rad30 (Pol \( \eta \)) and the Pol \( \zeta \) subunits Rev3 or Rev7 increase the mutation rate (Rajpal, Wu and Wang 2000; Pavlov, Nguyen and Kunkel 2001), they should be maintained at low levels under non-mutagenic conditions. Indeed, all budding yeast TLS polymerases (Pol \( \eta \), Pol \( \zeta \) and Rev1) need to be removed from the replication fork as quickly as possible after their task is performed; otherwise, they will frequently generate mistakes on the undamaged template due to their poor fidelity (Zhang et al. 2006; McCulloch et al. 2007). In some cases, they can even produce complex mutations (e.g. tandem base pair substitutions and clusters of multiple, closely spaced mutations) as was shown recently for Pol \( \zeta \) (Stone, Lujan, and Kunkel 2012). For this reason, TLS polymerases are actively removed from the cell when they are no longer required. Rev1 and Pol \( \eta \) are short-lived proteins in vivo, with half-lives of approximately 20 min, at least for their tagged versions (Skoneczna et al. 2007; Wiltrout and Walker 2011). The Rev1 protein is additively regulated during the cell cycle, and is most unstable during G1 phase (Wiltrout and Walker 2011). Because at least two TLS polymerases are degraded in a ubiquitin-dependent manner, it is not surprising that the mutator effect was also observed in yeast strains with reduced proteasomal activities (Podlaska et al. 2003; McIntyre et al. 2006). Furthermore, Pol \( \eta \) was shown to be transiently stabilized after UV irradiation (Skoneczna et al. 2007), when its cognate substrates (e.g. T-T dimers) emerge in the DNA. Pol \( \eta \) specializes in the bypass of T-T dimers and inserts appropriate nucleotides (i.e. AA) relatively accurately opposite these lesions (Washington et al. 2000; Gibbs et al. 2005).

Sumoylated PCNA-, Srs2- and replicative polymerase-dependent DNA synthesis on damaged templates The PCNA of \textit{S. cerasvisea} has two
SUMO acceptor sites: a major one at Lys164 (that may also be ubiquitinated) that is crucial for DNA damage tolerance and a minor one at Lys127 that does not appear to play a role in DNA damage tolerance activity but is required for the establishment of sister chromatid cohesion during S phase (Hoege et al. 2002; Stelter and Ulrich 2003; Moldovan, Pfander and Jentsch 2006). Because PCNA forms homotrimeric complexes, there are three Lys164 residues available for potential modification. Sumoylation of one Lys164 facilitates modification of its neighboring Lys164 in the complex, but the ligand for subsequent modification may not necessarily be the same as the primary ligand. Therefore, various modifications can exist for one complex, including poly-SUMO chains (Windecker and Ulrich 2008). In addition to PCNA monoubiquitination, Siz1-mediated PCNA sumoylation is required for TLS stimulation. Epistatic analysis of relationships between SIZ1, MMS2 and RAD5 with respect to UV sensitivity demonstrated that in the absence of PCNA sumoylation, Mms2-Ubc13 and Rad5 independently influence TLS (Halas et al. 2011).

The fraction of sumoylated PCNA increases during S phase, as well as during extensive DNA damage. Sumoylated PCNA is able to bind replicative polymerases, an event that permits synthesis over the lesion. Sumoylated PCNA also has high affinity for Srs2 (Kolesar et al. 2012), whose binding prevents homologous recombination (HR) (Papouli et al. 2005; Pfander et al. 2005). Srs2 is a 3′→5′ DNA helicase (Rong and Klein 1993) that unwinds hairpin intermediates that can be formed by triplet repeats in the DNA and promotes fork reversal in these repetitive sequences, thereby preventing their instability and fragility and consequently contributing to protection against DNA damage.
et al. share the accessory subunits Pol31 and Pol32, one could not easily as expected. This study demonstrated an additional role for Pol32 in TLS beyond its role as a Pol ζ subunit. This newly reported role is related to Pol θ because mutants lacking the C-terminal part of Rev3, responsible for Pol ζ catalytic activity and for interaction with the accessory subunits, are partially proficient in Pol32-dependent UV-induced mutagenesis (Siebler et al. 2014). The polymerase switch scenario was also demonstrated for other polymerases. When Pol η manages to overcome the Tor-T dimers that blocked DNA replication by DNA Pol θ or Pol ε another polymerase switch occurs that allows normal replication to proceed (McCulloch et al. 2004a). Thus, DNA synthesis on templates containing lesions usually requires the cooperation of two polymerases.

Daraba et al. recently demonstrated that this polymerase switch may be achieved by degradation. They showed that upon DNA damage, Def1 promotes the ubiquitination of the Pol13 catalytic subunit of replicative polymerase δ. This ubiquitination is followed by proteasomal degradation. In contrast, the Pol31 and Pol32 subunits of polymerase δ are not affected and are able to form a complex with Rev1; thus, this TLS polymerase may perform DNA lesion bypass at stalled replication forks (Daraba et al. 2014). Ubiquitinated PCNA- and specialized polymerase-dependent TLS. Following DNA damage, PCNA is monoubiquitinated at Lys164 by the Rad6–Rad18 complex (Hoege et al. 2002). This modification enables PCNA to interact with TLS polymerases and provides access to the damaged template (Fig. 4). TLS is performed in various arrangements by polymerases operating independently (e.g. Pol η and Pol ζ) or in tandem (Pol η then Pol ζ or Pol ζ with Rev1). The final result of synthesis depends on the polymerases engaged in the bypass as well as on the specific lesion (Harasck, Prakash and Prakash 2000, 2003; Gibbs et al. 2005; Northam et al. 2006; Parker et al. 2007; Stone, Lujan, Kunkel 2012). Notably, some TLS polymerases bypass specific DNA lesions with perfect reproduction of the initial content of the template, displaying high effectiveness towards the cognate substrate [e.g. Pol η inserts C opposite 7,8-dihydro-8-oxo-guanine (Harascka, Prakash and Prakash 2000) and A–A opposite a T–T dimer (Gibbs et al. 2005)]. Other polymerases synthesize sequences that differ from the original sequence. Moreover, the sequence context of the lesion influences the relative likelihood that specific nucleotides will be used for DNA synthesis by different TLS polymerases, as was shown by Chan, Resnick and Gordenin (2013).

Polyubiquitinated PCNA- and Rad5-dependent damage avoidance pathway. Following DNA damage, Mms2 and Ubc13 proteins are redistributed from the cytoplasm to the nucleus, where they associate with Rad5. The Rad5–Mms2–Ubc13 complex is responsible for ubiquitination of the PCNA Lys164 residue (Ulrich and Jentsch 2000). Ubiquitins in the chain are linked via Lys63; however, it is not known if the polyubiquitin chain is built sequentially or transferred en block on PCNA. Polyubiquitinated PCNA stimulates the error-free PRR damage avoidance pathway (Branzei, Seki and Enomoto 2004; Branzei, Vanoli, Foiani 2008) (Fig. 3). Using this pathway, the lesion in the DNA template can be omitted as follows: (1) by template switch or sister chromatid junctions (SCJ) using the HR proteins Rad51, Rad52 and Rad54, as well as the Sgs1–Top3 complex, permitting resolution of cross-structures generated during repair (Gangavarapu et al. 2006; Branzei and Foiani 2007; Ball et al. 2009); or (2) by Rad5-dependent fork regression (‘a chicken foot’ DNA structure), followed by nascent-strand annealing and DNA synthesis (Blastyak et al. 2007). Both branches of this pathway are error free.

The 9–1–1/RPA-dependent pathway involved in gap filling and telomere maintenance. As previously mentioned, PCNA serves as a platform for various proteins operating at the replication fork. The role of PCNA modification in the tunneling of cellular responses to DNA damage was also described. However, having one type of sliding clamp per cell is not sufficient (Fig. 3). In yeast, another PCNA-like sliding clamp exists that is designed especially for interaction with various proteins activated after DNA damage and engaged in DNA repair and cell-cycle checkpoints. This 9–1–1 complex (the Ddc1–Rad17–Mec3 complex, a homolog of human Rad9–Rad1–Hus1 complex) is a heterotrimeric checkpoint clamp consisting of the Ddc1, Rad17 and Mec3 subunits in S. cerevisiae and the Rad9, Rad1 and Hus1 subunits in Sc. pombe. The 9–1–1 complex is an early response factor to DNA damage that activates checkpoints (Majka and Burgers 2003). It can be loaded onto DNA using two different configurations: at the 5′ end of ssDNA coated with RPA or at both the 5′ and 3′ junctions of naked DNA at gaps left on the lagging strand after replication (Majka et al. 2006a). This complex subsequently stimulates Mec1 kinase and/or checkpoint sensor Dpb11, depending on the cell-cycle phase (Majka, Niedziela–Majka and Burgers 2006b; Navadgi-Patl and Burgers 2009; Puddu et al. 2011).
Figure 4. DNA synthesis on damaged template. The localization of lesion in the DNA template influences the choice of DNA damage repair/tolerance pathway. The damage of already replicated DNA strand is subject to DNA repair by pathway dedicated to deal with particular lesion, such as BER, NER, HR, etc (left panel of the figure). When DNA lesion happens in ssDNA, e.g. in the replication fork (the middle part of the figure) or in ssDNA gaps left after replication (right panel of the figure), the DNA synthesis process has to be applied. However, replication over the lesion is not an easy task, and typically requires switching of polymerases. Various TLS polymerases are able to overcome the lesion that blocked DNA replication; switching back to DNA replicative polymerase resumes normal replication (one TLS polymerase scenario, A). Sometimes, the polymerase that is able to add the nucleotide opposite the lesion is unable to elongate synthesis. Therefore, resumption of synthesis requires the cooperation of three DNA polymerases (B). To fill the gap one more, RPA-dependent step is needed: the recruitment of a sliding clamp to the 3'-OH end of the DNA gap. Then the repair can proceed as described above.

The 9–1–1 complex promotes both Dna2-Sgs1 and Exo1-dependent resection in response to uncapped telomeres (Ngo et al. 2014). The support of resection by the 9–1–1 complex is strongly inhibited by the DNA damage-dependent checkpoint protein Rad9. However, the 9–1–1 complex together with the Exo1 nuclease also participates in the error-free branch of the DNA damage tolerance pathway where it promotes template switching in a manner that is distinct from its canonical checkpoint functions and uncoupled from the replication fork. Recent work by Karras et al. (2013) revealed cooperation in the error-free pathway between the two related clamps PCNA and 9–1–1. The 9–1–1 complex also plays a major role in DNA repair by interacting with and stimulating activity of specialized but not replicative DNA polymerases involved in TLS by filling the gaps left behind the replication fork (Kai and Wang 2003; Cardone, Brendel and Henriques 2008). Additionally, it was shown that the Mcm10 protein that was proposed to play a role in replication fork restart and DNA repair interacts with the Mec3 subunit of the 9–1–1 clamp in response to nucleotide shortages or UV-induced replication stress (Alver et al. 2014). The 9–1–1 checkpoint clamp is also involved in preventing the deleterious effects of dUTP-related apurinic/apyrimidinic sites (called also abasic or AP sites) that may cause replication-related genetic instability (Collura, Kemp and Boiteux 2012). Another role for the PCNA-like sliding clamp is based on its interaction with the Rad51 and Dmc1 proteins, which stimulate efficient repair of meiotic DSBs by facilitating proper assembly of the meiotic recombination complex (Shinohara et al. 2003).

Similar to PCNA, the 9–1–1 complex is ubiquitinated; however, published data disagree on whether this modification is dependent on Rad6 and if Lys197 or Lys164 of Rad17 is the
major acceptor site for ubiquitin. It is also not clear if ubiquitination of the 9–1–1 complex participates in its known functions in DNA repair stimulation and cell-cycle control or merely labels it for proteasomal degradation (Fu et al. 2008; Davies, Neiss and Ulrich 2010). Nevertheless, an interaction between Rad18, the E3 ubiquitin ligase and subunits of the 9–1–1 complex was detected (Fu et al. 2008). Moreover, Fu et al. showed that the Rad6–Rad18 complex involved in PRR ubiquitination is required for increased transcription of a large number of yeast genes in response to DNA damage. They also suggested a crucial role for the 9–1–1 complex in DNA repair and checkpoint control under DNA damage stress, possibly through the recruitment or maintenance of the 9–1–1 clamp at lesion sites.

Surprisingly, a link between the 9–1–1 clamp and nucleosome assembly was recently found. The rad17Δ deletion is synthetic lethal with the deficiency of the histone chaperone Rtt106. Moreover, multiple genetic interactions between the 9–1–1 clamp and DNA replication-coupled nucleosome assembly factors, including Rtt106, Caf-1 and lysine residues of H3–H4 were found. Furthermore, rad17Δ cells showed (1) defects in the deposition of newly synthesized histones H3–H4 onto replicated DNA; (2) a reduction in the Asf1–H3 interaction due to an increased association of Asf1 with checkpoint kinase Rad53; and (3) an increase in the interaction between histones H3–H4 with histone chaperone Caf-1 or Rtt106. All of these results suggest a role for 9–1–1-dependent regulation in DNA replication-coupled nucleosome assembly via histone chaperone interactions (Burgess, Han and Zhang 2014).

Crosstalk between RFC complexes adapts cellular responses to various types of DNA damage The PCNA and 9–1–1 clamps play a crucial role during DNA replication and repair, serving as platforms capable of enabling access to the DNA by various proteins. These clamps are engaged in unwinding DNA, sensing cellular stresses and deciding which repair pathway will be employed under particular conditions. However, the clamps themselves are also subject to regulation. There are several complexes in the cell that control the interaction of PCNA and 9–1–1 with DNA (Fig. 3). The first complex that was shown to possess the ability to load and unload PCNA onto DNA was RFC. In eukaryotic cells, four independent sliding clamp loaders exist (Majka and Burgers 2004). All interact with Mps3, a protein attached to the nuclear envelope, suggesting that these clamp loaders are recruited to the nuclear envelope, e.g. in response to DNA damage (Haas et al. 2012). RFCs are heteropentamers consisting of four smaller Rfc2–5 subunits (Yao et al. 2006) and one large subunit that is unique to each complex. Thus, Rfc1 from the classical RFC complex may be substituted for Ctf4, Elg1 or Rad24, creating alternative RFC complexes (Green et al. 2000; Mayer et al. 2001; Kanelis, Agyei and Durocher 2003). The RFCs work as chromatin-associated factors that load and unload sliding clamps and promote DNA replication and repair; however, each of them has a special predisposition to operate under particular circumstances. The canonical RFC is responsible for PCNA loading and unloading on/off DNA during undisturbed replication (Bowman, O’Donnell and Kuriyan 2004), but it is also required for repair synthesis of large looped heteroduplexes in S. cerevisiae (Corrette-Bennett et al. 2004). Ctf18-RFC and Elg1-RFC can influence PCNA positioning on DNA. Ctf18-RFC unloads PCNA from DNA (Bylund and Burgers 2005) and promotes sister chromatid cohesion, while Elg1-RFC plays an opposing role in this process (Hanna et al. 2001; Maradeo and Skibbens 2009, 2010). Furthermore, Elg1 contributes to Ctf18 recruitment to chromatin. Elg1 and Ctf18 with the aid of Ctf4 may coordinate the relative movement of the replication fork with respect to the cohesin ring (Parnas et al. 2009).

The Elg1–RFC complex interacts effectively with the sumoylated form of PCNA and mediates a decrease in the level of sumoylated PCNA bound to chromatin (Parnas et al. 2010). It is still unclear whether Elg1–RFC actively unloads sumoylated PCNA from chromatin or recruits the demsumoylating enzyme Ulp1 to chromatin to reduce the level of sumoylated PCNA. As was noted in the previous section, PCNA sumoylation prevents recombination in the replication fork. Notably, the elg1Δ mutant displays a hyperrecombination phenotype (Ogiwara et al. 2007) and is synthetic lethal with deletions in genes involved in the HR pathway (Aroya and Kupiec 2005). In turn, Ctf18–RFC is involved in the initiation step of replication (Ma et al. 2010) and replication checkpoint activation (Kubota et al. 2011). Ctf18–RFC is essential for Mrc1-dependent activation of Rad53 and for the maintenance of paused replication forks, most likely by bridging Mrc1 and primed ssDNA fragments that serve as a signal of paused DNA synthesis (Crabbé et al. 2010). The Ctf18–RFC loading clamp facilitates the bypass of triplet repeats during replication (Gellon et al. 2011) to maintain the telomeres (Gao et al. 2014) and is also involved in preserving proper genome copy number in the cell during division (Alabudzinska, Skoneczny and Skoneczna 2011). The 9–1–1 complex is placed on DNA by the dedicated clamp loader Rad24–RFC in a reaction directed by RPA (Majka et al. 2006a). Interestingly, the Rad24–RFC DNA damage checkpoint clamp loader also unloads PCNA clamps from DNA. Therefore, Rad24–RFC may clear PCNA from DNA to facilitate the shutdown of replication when faced with DNA damage (Yao et al. 2006); however, it is less effective at performing this task than is canonical RFC (Thompson et al. 2012). Rad24–RFC functions as a checkpoint protein that is crucial for DNA processing, recombination partner choice and cell survival after various stresses and the subsequent DNA repair (Aylon and Kupiec 2003; Aylon et al. 2003).

Cells lacking one of the alternative RFC complexes exhibit a strong genome instability phenotype (Aroya and Kupiec 2005; Banerjee, Sikdar and Myung 2007). The alternative RFCs channel cellular responses to DNA-borne stress into two major repair pathways: first, employing PCNA to enable replication resumption, and second, employing the cohesion complex to preserve the integrity of chromosomes. Moreover, alternative RFCs function in a cell-cycle-dependent fashion, adjusting the method of repair to the cell-cycle phase and possibly to cell ploidy (Delaçôte and Lopez 2008; Li and Tye 2011). However, the molecular basis through which RFCs exert these diverse effects remains to be uncovered.

RECOMBINATION AND GENOMIC STABILITY

In general terms, DNA recombination can be defined as a DNA transaction pathway in which a strand exchange occurs between different DNA molecules. HR pathway uses a homologous strand as a template for the repair of a damaged strand. The homologous strand might be a sister chromatid, a homologous chromosome or an ectopically located sequence. Non-HR ligates DNA molecules without the step of a homology search for DNA replication priming. Conserved recombination pathways have evolved as a system for reactivating blocked replication forks and for repairing DSBs and interstrand cross-links, lesions that would preclude replication fork progression.
Regulation of the choice between HR and NHEJ

DSBs can arise spontaneously due to endogenous and exogenous factors, such as replication through a nick, metabolism-derived reactive oxygen species (ROS), topoisomerase failures and ionizing radiation. These lesions, however, can also occur during programmed gene rearrangements and other recombination events, such as mating-type switching in budding yeast or during meiosis (reviewed in Symington and Gautier 2011). The DSB repair has been studied extensively in the budding yeast. These lesions are processed through two competing pathways, depending on the cell type, the stage in the cell cycle and the structure of the DNA ends: HR and NHEJ. HR is initiated when the DSB end is resected by nucleases and helicases to generate 3′-ssDNA overhangs. In budding yeast, the binding of the Mre11-Rad50-Xrs2 (MRX) complex and the Sae2 protein to DNA initiates 5′→3′-end resection, which is further processed by Exo1 and Dna2–Sgs1 nucleases (reviewed in Mimitou and Symington 2011). The resulting long ssDNA bound to RPA activates the Mec1 (Sc. pombe rad3 and human ATM) DNA damage checkpoint, which coordinates DNA repair reactions with cell-cycle progression (Zou and Elledge 2003). Thereafter, the various HR processes differ in the processing of intermediates, resulting in different error-free and mutagenic repair pathway outcomes. These HR subpathways are further described below in the section ‘HR subpathways’. In the NHEJ pathway, DSB ends are blocked from 5′-end resection by bound Ku proteins (Yku70 and Yku80), which form a heterodimer. The Ku heterodimer promotes the direct ligation of the DSB ends by the specialized ligase complex Dnl4–Lif1, requiring ‘clean’ DNA ends, i.e. 3′-OH and 5′-phosphate groups, for ligation. If the DSB ends are not clean, then additional processing by nucleases and a dedicated polymerase (Pol α) are needed for ligation to occur, and during this process, small deletions and insertions might be introduced at the junction site. Therefore, this pathway is considered error prone. Indeed, many lines of evidence suggest that in budding yeast increased mutagenesis can be dependent on NHEJ (Halas et al. 2009; Lehner et al. 2012; Shor, Fox and Broach 2013). HR and NHEJ are separate pathways that are differently regulated (see below), but the MRX complex is shared by both pathways, because, in addition to its important role in HR, MRX also functions as a DNA end-bridging factor in NHEJ (Chen et al. 2001). Nevertheless, Ku proteins and Dnl4 inhibit DNA end resection (Zhang et al. 2007), and, in turn, DNA end resection inhibits NHEJ ligation, reflecting the end requirements of Dnl4–Lif1-mediated ligation. Consistently, defects of sae2 null cells in DSB processing after exposure to ionizing radiation are suppressed by inactivation of YKU70, and this suppression requires both nucleases mediating the extensive end resection, Exo1 and Sgs1–Dna2 (Mimitou and Symington 2010). As previously mentioned, these two pathways are also differently regulated in budding yeast cells according to cell type and cell-cycle stage. In haploid cells, both HR and NHEJ pathways are efficient, whereas in diploid cells NHEJ is inhibited due to the mating-type control-mediated repression of the NEJ1 (LIF1) gene encoding the critical NHEJ regulator (Frank-Vaillant and Marcand 2001; Kegel, Sjöstrand and Aström 2001). The choice of a DSB repair pathway is also dependent on the phase of the cell cycle. In both haploid and diploid cell types, DSB repair through HR primarily occurs during the S and G2 phases when DNA has replicated and one sister chromatid is available as a template to repair the other sister chromatid. This control is mediated through cyclin-dependent Cdk1 kinase, which phosphorylates many DNA repair proteins in the cell (reviewed in Ensink and Kolodner 2010). However, the most critical step to initiate HR involves overcoming the Ku-mediated inhibition of end resection (reviewed in Symington and Gautier 2011). This is accomplished through two Cdk1-dependent mechanisms: one pathway that is associated with the regulation of Ku/Dnl4-Lif1 affinity for DNA ends (Zhang et al. 2009) but has not yet been characterized, and another pathway involving the activation of Sae2 through phosphorylation at Ser267 (Huertas et al. 2008). Interestingly, the CDK-dependent regulation of Ctp1, the human ortholog of Sae2, is evolutionarily conserved (reviewed in Symington and Gautier 2011). The requirement for the early end resection HR factors, Mre11 and Sae2, depends on the structure of the DNA ends in budding yeast. These factors are not required for the resection of clean DSB ends, but are necessary for the resection of dirty or complex ends, induced through ionizing radiation, DSB ends adopting secondary structures or DSBs with a protein covalently bound to the 3′ end, e.g. the binding of topoisomerase to a DNA end after treatment with camptothecin (reviewed in Symington and Gautier 2011). The end resection step during DSB repair has also been studied in fission yeast, and the mechanism is essentially conserved between the two yeast species, although there are some interesting differences (Langerak et al. 2011). Similar to the HR in S. cerevisiae, in fission yeast cells defects in the initiation of end resection are suppressed in the absence of Ku proteins.

HR subpathways

The RPA-coated ssDNA generated during the extensive end resection mediated through HR nucleases is processed via several HR subpathways, depending on the homologous DNA availability (reviewed in San Filippo, Sung and Klein 2008). These sub-pathways include (1) the double HJ pathway, also known as the DSBs repair pathway, (2) synthesis-dependent strand annealing (SDSA), (3) break-induced replication (BIR), (4) single-strand annealing (SSA) and (5) the error-free lesion bypass pathway. In these pathways, except SSA, RPA covering ssDNA is replaced by the Rad51 recombinase in a reaction mediated by Rad52 (Sung 1997). The Rad51 protein combined with ssDNA forms a nucleoprotein filament that catalyzes homologous pairing and strand invasion into a homologous duplex (reviewed in San Filippo, Sung and Klein 2008). The invading strand is used by a DNA polymerase for priming DNA synthesis to extend the ensuing displacement loop (D-loop) with the assistance of the translocases Rad54 or Rdm54 (reviewed in San Filippo, Sung and Klein 2008; Symington, Rothstein and Lisby 2014).

In the double HJ pathway, the strand displaced by the extending D-loop anneals to the second ssDNA overhang on the other side of the DSB (second-end capture) and primes DNA synthesis to fill in the gap. Ligation generates double HJs (crossed-strand structures). The resolution of double HJs by structure-selective endonucleases (resolvases) in different orientations leads to either crossover or non-crossover of flanking markers. HJ intermediates are also dissolved by branch migration, which is mediated by the Sgs1-Top3-Rmi1 (STR) complex (reviewed in Ashton et al. 2009, but has not yet been characterized, and another pathway involving the activation of Sae2 through phosphorylation at Ser267 (Huertas et al. 2008). Interestingly, the CDK-dependent regulation of Ctp1, the human ortholog of Sae2, is evolutionarily conserved (reviewed in Symington and Gautier 2011). The requirement for the early end resection HR factors, Mre11 and Sae2, depends on the structure of the DNA ends in budding yeast. These factors are not required for the resection of clean DSB ends, but are necessary for the resection of dirty or complex ends, induced through ionizing radiation, DSB ends adopting secondary structures or DSBs with a protein covalently bound to the 3′ end, e.g. the binding of topoisomerase to a DNA end after treatment with camptothecin (reviewed in Symington and Gautier 2011). The end resection step during DSB repair has also been studied in fission yeast, and the mechanism is essentially conserved between the two yeast species, although there are some interesting differences (Langerak et al. 2011). Similar to the HR in S. cerevisiae, in fission yeast cells defects in the initiation of end resection are suppressed in the absence of Ku proteins.

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Rad51 presynaptic filaments (Krejci et al. 2003; Veaute et al. 2003). As mentioned in section ‘Postreplication repair: a specialized pathway to repair replication stall-borne problems’, Srs2 binding to SUMOylated PCNA also limits the access of DNA polymerases to recombination intermediates in a mechanism independent from the Srs2 interaction with Rad51 (Burkovics et al. 2013). Second, the Mph1 helicase dissociates Rad51-dependent D-loops with high efficiency (Prakash et al. 2009).

In both the double HJ and SDSA pathways, DNA replication utilizes only leading-strand synthesis machinery to extend both the first and second ends (see below). Single-ended DSBs, e.g. at telomeres or broken replication forks, are resected similarly to two-ended DSBs, and the resulting 3′-ssDNA overhangs invade a homologous duplex and prime DNA synthesis only from one side. BIR continues to the end of the invaded duplex and engages the machinery of both leading- and lagging-strand DNA synthesis (Lydeard et al. 2010). BIR leads to the non-reciprocal exchange of a part of a donor chromosome to a recipient chromosome (Kraus, Leung and Haber 2001); however, this pathway is also highly mutagenic for other reasons, as described below. Sometimes a DSB is flanked by direct repeats. Following end resection, exposed complementary 3′-ssDNA overhangs anneal, forming a duplex in the new strand configuration, with the concomitant deletion of the intervening sequence. This is a SSA pathway, which is mechanistically different than the other HR pathways, because it does not require Rad51 for homologous strand pairing. However, the strand annealing reaction is not spontaneous (it is inhibited by RPA bound to ssDNA); it is mediated by an additional Rad52 protein function that is specific for ssDNA annealing (reviewed in Symington, Rothstein and Lisby 2014). This pathway also requires the activity of the Rad1–Rad10 endonuclease (Fishman-Lobell and Haber 1992), which cuts branched DNA structures at the transition between dsDNA and ssDNA to remove ssDNA flaps that fail to anneal (reviewed in Lyndaker and Alani 2009). The SSA pathway is efficient, even for end resection and annealing between short (5–25 nt) direct repeats flanking a DSB. This ‘small’ variant has been described as a microhomology-mediated end joining (MMEJ) pathway in Ku-deficient cells (Ma et al. 2003). However, the MMEJ pathway is not well characterized and its genetic requirements remain controversial (see also Symington and Gautier 2011). In S. cerevisiae, this ‘small’ SSA variant might have a modified requirement for annealing (reviewed in Decottignies 2013) because instead of Rad52, Rad59 is necessary for the annealing of short homologous sequences (Sugawara, Ira and Haber 2000). An SSA-like pathway has also been studied in fission yeast as a microhomology-mediated recombination pathway repressed by Ku proteins (Decottignies 2007). Indeed, repetitive elements are numerous in yeast genomes, consequently SSA could mediate large-scale rearrangements that cause deletions of sequences located between the repeats.

The last HR subpathway for error-free lesion bypass is one of the two main mechanisms of DNA damage tolerance (the other mechanism is TLS, which occurs either in an error-free or error-prone manner as mentioned in the section ‘Postreplication repair: a specialized pathway to repair replication stall-borne problems’) in response to stalled replication forks. Various endogenous and exogenous factors might cause replication fork stalling (reviewed in Aguillera and Gómez-González 2008; Branzei and Foiani 2010), including secondary DNA structures often adopted by repetitive sequences, highly transcribed DNA, tightly bound proteins forming replication fork barriers and a diverse array of template lesions. As mentioned in section ‘Postreplication repair: a specialized pathway to repair replication stall-borne problems’, a recombination-dependent mechanism mediates transient template switching, which uses the sequence of the newly synthesized sister chromatid to bypass the lesion, providing adequate time for damage removal and facilitating both gap filling and the restart of replication forks stalled by replication stress. Template switching is associated with the formation of SCJ, in the proximity of replication forks, that are cruciform pseudo-double HJ structures (Liberi et al. 2005; reviewed in Vanoli et al. 2010). These recombinant intermediates accumulate in a Rad51/Rad52-dependent manner in MMS-treated cells lacking the Sgs1 helicase. The pathway utilizes a subset of recombination proteins (Rad51, Rad55, Rad57, Exo1 and Rad59) and both replicative polymerases, Pol ε and Pol δ (Vanoli et al. 2010). Recently, template switch intermediates have been characterized using the 2D gel electrophoresis and electron microscopy (Giannattasio et al. 2014). The choice between different bypass mechanisms is regulated by ubiquitin and SUMO modifications of PCNA at sites of stalled replication forks (reviewed in Moldovan, Pfander and Jentsch 2007) and mentioned in the previous sections). In S. cerevisiae, the template switch mechanism depends on the Rad18 ubiquitin ligase- and Rad5-mediated polyubiquitylation of PCNA, whereas Ubc9-dependent SUMOylation of PCNA and Mms21-dependent SUMOylation of Smc5-Smc6 and Sgs1-Top3 complexes are required to remove the cruciform recombinant intermediates (Liberti et al. 2005; Branzei et al. 2006; Branzei, Vanoli, Foiani 2008; Sollier et al. 2009). Thus, this process is strictly controlled, highlighting its importance for the maintenance of genome stability in conditions of replication stress. Interestingly, González-Prieto et al. (2013) have found evidence for the Rad52-mediated recruitment of Rad51 to chromatin during the unperturbed replication. Upon exposure to an alkylating agent, the recruited Rad51 facilitates DNA synthesis on a damaged template in a DSB repair-independent process, most probably by promoting gap filling of single-stranded lesions that accumulate during replication of alkylated DNA. Future research will determine how the Rad52-dependent recruitment of Rad51 to unperturbed replication forks is linked to the above-mentioned lesion bypass mechanisms. As an aside, in higher eukaryotic cells, the Rad51 ortholog and other HR factors are indispensable for DNA replication, and the inactivation of genes encoding these proteins is lethal, whereas equivalent recombination proteins are dispensable for viability in both yeast species (reviewed in San Filippo, Sung and Klein 2008; Errico and Costanzo 2010).

Mutations dependent on recombination

Although HR is typically an error-free process, error-free recombination products are only produced during allelic-strand exchanges between sister chromatids, and as mentioned above, recombination between sister chromatids in the S and G2 phases of the cell cycle is favored by the cell-cycle mechanisms that regulate recombination. Aside from recombination between sister chromatids, there are many sources of genetic alterations in the final recombination products.

First, Rad51-mediated strand invasion of homologous sequences containing few heterologies can occur, thus generating a heteroduplex DNA molecule bearing mismatches typically processed by the MMR pathway (Chen and Jinks-Robertson 1998). The MMR-mediated repair of these mismatches results in gene conversion, defined as the unidirectional (non-reciprocal) transfer of information from one DNA duplex to another. If mismatches are not repaired, then the mismatched strands will segregate during the next round of replication, resulting in a
sectored colony, or when there are too many mismatches in the heteroduplex DNA intermediate, the MMR-mediated anti-recombination pathway blocks the formation of recombinant products (heteroduplex rejection). Thus, the final recombinants could include gene conversion changes introduced by the MMR of heteroduplex DNA molecules initiating recombination.

Second, the two HR pathways, double HJ and SDSA, share a common intermediate, the D-loop, in which only donor (displaced) strands are copied (Ira, Satory and Haber 2006). The modified replication forks in the D-loop use only the leading-strand synthesis machinery to extend the first and second ends. Importantly, D-loop extension does not require the replicative helicase complex comprising the heterohexameric Mcm2–7 (MCM) helicase proteins and Cdc45 (and presumably GINS proteins) (Wang et al. 2004). However, a recent study shows that the proper functioning of GINS in the replisome is necessary for the fidelity of DNA replication in budding yeast (Grabowska et al. 2014). Accordingly, several lines of evidence suggest that the replication forks in D-loops, which form during HR-mediated DSB repair, replicate DNA with significantly decreased fidelity (reviewed in Malkova and Haber 2012). In the region surrounding an induced DSB, mutations arise at a rate 1000-fold higher than the spontaneous mutation rate. Although the majority of these mutations are base substitutions, other types of mutations can occur through multiple microhomology-mediated template switches between various, even highly diverged, sequences during replication, indicating that D-loop replication is less effective than normal replication. Consistently, template switch mutations are eliminated in strains with a proofreading-defective Pol δ because defective polymerases are less likely to dissociate from the template than the wild-type enzyme. Thus, mutations also result from errors in the replicative polymerases Pol δ and Pol ε. In addition, the mutations detected in the region of the DSB repair where both strands have been newly synthesized are not dependent on either TLS polymerases, Pol ζ and Pol η, or Pol32, despite the dependence on Pol δ, or the MMR system (Hicks, Kim and Haber 2010). During DSB repair, DNA replication also occurs at regions outside the D-loop, where extensively resected ssDNA is filled in to complete the repair. This replication is also mutagenic, generating an increased proportion of frameshifts rather than base pair substitutions. The base pair substitutions are associated with the Pol ζ-mediated fill-in replication of ssDNA, and the frameshift mutagenesis is halved in rev3 null strains. Therefore, multiple polymerases are responsible for the mutagenesis occurring during fill-in replication. In addition, MMR defects do not change either the rate or the spectrum of the mutations (Hicks, Kim and Haber 2010). It is not clear why the MMR system is active on the heteroduplexes formed during strand invasion in HR pathways, but becomes inactive once the DNA replication associated with HR is initiated. Nevertheless, MMR is temporally coupled to replication in the S phase, whereas heteroduplex recombination is largely cell cycle independent (Hombauer et al. 2011). It is possible that the replication associated with recombination lacks a cell-cycle-dependent factor for the specific activation of MMR in the S phase. Interestingly, Rattray et al. (2002) showed that in the absence of Rad57, the DSB-induced recombination of inverted repeats significantly increases mutagenesis associated with repair events. Rad57 forms a heterodimer with Rad55, which mediates Rad51-dependent homologous DNA pairing and strand exchange, as an alternative to the Rad52 mediator function (Sung 1997) mentioned in the section ‘HR subpathways’. Therefore, it is conceivable that after some template switching events during DNA replication associated with DSB repair, the polymerase resumes replication on the correct template after a reiterated engagement of the extending strand with Rad51, suggesting that in budding yeast cells, the function of Rad51 contributes to the fidelity of DNA replication, as mentioned in the section ‘HR subpathways’ in relation to the recombination-dependent error-free lesion bypass.

The fidelity of DNA replication associated with DSB repair has also been studied in the context of concomitant exposure to MMS or UV and a short oligonucleotide template for repair (Yang et al. 2008). In this system, extensive end resection typically proceeds after DSB induction, but only a small region in the immediate vicinity of the DSB is available for D-loop replication, resulting in the persistence of ssDNA next to the DSB that has to be filled in. Not surprisingly, treatment with MMS or UV leads to a further 10-fold increase in mutagenesis, but in addition, in the region that has been rendered single-stranded prior to treatment, mutations occur in widely separated clusters of multiple mutations. The UV-induced mutations also show a strand-specific bias. Thus, lesions in exposed ssDNA are hypermutagenic. As mentioned above, although the active replication forks in BIR require most of the S-phase replication proteins (Lydeard et al. 2010), BIR is highly inaccurate and, in contrast to two-sided DSB events, the lack of replication fidelity extends over the entire path of the replication fork (Deem et al. 2011). The rate of frameshift mutagenesis increases 2500-fold during BIR compared to mutagenesis during normal replication, although MMR and proofreading activities are proficient. The frameshift mutations suggest that replication becomes unstable due to template-switching errors. Interestingly, the replication fork generated during BIR is unusual. The replication proceeds in a PiF1-dependent migrating D-loop structure that is extended by Pol δ (Saini et al. 2013; Wilson et al. 2013). Similar to the results mentioned above, BIR intermediates are a strong source of clustered mutagenesis induced after MMS treatment (Sakofsky et al. 2014). Furthermore, in conditions of replication stress or premature mitosis due to checkpoint deficiency, BIR leads to a surge of GCR events. The results of research on BIR mechanisms are particularly significant because these experiments model the fates of collapsed forks or eroded telomeres in the cell, suggesting that failed replication can lead to genome rearrangements and clustered mutagenesis. Studies in fission yeast have shown similar results on BIR. Iraqui et al. (2012) reported that a single collapsed fork leads to mutations and large-scale genomic rearrangements (deletions and translocations) due to erroneous DNA synthesis during the recovery of replication forks.

The third source of genetic modifications during the HJ pathway, but not the SDSA pathway, is the resolution step, in which resolvases catalyze the cleavage of HJ intermediates to generate either non-crossover or crossover (reciprocal exchange of the flanking sequences) products. The double HJ, leading to crossover in the final recombination products, does not have genetic consequences when the HJ intermediates have linked sister chromatids in the same allelic positions. However, under other circumstances, the resolution may result in genetic rearrangements, e.g. LOH in mitotic diploid cells when the HJ intermediates have linked homologous chromosomes. There are three mitotic resolvases in S. cerevisiae (reviewed in Sarbajna and West 2014): Yen1 (the homolog of the human GEN1 protein), Mus81–Mms4 and Sbx1–Sbx4 (however, the report by Muñoz-Galván et al. 2012 questioned the nuclease activity of Sbx1 in budding yeast). These enzymes differ in the range of branched molecules processed and the products of the catalyzed HJ cleavage. The results of several studies on the regulation of S. cerevisiae resolvases have indicated a number of mechanisms by which cells control the actions of these enzymes.
structure-selective nucleases during mitosis and meiosis via cell-cycle-dependent phosphorylation. For example, there is strong evidence for cell-cycle regulation of joint molecule resolution during recombination-based template switching events between sister chromatids (error-free lesion bypass mentioned in sections ‘Postreplication repair: a specialized pathway to repair replication stall-borne problems’ and ‘HR subpathways’). The pivotal element in this control circuit is the conserved Dpb11–Slx4 complex (in Sc. pombe, Rad4/Cut5 and Slx4, respectively) of scaffold proteins that control the joint molecule resolution between sister chromatids mediated by the Mus81–Mms4 endonuclease. Cell-cycle-dependent phosphorylation of Slx4 by Cdk1 promotes the Dpb11–Slx4 interaction. In mitosis, at the appropriate time to resolve joint molecules before cell division, phosphorylation of Mms4 by Polo-like kinase Cdc5 (in Sc. pombe Plol) stimulates the association of Mus81–Mms4 with the Dpb11–Slx4 complex, thereby promoting joint molecule resolution (Gritenaite et al. 2014). To close the regulatory circuit, the DNA checkpoint inhibits the interaction of Mus81–Mms4 with Dpb11–Slx4 and consequently prevents resolution (Gritenaite et al. 2014). Another line of recent research concentrates on the regulation of HJ resolution in meiosis, because crossovers are essential for the establishment of cohesion-mediated interactions between homologs and the ultimate segregation of these molecules (Matos et al. 2011; Matos, Blanco, West 2013; reviewed in Sarbajna and West 2014). Furthermore, there is a meiosis-specific pathway for crossover production that is presumably dependent on the recently described heterodimeric Mih1–Mih3 endonuclease (Ranjha, Anand and Cejka 2014; Rogacheva et al. 2014; reviewed in Kohl and Sekelsky 2013). The metabolism of HJs in the fission yeast is notably different from that of the budding yeast, since Sc. pombe cells lack a clear Yen1/GEN1 homolog and depend on the Mus81–Eme1 (the fission yeast orthologs of Mus81 and Mms4) resolvase for HJ processing in mitosis and meiosis. For example, Mus81–Eme1 is upregulated by DNA damage through Rad3 (human ATR; S. cerevisiae Mec1)–and Cdc2/Cdk1-dependent phosphorylation of Eme1 (Dehè et al. 2013), whereas, in S. cerevisiae, cell-cycle-dependent phosphorylation of Mus81–Mms4 is inhibited by DNA-damaging agents, because upon DNA damage the activated Mec1 checkpoint pathway delays expression/activation of cell-cycle kinases and cell-cycle progression. However, HJ resolution by endonucleases is not promoted in unchallenged mitotic cells. In S. cerevisiae during mitosis, HJs are rather processed by non-crossover dissolution mediated by the STR complex, as mentioned in the section ‘HR subpathways’. The STR complex suppresses crossover formation (Gangloff, Soustelle and Fabre 2000; Ira et al. 2003) and the accumulation of recombiant intermediates at damaged replication forks (Liberi et al. 2005). Biochemical studies, initially for the human ortholog BLM complex and subsequently for the S. cerevisiae complex, have shown that indeed the BLM/Sgs1 complex dissolves double HJ intermediates into products that are exclusively non-crossover (Wu and Hickson 2003; Raynard, Bussen and Sung 2006; Cejka et al. 2010). Surprisingly, the conserved Sgs1 helicase (the Sc. pombe ortholog Rqh1) has multiple functions linked to recombination-related DNA transactions. In mitotic cells, in addition to being a component of the crossover-suppressing STR complex, the helicase functions also as a prerecombination protein in end resection, as described in the section ‘HR subpathways’.

Recombination is a double-edged sword: it facilitates DNA replication and protects genome stability, but it can also lead to genomic rearrangements. In fission yeast, recombination promotes cell viability under replication stress at the expense of genetic stability (Lambert et al. 2005). Therefore, recombination must be tightly controlled through multiple mechanisms, such as those listed above, and coordinated with the cell-cycle checkpoint response (reviewed in Labib and De Piccoli 2011). Last but not least, structural maintenance of chromosomes (SMC) complexes, cohesin, condensin and Smc5–Smc6 are indispensable for chromosome organization, DNA transactions and cell viability (reviewed in Jepsson et al. 2014). The SMC complexes have already been implicated in the regulation of DSB repair by HR (reviewed in Symington, Rothstein and Libby 2014), whereas the Smc5–Smc6 complex in particular has been shown to be required for preventing the accumulation of inappropriate recombination structures arising during mitosis under replicative stress (Sollier et al. 2009; Choi et al. 2010) and during meiosis (Copsay et al. 2013; Lilienthal, Kanno and Sjögren 2013; Xaver et al. 2013). Future studies will focus on understanding how these factors link recombination and the maintenance of chromosome structure.

THE INFLUENCE OF PLOIDY ON GENOME STABILITY

One important aspect of the optimization of genome maintenance processes is the number of copies of genetic material found in various organisms or during various growth stages of a given organism. Having more than one copy of the genome helps to create a ‘reservoir of variability’ that is beneficial for adaptation. Second copy of the genome is also crucial for a HR, the usually accurate DNA repair mechanism. Therefore, it is not surprising that most eukaryotes are diploid or polyploid for the most of their lives. However, having multiple copies of the genome also has a downside. Sometimes, especially under conditions that cause DSBs, it allows for genome rearrangements that can result in excessive genome shuffling with disastrous rather than beneficial effects on the survival. In haploid cells, the presence of a second copy of the genome that enables HR is limited to the S and G2 phases of the cell cycle when DNA is duplicated. During G1, rearrangements may persist in the cell only if they do not perturb essential genes. Thus, sensitivity to DSBs is greater in haploid cells than in diploid cells. The distinction in mutation frequencies between haploid and diploid cells becomes comprehensible when the disparity in their DNA rearrangement survival rates and the differences in the availability of homologous sequences in their genomes are considered. In diploid S. cerevisiae cells, the frequency of spontaneous mutagenesis in forward mutation assays at CAN1/can1Δ and URA3/ura3Δ heterozygous markers is two orders of magnitude higher than in respective loci in haploid cells (approximately 10–4 versus 10–6, respectively) (Hiraoka et al. 2000; Ohnishi et al. 2004; Alabrudzinska, Skoneczny and Skoneczna 2011). The frequency of point mutations, such as base substitutions or frameshifts, remains similar in both cell types. The higher mutagenesis level in diploid cells is due to recombination events, including gene conversion, crossover or chromosome loss, as revealed by mutation spectra analysis (Hiraoka et al. 2000; Ohnishi et al. 2004). These differences may suggest that genome maintenance mechanisms specific for various ploidy levels exist. However, this part of the genome stability story needs further experimental support. One of the first results showing differences in budding yeast cellular responses to DNA damage with respect to ploidy were published by Li and Tye (2011). The authors showed that the mcm4-Δmcm4′ mutation leading to an MCM helicase defect caused replication stress in both haploid and diploid cells. However,
only diploid mutants exhibited a G2/M transition delay, displayed signs of severe genetic instability and subsequently, reduced viability. These unforeseen outcomes are due to differences in repair pathways choice; while haploid cells use the Rad6-dependent pathways that resume stalled replication forks, diploid cells use the DSB repair pathways that depend on the Rad52 and MRX complex (Li and Tye 2011). A study of the nature and frequency of chromosomal rearrangements at a marker gene placed in the telomeric region showed a decrease in chromosome rearrangement frequency and an increase in the complexity of the rearrangements occurring at the target gene with the increase in ploidy level. The presence of short DNA tandem repeat sequences seems to be a key requirement for deletion and reciprocal translocation processes to occur in diploids (Tourrette et al. 2007). A genomic screen for sensitivity to the Top2 inhibitor doxorubicin revealed a substantial difference between haploid and diploid cells in response to this agent. Experiments in which cells from homoygous diploid and haploid yeast gene deletion clone collections were exposed to doxorubicin revealed much higher levels of damage and cell death or severely reduced growth fitness (up to 5-fold) of diploid mutant clones compared to their respective haploid mutant clones. In addition, the toxic effect of doxorubicin was observed only in diploid cells of strains lacking such genes as bem1Δ, ctf4Δ, ctk1Δ, hfl1Δ, mup133Δ or tho2Δ. These mutant strains displayed severe G1/S phase cell-cycle progression defects following doxorubicin treatment, and some of the mutant strains were either significantly enhanced (ctk1Δ and hfl1Δ) or deficient (tho2Δ) in recombination (Westmoreland et al. 2009). As mentioned in the section ‘Regulation of the choice between HR and NHEJ’, the efficiency of NHEJ is significantly reduced in diploid cells due to mating-type locus-dependent regulation of NHEJ activity. Because the choice of one DSB repair pathway over the other may secure or endanger stable genome maintenance, the time window for NHEJ activity is limited to the G1 phase even in haploids. In contrast, HR, in which the sister chromatid is used to efficiently repair DSBs, occurs during the post-replication stage. Delacôte and Lopez (2008) suggested that the association between the cell-cycle checkpoint and the appropriate DNA repair pathway determines the maintenance of genome stability. In the deletion strains that are specifically sensitive to doxorubicin as diploid homozygous strains, this sensitivity is clearly correlated with defects in G1/S phase cell-cycle progression. Doxorubicin-induced DSBs produced in the G1 phase are thus not repaired by NHEJ because the DNA checkpoint does not function properly to compensate for the decreased efficiency of NHEJ in diploids and, consequently, the DSBs can progress through S phase and be processed by HR in the late S/G2 phase. However, in this case, HR cannot use the sister chromatid, which is also broken at the same locus, but must use ectopic homologous sequences dispersed in the genome, leading to genetic instability.

DIVISION ABNORMALITIES

Whatever the reason for DNA content change, it always arises in the progeny cells as a consequence of cell division imperfections, resulting in another enormous reservoir of genetic variability. Factors such as unrepaired DNA damage, faulty mitotic spindle organization, functioning or positioning, defective septum organization or perturbations in mitotic checkpoints may affect the final DNA content of the cells. Abnormal cell division drives two possible anomaly types: aneuploidization and ploidy shift.

Chromosome instability in yeast

Having two copies of the genome, diploid cells are oversensitive to factors causing rearrangements. For example, DSBs stress can lead to changes in the DNA sequence, including aneuploidization (i.e. the gain or loss of individual chromosomes). The latter phenotype can also result from dysfunction of genes involved in DSB recognition, signaling or repair. Recent discoveries reveal further causative agents that may lead to aneuploidy in S. cerevisiae, such as hydroquinone-induced delay at G2/M transition checkpoint, which is normally activated by the Hog1-Swe1 pathway (Shiga et al. 2010). Chromosome instability (CIN) is a widely observed phenotype in yeast and other fungi, including different pathogenic species (Hughes et al. 2000; Infante et al. 2003; Morrow and Fraser 2013; Harrison et al. 2014). Interestingly, various species differently tolerate aneuploidy, e.g. Sc. pombe are primarily euploid (with a species-specific number of chromosomes), because aneuploid cells of this organism display low viability (Mońlar and Szűcs 2013). The tolerance of CIN is influenced by two factors: (1) the character of the change, i.e. chromosome gain or loss events, and (2) the chromosome type, i.e. which chromosome is affected. In S. cerevisiae, monosomy (loss of one chromosome from diploid genome) can be tolerated, particularly in the case of chromosome III (Haber 1974). In standard laboratory conditions, the spontaneous rate of individual chromosome loss in S. cerevisiae varies between 10⁻⁴ and 10⁻⁶ events per cell division (Hartwell and Smith 1985; Klein 2001; Kumaran, Yang and Leu 2013). The same rate of chromosome loss has been observed in Sc. pombe (Bodi, Gysler-Junker and Kohli 1991). Notably, there is no strong correlation between the frequency of chromosome loss and the size of the chromosome. The largest yeast chromosome, XII, is lost at a frequency similar to chromosome III, one of the smallest chromosomes of S. cerevisiae. Both chromosomes show a much higher loss frequency compared with other chromosomes of similar sizes (Kumaran, Yang and Leu 2013). The frequency of chromosome XV gain under normal conditions is approximately 10⁻²–10⁻³ events per cell division. After antifungal drug treatment, the frequency increases by approximately one to two orders of magnitude (Howlett and Schiestl 2009). Diverse stresses can induce CIN. Proteotoxic stress caused by transient Hsp90 inhibition or heat shock markedly increased CIN producing cell populations with various karyotypes. This elevated CIN is most likely linked to the chaperone role of Hsp90 in the kinetochore complex assembly (Stemmann et al. 2002). Cells exposed to Hsp90 inhibitor exhibit changes in chromosome XV, resulting in a multi-drug-resistant phenotype and disturbed chromosome stoichiometries. These results strongly suggest that aneuploidy is a form of stress-induced mutation capable of fueling rapid phenotypic evolution and drug resistance and may be related to a Hsp90-dependent adaptive mechanism under stress conditions (Chen et al. 2012). However, Yona et al. showed that chromosomal duplications are only the first ‘quick fix’ reaction in response to acquired stress. They suggested that aneuploidy is only a transient stage involving a short-lived intermediate that facilitates further adaptation and gives cells time to develop more refined and sustainable solutions (Yona et al. 2012). This may explain why aneuploidy is so widespread among yeast strains (Hughes et al. 2000).

The concept of the selective advantage of chromosome gain or loss under various stresses was systematically analyzed in S. cerevisiae by two groups (Pavelka et al. 2010; Sheltzer et al. 2011). Experimental data showed numerous examples of aneuploidy in yeast cells, but regardless of the origin of this
biological phenomenon, the possession of more chromosomes did not necessarily benefit the yeast. The addition of an extra copy of a single chromosome initially decreased cellular fitness. Cells displayed increased chromosome loss, elevated mitotic recombination and defective DNA damage repair, causing aneuploidy-induced genomic instability that could facilitate the development of subsequent genetic alterations and drive genome instability (Sheltzer et al. 2011). However, quantitative growth assays revealed that some aneuploid strains with distinct karyotypes and genome contents up to 3n showed significantly improved growth compared with isogenic euploid strains when cultured in parallel and subjected to various suboptimal growth conditions or the presence of a panel of chemotherapeutic or antifungal drugs. Aneuploidy directly affects the cell proteome and generates significant phenotypic variations that could help the cell adapt to changing conditions. Thus, aneuploidy can drive evolution (Pavelka et al. 2010). It was also demonstrated for several independent translocation events that despite their common origin from the integration of the same linear DNA construct, each translocation mutant strain had a different phenotype, sporulation rate, level of gene expression and cell morphology. These phenotypic variations are the result of applying various methods to cope with an initial translocation event in individual yeast cells (Rossi, Noel and Bruschi 2010).

In parallel with these findings, the hypothesis was proposed for the fungi Candida albicans claiming aneuploidy as a form of large-scale mutation able to confert adaptive phenotypes under diverse stress conditions (Zacchi et al. 2010). Candida albicans strains lacking two of the four histone H4 variants showed a decrease in histone H4 dosage followed by a severe growth defect, unstable colony morphology and the production of fast-growing morphologically stable suppressors. These suppressors displayed an increased histone H4 gene copy number due to partial or whole chromosomal trisomies. In budding yeast, hypoacetylated histone H4 is present at centromere regions associated with the Cse4 protein. Inhibition of histone deacetylases in cse4 or hhf1–20 kinetochore mutants is lethal. An increase in the acetylation of H4 in a sir2 (histone deacetylase) mutant or in cells overexpressing deacetylase Sas2 led to increased rates of chromosome loss and to synthetic dosage lethality in kinetochore mutants. Therefore, the hypoacetylation of H4K16 at centromeres plays an important role in accurate chromosome segregation (Choy et al. 2011). In contrast, a malfunction in sister chromatid cohesion led to chromosome gain (Covo et al. 2014).

Other mechanisms, in addition to exogenous stress, could also lead to aneuploidization. Alterations in chromosome segregation can happen due to failed control of DNA quality (replication checkpoint) or cell division (mitotic checkpoint), improper spindle organization or orientation and defects in attachment of chromosomes to spindle filaments or in system correcting misattachments (spindle assembly checkpoint), as well as deficiencies in the mechanisms of chromosome segregation (anaphase-promoting complex checkpoint), cytokinesis or septation (Suijkerbuijk and Kops 2008; McCulley and Petes 2010; Thompson, Bakhoun and Compton 2010; Silva et al. 2011; Stirling, Crisp et al. 2012). The proteins engaged in the complex process of mitosis frequently influence the final DNA content in daughter cells. Abnormal DNA content imposed by defects in mitosis can be repaired, provided that the causative agent is eliminated. Otherwise, a dividing aneuploid cell undergoes mitotic arrest triggered by unbalanced DNA content and generates a damage signal (prolonged spindle assembly checkpoint activation) for its elimination by mitotic catastrophe (Vitale et al. 2011). If cell death is not executed at this point, various scenarios may occur, including senescence followed by cell death, continuation of aberrant divisions leading to aneuploidization or cell death (depending on whether daughter cells receive all essential genes), mitotic slippage or cytokinesis failure leading to polyploidization. For example, in S. cerevisiae failure to establish the cohesion complex in ctf18Δ or eco1 mutants causes aneuploidization (Spencer et al. 1990; Kouprina et al. 1993; Unal, Heidinger-Pauli and Koshland 2007). However, inability to dissolve the cohesion complex triggers polyploidization (Bermúdez-López et al. 2010). Lack of the ESP1 gene encoding separase or an excess of Pds1 encoding securin aborts sister chromatid separation and also causes polyploidization (Lu and Cross 2009). Mitotic exit network components directly control cytokinesis by targeting the Inn1, Cyk3 and Cha2 proteins to the bud neck. Dysfunction of any of these proteins results in aneuploidization (Meitinger et al. 2010). Another process that contributes to proper chromosome segregation involves sensing chromosome attachment to the mitotic spindle (reviewed in Biggins 2013). When chromosomes are attached improperly, a signal of no tension is sensed to activate the spindle assembly checkpoint (SAC). Components of this checkpoint include the Mad1, Mad2, Mad3, Bub1, Bub3 and Mps1 proteins (Hoyt, Totis and Roberts 1991; Li and Murray 1991). The SAC mediates cell-cycle arrest by inhibiting the anaphase-promoting complex (APC) via inhibition of Cdc20 (Li et al. 1997; Kim et al. 1998). This prevents the APC from promoting the ubiquitylation of cyclin B and securin, two key substrates required for mitotic progression (Peters 2006). The checkpoint signal that prevents cells with defective spindles from initiating chromosome segregation is generated at the kinetochore. Although all SAC proteins except Mad3 localize to the kinetochore (Gillett, Espelin and Sorger 2004), the time of binding and the roles of the specific proteins differ. Mad1 and Mad2 are specifically recruited to unattached kinetochores (Gillett, Espelin and Sorger 2004). Although Mad1 is stably bound to unattached kinetochores, Mad2 exists in two different pools. The Mad2 in a ‘closed’ conformation (Mad2-C) remains stably bound to Mad1 and serves as a receptor for soluble Mad2 in an ‘open’ conformation (Mad2-O) (Luo et al. 2002; De Antoni et al. 2005). Mad2-O cycles onto the kinetochore, where it binds to Mad2-C, promoting the conversion of Mad2-O to Mad2-C. In the ‘closed’ conformation, the Mad2 protein exposes the dimerization surface, which is also required downstream of kinetochores to mount a checkpoint response. Interestingly, downstream of kinetochores, the dimerization surface does not mediate Mad2 dimerization but interaction with Mad3 (Mariani et al. 2012). Because Mad1 and Cdc20 have similar Mad2-binding motifs, Mad2-C is able to bind to Cdc20 as well (Luo et al. 2002; Mapelli et al. 2007). Thus, the kinetochore Mad1-bound pool of Mad2-C promotes the formation of a soluble Mad3-Mad2-Cdc20 complex (stabilized by the Mad3–Mad2 interaction), subsequently promoting APC inhibition. The Bub1 and Bub3 checkpoint proteins always localize to kinetochores during mitosis, but similar to Mad1 and Mad2, they are also required for regulation of kinetochore biorientation (Warren et al. 2002; Lee and Spencer 2004). Due to this additional function, mutation of these genes results in the strongest segregation defects (Warren et al. 2002; Kawashima et al. 2010; Storchova et al. 2011). A mutation in the SGO1 gene encoding the spindle checkpoint component involved in sensing the presence of that tension leads to chromosome loss as well (Indjejian, Stern and Murray 2005; Fernius and Hardwick 2007). The iron-responsive transcription factor Aft1 associates with the kinetochore complex through Imi3 and, similar to Imi3, is required for the increased association of cohesin
with pericentric chromatin, which in turn is required to resist microtubule tension. Thus, oft1Δ cells display chromosome segregation defects (Hamza and Baetz 2012). Another protein involved in activation of the SAC in response to a defect in mitosis is histone H3. A histone H3 mutation impairs the ability of yeast cells to activate the checkpoint in a tensionless crisis, leading to missegregation and aneuploidy. This defect results from an attenuated H3–Sgo1 interaction that is essential for pericentric recruitment of Sgo1. Histone H3 seems to be a key factor in transmitting tension status to the SAC (Luo et al. 2010).

**Ploidy shifts in yeast**

Studies have shown that ploidy reduction towards diploidy occurs after several hundred generations in both triploid and tetraploid lines of *S. cerevisiae* (Gerstein et al. 2006; Gerstein, McBride, Otto 2008). However, after a sufficient number of generations (approximately 1800 generations), haploid strains also convert into diploid strains (Gerstein et al. 2006). These results suggest that the diploid state is favorable for standard laboratory *S. cerevisiae* strains maintained under typical conditions. However, ploidy change occurs quite often in diploid cells.

Similar to aneuploidy, ploidy changes in yeast cells can be provoked by several mechanisms: (1) defects in condensation of not fully replicated or unrepaird damaged DNA that persist till mitosis; (2) defects in interaction between DNA, kinetochore and spindle microtubules; (3) perturbations in mitotic spindle organization, positioning, orientation or microtubule dynamics; (4) disturbances in septum organization; (5) malfunction of control pathways of chromosome division and (6) various exogenous factors, such as gamma irradiation, certain anticancer drugs or heat shock, that can affect the aforementioned factors. Two experimental approaches revealed this intriguing phenomenon, which is exclusive to diploid cells. Saccharomyces cerevisiae diploid cells constantly exposed to rearrangement stress as a consequence of deletion of RAD52 (engaged in HR) or CTF18 (regulating PCNA and cohesion complex access to DNA) led to GCR, which can reduce the genome to the haploid level. This process occurs sequentially in rad52/rad52 cells, resulting in the loss of the chromosomes one by one during successive divisions (Song and Peters 2012), whereas ctf18/ctf18 cells reduce their genome in a single step (Alabrudzinska, Skoneczny and Skoneczna 2011). Both haploidization types result in the limitation of rearrangement events. Due to the shortage of homologous sequences in the haploid genome, GCR frequency is low in haploid compared to diploid cells (10⁻⁹ versus 10⁻⁶, respectively) (Chen and Kolodner 1999; Hiraoka et al. 2000; Ohnishi et al. 2004). It is still not clear if the tendency to reduce the ploidy level observed in diploid strains bearing rad52 or ctf18 mutations is incidental or a way to avoid further genome destabilization. However, haploid strains that appeared in the process of ploidy reduction have much lower mutation rates than initial diploids; moreover, they are fertile and more vital, so they likely dominate the cell population and have better chances for survival (Alabrudzinska, Skoneczny and Skoneczna 2011). Although it should be noticed that the frequency of haploidization is not high, approximately 90% of cells in the initial diploid population drift in the opposite direction and increase their DNA content to the aneuploid or polyploid level. These cells show elevated mutation rates, decreased fitness and escalating division problems, leading to permanent cell-cycle arrest and finally to cell death (Alabrudzinska, Skoneczny and Skoneczna 2011).

Ploidy changes have also been reported for other yeast mutants, but those cases involved the conversion of haploids into diploids. These changes accompany deletions of a number of genes. Haploid cells devoid of the ZDS1 gene (which encodes a factor involved in mitotic exit) will become diploid cells when exposed to Ca²⁺ ions (Miyakawa and Mizunuma 2007). Mutations in any of three adjacent residues (L97, Y98 or G99) near the C-terminus of histone H4 led to polyploidy as well (Yu et al. 2011). The deletion of the genes encoding components of the RSC chromatin remodeling complex required for GI/S transition (SFH1 or RSC3) resulted in a ploidy shift that could be rescued by CLB5 deletion (S-phase cyclin) or by transient depletion of the replication origin licensing factor Cdc6 (Campsteijn, Wijnands-Collin and Logie 2007). Increased ploidy is also observed in 1pl1Δ and bem2Δ strains (Chan and Botstein 1993). 1pl1 is an Aurora kina- nase in the chromosomal passenger complex required for spindle pool body cohesion, regulation of chromosome segregation, spindle checkpoint and cytokinesis. The Rho GTPase-activating protein Bem2 is involved in the control of cytoskeleton organization and is required for bud emergence (Chan and Botstein 1993). These data highlight the role of regulatory systems in the control of cellular ploidy.

Schizosaccharomyces pombe is a valuable model for studying the mechanisms governing the accuracy of chromosome segregation. Because fission yeast cells divide equally, much like the cells of higher eukaryotes, these organisms are often used as models for studying chromosome segregation. However, both *S. cerevisiae* and *S. pombe* undergo closed mitosis, whereas the majority of higher eukaryotes perform open mitosis, during which the nuclear envelope is disassembled. The release of the nuclear content enables spindle microtubules to access kinetochores (Kanoh 2013). Nuclear envelope breakdown does not occur during closed mitosis. This feature of yeast cells changes the nature of the chromosome segregation process. In *S. pombe*, telomeres are tethered to the nuclear envelope in interphase. Chromosome tethering is required for various nuclear processes, such as transcription, transport and DNA repair. But chromosome segregation in mitosis requires efficient chromo- some movements to assure equal division. Cdc2-dependent phosphorylation of the telomere-binding protein Rap1 during mitosis promotes the transient dissociation of telomeres from the nuclear envelope, which is required for accurate chromosome segregation (Fujita et al. 2012). The tethering of chromo- somes ends has an additional function: it protects telomeres from illegitimate end-to-end fusions that would otherwise lead to dicentric chromosome formation. Dicentric chromosomes trigger breakage–fusion–bridge cycles and subsequent genome instability (Almeida and Godinho Ferreira 2013). The creation of chromosome fusions in telomerase mutant strains depends on the MMEJ SSA repair pathway and requires MRN/Ctp1. In strains lacking the telomere regulator Taz1/TRF2, end-joining reactions occur via NHEJ (Almeida and Godinho Ferreira 2013).

Studies have shown that the putative Ras1 effector in *S. pombe*, Scd1, alters microtubule dynamics within the spindle to affect spindle assembly and chromosome capture. Scd1 physically associates with Moe1, a factor that contributes to the inherent instability of microtubules and is required for proper spindle function. Defects in both Ras1 and Scd1 lead to genome instability (Segal and Clarke 2001). Recent studies have impli- cated two other pathways that contribute to the regulation of accurate chromosome segregation: Chk1/Wat1-dependent regulation of the mitotic spindle microtubule structure (Verma et al. 2014) and histone variant H2A.Z-dependent regulation of cohe- sin dynamics. Indeed, it has been shown that H2A.Z and the SMC complex ensure genome integrity through accurate chro- mosome segregation (Tapia-Alveal et al. 2014).
Mechanisms of ploidy change in yeast

Ploidy shifts occur frequently in yeast cells. One can assume that most of these events happen when cells evade death initiated by improper division. Cells that are arrested at the mitotic checkpoint are expected to die if they unable to restore proper mitosis. Ongoing aberrant division must result in abnormal DNA content. Division arrest may be initiated by severe DNA damage, spindle perturbation or even heat shock and should not be revoked until the problem is solved. If the signal recognizing the presence of anomalies does not disappear and division arrest persists, the cell normally is marked for removal by mitotic catastrophe similar to the process in mammalian cells (Vitale et al. 2011). However, when cell-cycle control fails or there is a mutation predisposing the cell to genomic instability, the cell may avoid execution and the aberrant division may be completed with various consequences, such as the gain or loss of single chromosomes, haploidization or entering into sequential steps of illegitimate polyploidization and depolyploidization (Fig. 5).

What mechanisms could lead to a ploidy shift? In general, it is believed that genome duplication may occur by two independent mechanisms: ‘mitotic slippage’ or ‘cytokinesis failure’. Aneuyploidization can be provoked by DNA discontinuities, by errors in chromosome attachment (e.g. monotelic, syntenic or merotelic chromosomes) (Fig. 6), or by non-synchronous and delayed spindle movement.

However, in other fungi, the ploidy shift also results from the successive loss of chromosomes through different stages of aneuploidy, ending with haploid state. This mechanism has been described for different species of imperfect fungi after treatment with benomyl (Stahl and Isser 1992; Samsonova et al. 1996) and γ-radiation (Mortimer, Contopoulou and Schild 1981), and after protoplasts fusion (Fournier et al. 1997; Büttner et al. 1990) or in early zygote progeny (Kurischko 1986). The polyploid and subsequent ploidy reduction via multipolar spindle formation has been observed in the tetraploid cells of C. albicans (Suzuki et al. 1986). Indeed, other unexpected ploidy changes have also been described in the ‘obligate diploid’ C. albicans. This fungus can become haploid through a chromosome loss mechanism, and the haploids of different mating types efficiently mate to regenerate the diploid form. Homozygous diploids arise spontaneously through autodiploidization. Haploids and autodiploids can become haploid through a chromosome loss mechanism, however, it cannot be excluded at this point. A similar mechanism comprising micronuclei formation, heterochromatinization and fragmentation of micronucleated chromatin in the final step during haploidization in a process resembling apoptosis (named programed DNA elimination) was observed in other eukaryotic cells (Zelesco and Graves 1983; Gernand et al. 2005). In mammalian cells, the formation of micronuclei containing aggregated, DSB-fragmented DNA, kinetochores and centromeres is widely used to score genomic instability, genotoxic exposure or replication stress (Norppa and Falck 2003; Huang et al. 2011; Xu et al. 2011). Determining whether nucleoautophagy that occurs in yeast cells (Krick et al. 2008; Robert et al. 2011) may work as a similar mechanism requires further study. Gerstein et al. provided additional evidence supporting the first haploidization mechanism described above. In their experiments, polyploid S. cerevisiae cells were cultured for hundreds of generations, resulting in ploidy reduction towards diploidy. Remarkably, the chromosome loss was not random. Instead, nearly complete sets of chromosomes were lost at once, with some additional chromosome missegregation events. The authors proposed a mitotic mechanism for the elimination of an entire set of chromosomes in S. cerevisiae, which would reduce the ploidy level (Gerstein et al. 2006; Gerstein, McBride, Otto 2008).

TRANSCRIPTION CAN AFFECT GENOME STABILITY

Although most mutagenic changes are introduced into DNA during its replication or due to erroneous DNA repair, the importance of transcription as a cause of genetic instability is increasingly being recognized. It is hardly surprising that transcription can affect genome stability because RNA synthesis involves significant temporary changes in DNA strand conformation and, especially when passing through highly expressed genes, transcription machinery may often come into conflict with other protein complexes operating on the DNA, primarily the replication complex. Remarkably, approximately 40 human genes are between 1 and 2.3 million base pairs in length (Scherer 2008). Because the rate of transcription in eukaryotes is 18–42 nt per second (Pérez-Ortin, Alepuz and Moreno 2007), in such extreme cases the gene transcription can last for more than 10 h and span several cell division cycles (Tennyson, Klamut and Worton 1995). Thus, encounters between the transcription and replication machineries are inevitable. For this reason, those loci experience frequent mutagenic events (Helmrich, Ballarino and Tora 2011).

Intensively transcribed DNA stretches are also prone to mutagenesis inflicted by transcription in S. cerevisiae cells.

The first indications that transcription could destabilize the genome came from early studies on bacteria demonstrating with the reversion assay that increased transcription from a certain region was accompanied by elevated mutagenesis of that region (Fliermans and Dworkin 1971). Later, this phenomenon was found and studied in both prokaryotic and eukaryotic models with a substantial contribution from S. cerevisiae. Its densely packed, mostly intronless genome makes studies of this phenomenon especially convenient. As mentioned above, the RNA polymerase complex (RNAP) may collide with the replication fork. This is most likely to occur when the complexes move in opposite directions (head-on collisions); however, in bacteria in which replication is more than 10 times faster than transcription (approximately 600 nt per second), collisions can also occur when they move codirectionally. Transcription involves the separation of DNA strands; thus,
Figure 5. The DNA content disturbances mechanisms in yeast. When cells arrested at the mitotic checkpoint are unable to restore proper mitosis, they supposedly die due to mitotic catastrophe. If cells evade death at this point, the ploidy shifts may occur. Aberrant division results in various consequences, such as the gain or loss of single chromosomes, haploidization or entering into sequential steps of illegitimate polyploidization and depolyploidization. Genome duplication may occur by 'mitotic slippage' or by 'cytokinesis failure' leading to binucleated or polyploidial cell, respectively. Polyploidal cells frequently undergo aneuploidization. Ploidy reduction may occur by multipolar spindle formation and subsequent loss of chromosome set in one step, by successive loss of chromosomes leading to haploid state, or by selective elimination of the parental chromosomes from the cell via nucleoautophagy.
the non-transcribed strand (NTS) is exposed to potential attack by mutagenic agents. During transcription, newly synthesized RNA may form a heteroduplex with the transcribed strand (TS), whereas the NTS remains single stranded and may form alternative secondary structures. Moreover, the movement of RNAP along the DNA strand distorts its double helix uniformity. All of these transcription-associated incidents make the transcribed DNA regions prone to mutagenic changes introduced through various mechanisms that have been described in previous sections of this review. These mechanisms are collectively classified as transcription-associated genome instability (TAGIN) and can be divided into two broad categories: transcription-associated mutagenesis (TAM) and transcription-associated recombination (TAR).

Sources and mechanisms of TAGIN

During transcription, DNA strands are separated by the RNAP complex, creating the transcription bubble. This enforces positive supercoiling in front of the complex and negative supercoiling behind it (Liu and Wang 1987). Although this torsional stress is constantly relaxed by topoisomerases (Wang 2002), the journey of RNAP along the transcribed genome region is accompanied by local distortion of the native DNA helix. Nascent RNA molecule forms hybrid 9 nucleotides in length with the strands within the double helix (Roberts and Crothers 1992), thus, this phenomenon might represent yet another transcription-related source of torsional stress, resulting in negative supercoil accumulation with all its associated consequences (Fig. 7A).

One of the structures that may form in the trail of the RNAP moving along the DNA is an R-loop (Fig. 7B). This structure consists of a stretch of ssDNA of the NTS and a hybrid between the newly synthesized RNA and TS. The formation of an R-loop is favored by the negative DNA supercoils that facilitate invasion of the DNA helix by the RNA strand. RNA:DNA heteroduplexes are thermodynamically more stable than the respective DNA strands within the double helix (Roberts and Crothers 1992), especially when the NTS strand is G-rich (Ginno et al. 2013); thus, one might wonder how the cell is able to dissociate them once they form. Yet under normal conditions, there are various processes that are synchronized with transcription that can counteract R-loop formation. In prokaryotes, transcription is coupled to translation (Gowrishankar and Harinarayanan 2004), which is able to pull RNA off the transcribed operon. In eukaryotes, RNA is post-transcriptionally processed and exported from the nucleus (Li and Manley 2006). Incidentally, although R-loop formation is considered to be a threat to genome stability (see below), it has also been shown to play a positive regulatory role in transcription in human cells (Ginno et al. 2013).

The consequences of collisions between transcription and replication may be even more serious. Distinct from bacteria, in eukaryotes, transcription and replication are usually temporally separated within the cell cycle, although some genes, including those encoding rRNA, tRNA and histones, are transcribed during S phase. Replication and transcription seem to be also spatially separated to some extent (Wei et al. 1998). Nevertheless, collisions are unavoidable. Intuitively, head-on conflicts would seem to inflict more damage than codirectional ones. Transcription and replication travelling in the opposite direction force the buildup of positive supercoils between them, which may impede replication fork movement. Indeed, replication fork pauses...
have been observed in yeast at sites of head-on encounters between replication and transcription of rRNA genes (Deshpande and Newlon 1996). These collisions can lead to replication fork arrest or reversal, resulting in formation of potentially harmful structure called ‘chicken-foot’, as was shown for Escherichia coli (Postow et al. 2001). This type of structure has been demonstrated in S. cerevisiae when the replication block leads to activation of the PRR damage avoidance pathway via fork regression (Blastyák et al. 2007).

Transcription-associated genome instability

Transcription as a source of point mutations and small insertions or deletions

The first suggestions of the occurrence of TAM in S. cerevisiae appeared in the early 1990s. It was shown that the induction of gene expression could increase the mutation rate of that gene by as much as 150-fold (Korogodin et al. 1991). This finding was later supported by results obtained with pGAL- and pTET-inducible promoter systems by Datta and Jinks-Robertson (1995). More recently, it was shown that the mutagenesis rate is directly proportional to the level of transcription (Kim et al. 2007). By cloning the reversion system in either orientation, the authors demonstrated that the overall mutation rate was similarly increased regardless of the direction of replication fork movement, but an influence on the mutation spectra was observed. An impact of the transcription level on the mutation spectra was also demonstrated in another study. While low transcription levels resulted mostly in base substitutions, high transcription levels predominately led to short insertion–deletion mutations (Lippert et al. 2004), with two nucleotide deletions comprising 21% of all mutations. Therefore, the authors deemed 2-nt deletions as a TAM signature.

The TAM phenomenon was mostly studied on individual loci, sometimes artificially generated that may be prone to increased instability due to the heterologous context or might in some other way be sequence specific. Therefore, one would wonder if this phenomenon manifests across the whole genome. This question was addressed in a study analyzing the whole genomic sequence and transcriptome data (Park, Qian and Zhang 2012). The authors compared genome sequences of closely related S. cerevisiae and S. paradoxus as well as human and macaque sequences selecting only differences present in transcribed yet selectively neutral intron sequences. Then the authors compared the frequency of mutational changes within those sequences with their transcription levels and were able to demonstrate simultaneously for numerous loci that positive correlation between transcription and mutagenesis does exist.

A characteristic feature of TAM is its asymmetry. It was demonstrated in E. coli by transcription-dependent preferential deamination of cytosines on the NTS (Beletskii and Bhagwat 1996). These data were more recently substantiated by a demonstration of strand asymmetry in the incidence of spontaneous oxidative and alkylationing lesions (Klapacz and Bhagwat 2005; Fix, Canugovi and Bhagwat 2008). In a whole genomic comparison study of two related yeast species, all types of mutations were found in correlation with increased transcription. However, the most frequent base substitutions were C→T, A→G, G→T and A→T (Park, Qian and Zhang 2012), suggesting that TAM is a result of cytosine or adenine deamination and the consequence of oxidative damage because 8-oxo-guanine (the major oxidative lesion) can mispair with adenine (Cheng et al. 1992). Another type of lesion that seems to be associated with a high level of transcription is the generation of AP sites (Morey, Greene and Jinks-Robertson 2000). Such sites are repaired by preferential incorporation of dUTP, leading to mutation and allowing detection of the lesions (Kim and Jinks-Robertson 2009). However, the mechanisms of transcription-associated base removal from the DNA strand are currently unknown. Thus, it seems clear that the predominant source of TAM is DNA damage.

Transcription-associated recombination

Transcription can also invoke higher order changes in the genome. It was demonstrated in S. cerevisiae that TAM may be the consequence of the activity of all three RNAPs (Keil and Roeder 1984; Pratt-Hyatt et al. 2006). Conflicts between transcription and replication are considered the main source of TAM. Head-on conflicts stimulate TAR more than codirectional conflicts (Prado and Aguilera 2005) (Fig. 7); however, a genome-wide study revealed correlations of both types of conflict with high levels of transcription in yeast (Azvolinsky et al. 2009). These conflicts can lead to replication fork arrest (Bermejo, Lai and Foiani 2012), thereby increasing the probability of replication fork collapse and causing recombinogenic DSBs and ssDNA gaps (Aguilera and Gómez-González 2008; Branzei and Foiani 2010). Likewise, recombination may be mediated by R-loops (González-Aguilera et al. 2008; Stirling 2012), since in yeast, deficiency of the THO complex that binds to the nascent mRNA during transcription elongation (Rondón, Jimeno and Aguilera 2010) shows a strong hyperrecombination phenotype (Piruat and Aguilera 1998). However, the exact mechanism(s) of recombination originating from R-loop formation or transcription–replication conflicts is unknown.

One of the regions of potential intense conflicts between replication and transcription is the rDNA region of the genome. In S. cerevisiae, this region consists of more than 100 identical tandem repeats located on chromosome XII (James et al. 2009). Each repeat encodes 35S rRNA and on the opposite strand, SS rRNA. The direction of transcription of both of rRNAs is outward from the ARS element that lies between both sequences. This configuration prevents head-on collision between replication and transcription within a single rDNA unit but would not prevent conflicts with transcription running from the adjacent unit. However, each repeat is separated by a short sequence called the replication fork barrier. The binding of Fob1 protein to the barrier blocks the progression of the replication fork in one direction, into 35S rDNA sequence of the neighboring rDNA unit (Kobayashi 2003; Mohanty and Bastia 2004).

Multiple copies of rDNA make this region exceptionally susceptible to lesions that can instigate recombination and detrimental diversification of the number of repeats within the cell population. Nevertheless, maintaining the optimal number of rDNA copies also depends on recombination (Kobayashi 2011). A study demonstrated that the transcription of rDNA is necessary to accomplish this maintenance (Kobayashi et al. 1998). In this case, tightly controlled TAR is beneficial to the S. cerevisiae cell.

TAM as an adaptive mechanism

Contrary to mutations introduced during replication, those imposed by transcription can occur in non-dividing cells under starvation conditions. Notably, starvation-induced expression of specific genes leads to increased mutagenesis within their sequences and provides potential benefits to cells under strong selection pressure (reviewed in Wright 2004). In fact, the first paper reporting data indicating the existence of TAM in budding yeast (Korogodin et al. 1991), mentioned on the beginning of this
section, describes exactly this type of experimental setup. It is likely that TAM is common in starved or resting cells because in a genome-wide study of stationary-phase cells, a correlation was found between the level of transcription and DNA turnover (probably reflecting the repair reactions) at various loci (de Morgan et al. 2010).

Transcription-coupled DNA repair

In addition to mutagenesis, one of the mechanisms of DNA repair is coupled to RNA synthesis, which increases the complexity of connections between transcription and genome stability even further. Various DNA lesions, including UV-induced photoproducts and bulky adducts present on the TS, create a physical barrier to movement of the RNAP along the DNA strand. The stalled complex recruits nucleotide–excision repair (NER) machinery (Hanawalt and Spivak 2008). Recently, it was demonstrated in yeast that AP sites on the TS are repaired by the NER pathway, indicating that non-bulky lesions also trigger TCR (Kim and Jinks-Robertson 2010). Therefore, transcription-coupled DNA repair (sometimes called TC-NER) preferentially repairs the transcribed DNA strand. In yeast, the Rad26 protein is required for efficient TC-NER (van Gool et al. 1994).

MITOCHONDRIA AND GENOME STABILITY

Mitochondria are double-membrane-bound organelles that are found in all eukaryotic cells. These organelles are thought to have evolved through endosymbiosis of α-proteobacteria-like cells that colonized a primordial eukaryotic cell for mutual benefit (Gray 1999). Mitochondria play an important role in the transformation of the energy stored in the bonds of carbohydrate molecules into the ATP synthesis via oxidative phosphorylation during respiration. Oxidative phosphorylation is accomplished in the inner mitochondrial membrane through the sophisticated assembly of four respiratory multisubunit protein complexes (although both budding and fission yeast lack complex I) of the electron transport chain and the fifth respiratory complex of ATP synthase (Saraste 1999). During evolution, mitochondria lost most of the ancestral proteobacterial genome, reflecting gene transfer to the nucleus and the loss of redundant information. However, these organelles preserved a residual genome encoding a few subunits of the respiratory complex, together with ribosomal RNAs and tRNAs necessary for mitochondrial genome expression. Budding and fission yeast share some features of their mitochondrial biology (see Table 2), but, as with many other aspects of Sc. pombe cell biology (Rhind et al. 2011), fission yeast mitochondria are more similar to mammalian mitochondria (Schäfer 2003). Specifically, the mitochondrial genome of Sc. pombe cells is compact compared to S. cerevisiae mtDNA (19 kb versus 75 kb), with few transcriptional initiation sites, short intergenic regions and the tRNA punctuation mode for mitochondrial messenger maturation, as in higher eukaryotic cells. Surprisingly, despite differences in genome organization, the sets of mtDNA genes in these two yeast species are remarkably convergent in spite of 500 million years of evolution after the two lineages diverged (Dashko et al. 2014). The most significant difference between the two yeast species is dependence on wild-type mtDNA, and consequently, dependence on respiration. Budding yeast cells are petite positive, i.e. able to survive without the wild-type mtDNA (so called rho+ mitochondrial genome) and the ability to respire. These mutants form smaller colonies than respiring cells on a glucose-limiting medium. In contrast, fission yeast cells are petite negative (Bulder 1964). The petite-positivity of budding yeast cells has greatly facilitated studies on the mechanisms responsible for the maintenance of mtDNA in this species. Cytoplasmic petite mutations occur spontaneously in budding yeast at a frequency of 1% per generation (Ephrussi 1953). There are several varieties of cytoplasmic petites (reviewed in Dujon 1981; Contamine and Picard 2000). (1) The most frequent varieties are those that possess inactive mtDNA products of mitochondrial genome rearrangements, including partial deletions and amplifications of remaining fragments (rho− mutants). (2) The mitochondria of another class of petites are entirely devoid of mtDNA (rho0 mutants). These yeasts are thought to arise spontaneously due to a failure to correctly segregate mtDNA with the mitochondria during their transmission to the incipient bud. (3) Further, there are cytoplasmic petites that have lost the ability to respire due to

**Table 2. Comparison of budding and fission yeast cells in respect to their energy-yielding metabolisms and mitochondrial genomes in reference to the human mtDNA (based on Kühl 2010 and Flores et al. 2000).**

| Feature                                      | Saccharomyces cerevisiae | Schizosaccharomyces pombe |
|----------------------------------------------|--------------------------|---------------------------|
| Metabolism                                   |                          |                           |
| Ability to ferment glucose and produce ethanol. | Yes                      | Yes                       |
| Ability to use ethanol as a sole carbon source. | Yes                      | No                        |
| Mitochondrial genome                         |                          |                           |
| Petite-positivity                            | Yes                      | No                        |
| Introns in mitochondrial genesa              | Yes                      | No                        |
| tRNA coded in mtDNA                          | 15S, 21S                 | rrs, rnl                  |
| RNA component of RNase P coded in mtDNAb     | 9S                       | rnp                       |
| Ribosomal subunit genec                      | VAR1                     | rps3                      |
| Major ORFs                                   | 8                        | 8                         |
| Complex I                                    | 0                        | 0                         |
| Complex III                                  | COB (CYTb)               | cytb                      |
| Complex IV                                   | COX1, -2, -3             | cox1, -2, -3              |
| Complex Vd                                   | ATP6, -8, -9             | atp6, -8, -9              |

aIn human mtDNA, there are no introns interrupting mitochondrial genes.
bThere is no RNase P RNA coded in human mtDNA.
cThere is no gene for a ribosomal protein in human mtDNA.
dIn human mtDNA, there are seven genes coding for subunits of complex I.
eIn human cells, subunit 9 of complex V is encoded by a nuclear gene.
a point mutation or mutations in only one or two of the mitochondrial genes necessary for respiration (mit− mutants). In addition to respiratory deficiencies, the rho− and rho0 mutants display pleiotropic phenotypes associated with their inability to synthesize mitochondrial proteins. In contrast, mit− mutants are proficient in mitochondrial gene expression, although these yeasts are also unable to respire. It is not easy to distinguish between rho−/rho0 and mit− mutants upon first sight; therefore, to estimate point mutagenesis in yeast mtDNA, researchers test yeast cultures to determine the frequency of mutants that are respiratory proficient, but resistant to inhibitors that target specific mitochondrial processes. Often, the resistance phenotypes are associated with point mutations in mtDNA. Several mitochondrial inhibitors have been used in numerous studies for the isolation of mitochondrial point mutations, but erythromycin is the most specific for S. cerevisiae mtDNA (Baruffini, Ferrero and Foury 2010). Resistance to erythromycin is usually conferred through mutations at several positions in the mitochondrial 21S rRNA gene (Sor and Fukuhara 1982; Vanderstraeten et al. 1998). Notably, mitochondrial point mutations (whether mit− or mitochondrial drug resistance mutations) occur rarely compared with rho− deletions (see above). The frequency of spontaneous and induced mitochondrial point mutations is several orders of magnitude lower than the incidence of petite mutants (Foury and Vanderstraeten 1992; O’Rourke et al. 2002). This difference suggests that point mutations and rho− deletions in mtDNA result from distinct mechanisms. In contrast to S. cerevisiae, fission yeast cannot survive the loss or even a large deletion of mtDNA. The metabolism of these yeasts strictly depends on respiration, even when grown on glucose (Flores et al. 2000). Nevertheless, several genetic modifications were shown to convert fission yeast into petite-positive organisms: (1) nuclear mutations (ptp1−1 or ptp2−1 (Hafferl and Fox 1992), until recently unidentified, (2) expression of S. cerevisiae YME1 gene, encoding a mitochondrial protease associated with the inner membrane (Kominsky and Thorsness 2000) and (3) an mtDNA mutator allele of the mitochondrial gene rps3/urf A, encoding a putative mitoribosomal protein (Seitz-Mayr and Wolf 1982; reviewed in Schäfer 2003). Mutations suppressing the mtDNA-loss lethality of another petite-negative yeast species Kluyveromyces lactis have been found to be alleles of genes encoding subunits of the mitochondrial F1-ATPase (reviewed in Clark-Walker 2003). These mutations increase the affinity of F1-ATPase for ATP in the hydrolysis reaction (Clark-Walker 2003). Many lines of evidence indicate that the robust ATP hydrolysis is critical for the viability of cells lacking the functional mtDNA, because the mitochondrial inner membrane potential, Δψ, in petite cells is maintained, in lieu of the respiratory chain activity, by hydrolysis of ATP imported from cytosol (Giraud and Velours 1997; Clark-Walker 2003). Consequently, in petite cells, ATP is imported from cytosol into mitochondria by the reversal of the ATP/ADP translocator that generates in the process an electric potential sufficient to maintain the mitochondrial protein import. The mitochondrial protein import, in turn, is essential for cell viability (reviewed in Chacinska et al. 2009), because the mitochondria are essential organelles even in cells of petite-positive species that do not require respiration (the essential function of mitochondria is described in the section ‘Mitochondrial influence on the nuclear genome’). Thus, it is likely that similar mutations to those from K. lactis conferring petite-positivity can be isolated in the fission yeast. However, although the feature of petite-positivity in S. cerevisiae has significantly contributed to the progress of studies on mtDNA stability, the potential advantage of using petite-positive strains of the fission yeast for studying the mechanisms of mtDNA maintenance remains unknown. Thus, most data on the stability of mtDNA in yeast come from studies on budding yeast, and, therefore, these results are reviewed below, with occasional forays into the S. pombe mitochondria if such data are available.

Nucleoids

Mitochondrial DNA in eukaryotic cells is packaged into compact nucleoprotein structures called nucleoids (Chen and Butow 2005; Kucej and Butow 2007). Nucleoids form dynamic complexes, and the number of nucleoids, mtDNA molecules/nucleoids and nucleoid protein content varies depending on growth conditions. An aerobically grown budding yeast cell might contain 40–60 nucleoids, whereas under conditions of limited aeration, the number of nucleoids is less than 10 per cell. The number of mtDNA molecules per nucleoid is inversely related to the number of nucleoids: the more nucleoids per cell, the less mtDNA molecules per nucleoid. The protein components of mitochondrial nucleoids have been identified in various organisms and are vastly divergent for organisms from different lineages (Kucej and Butow 2007). Surprisingly, in addition to the obvious proteins implicated in DNA transactions (e.g. replication and transcription) and DNA packaging, and in addition to the chaperonins and proteases involved in regulating other proteins, nucleoid components include coopted metabolic enzymes that, in a few instances, have been shown to be bifunctional (see below). This group of nucleoid proteins is the most lineage specific (Kucej and Butow 2007). Nevertheless, the main packaging proteins in nucleoids, HMG (high-mobility group) proteins, are conserved in eukaryotic cells. Abf2 is the HMG protein in S. cerevisiae mitochondria, whereas the mitochondrial transcription factor A (TFAM) is the HMG protein in human mitochondria.

In contrast to TFAM, the S. cerevisiae nucleoid protein does not play a specific role in regulating transcription in yeast. Instead, Abf2 binds mtDNA with some preference for GC-rich sequences (GC clusters in AT-rich sequences are dominant features of S. cerevisiae mtDNA) and determines the compaction and structure of mitochondrial nucleoids (Newman et al. 1996). Cells lacking the ABF2 gene lose rho+ mtDNA when grown on glucose-rich medium (Diffley and Stillman 1991), but these cell maintain rho− genomes on media with either non-fermentable or non-repressing substrates as sole carbon sources (Chen et al. 2005), or under conditions of general amino acid control pathway induction in cells grown without the amino acids isoleucine, leucine and valine (Zelenaya-Troitskaya, Perlman and Butow 1995). In the former case, it was shown that another mitochondrial nucleoid protein, Aco1, whose expression is repressed in glucose-rich media, partially replaces Abf2 in the nucleoid. In the latter case, another component of the mitochondrial nucleoid replaces the Abf2 mtDNA-packaging function, the Ivv5 protein. Ivv5 is the acetohydroxy acid reductoisomerase that catalyzes one of the reactions in the branched chain amino acid biosynthesis pathway. The Aco1 protein is a bifunctional iron–sulfur cluster (ISC)-containing enzyme, which functions as both a mitochondrial aconitase in the tricarboxylic cycle and a nucleoid protein for the maintenance of rho− genomes (Chen et al. 2005). These two functions are accomplished by two separate domains of the protein. Ivv5 binds to dsDNA independently of Abf2, and a deficiency of this protein destabilizes rho− mtDNA (Macierzanka et al. 2008). However, when Aco1 replaces the Abf2 mtDNA-packaging function in nucleoids, mtDNA is less protected and becomes sensitive to nuclease attack (Newman et al. 1996) and oxidative...
damage (O’Rourke et al. 2002). It has also been shown that, under these conditions, microsatellite repeats and longer direct repeats in mtDNA are destabilized, although point mutations are detected at wild-type levels (Sia et al. 2008). These results indicate that the lack of Abf2 in mitochondrial nucleoids increases polymerase slippage events during mtDNA replication, and this defect might lead to mitochondrial genome rearrangements. Therefore, this mechanism most likely reflects the rho⁺ instability observed in cells lacking Abf2. However, there is also evidence that Abf2 is a positive factor in some HR pathways. In a study using 2D gel electrophoresis, Abf2 promoted HJ intermediates in rho⁻ mtDNA in vivo but not recombination intermediates detected in the mtDNA of rho⁻ petite strains (MacAlpine, Perlman and Butow 1998). Thus, Abf2 not only helps to prevent genome rearrangements during replication, but it may also be a positive factor of a mitochondrial recombination process that helps to stabilize rho⁺ genomes. Interestingly, the report by Taylor et al. (2005) has shown that moderate overexpression of ABF2 increases the mtDNA copy number. Abf2 overproduction also increases the level of recombination intermediates (MacAlpine, Perlman and Butow 1998), suggesting that elevated Abf2-dependent recombination stimulates mtDNA replication.

Unfortunately, mitochondrial nucleoids from fission yeast cells have not yet been studied. Moreover, there are many predicted HMG proteins in the thermotrich genome (PomBase, http://www.pombase.org/), but none of these proteins have been identified as an obvious ortholog of Abf2 or TFAM, and none have been found to have a clear cut mitochondrial targeting sequence (based on MitoProt prediction; MitoProt, http://ihg.gsf.de/ihg/mitoprot.html/, last date accessed 12 November 2014). Therefore, the protein content in the mitochondrial nucleoids of fission yeast requires further study, which will likely reveal more instances of ‘evolutionary tinkering with nucleoids’ (Kucej and Butow 2007).

Replication of mtDNA

Saccharomyces cerevisiae cells harbor the mitochondrial DNA polymerase, Mip1 (Pol γ), which mediates mtDNA replication. The MIP1 gene deletion leads to the loss of mtDNA (rho⁺ strains; Foury 1989; Merz and Westermann 2009), indicating that this polymerase is a major, if not unique (see below), replicative polymerase in the mitochondria of budding yeasts. Similar results were obtained for the Saccharomyces pombe pogo1 gene, encoding the fission yeast ortholog of MIP1 (Chu et al. 2007). Mip1 functions as a single catalytic subunit enzyme (Lucas et al. 2004; Viikov, Väljamäe and Sedman 2011), in contrast to the mitochondrial polymerase from mammalian cells, which requires an accessory subunit (reviewed in Kaguni 2004). The mitochondrial DNA polymerase from the fission yeast, Pog1, is likely also a single subunit polymerase. The catalytic subunits of polymerase γ are phylogenetically related to the prokaryotic PolA family and well conserved in eukaryotic cells (Kaguni 2004), suggesting that Pol γ was an early invention in the evolution of mitochondrial endosymbiosis. In vitro studies have shown that Mip1 is a highly processive DNA polymerase with strong strand-displacement activity (Viikov, Väljamäe and Sedman 2011). Strand-displacement activity is important for the long-patch base-excision repair (BER) pathway, as described below, underscoring that Pol γ is responsible not only for mtDNA duplication, but also plays a role in various mitochondrial DNA repair pathways that require polymerase activity. Pol γ has three main activities: a polymerase activity, a 3′→5′ exonuclease proofreading activity and a 5′-dRP lyase activity (reviewed in Kaguni 2004). The high fidelity of mtDNA replication (Table 1) is largely determined by the N-terminal domain, which exhibits 3′→5′ exonuclease activity for the excision of misincorporated bases before polymerase extension. A mutation in this domain results in the substitution of a conserved residue, mip1-D347A, leading to a several fold increase in the frequency of point mutations in mtDNA and a several fold increase in the incidence of rho⁻ petite mutants (Foury and Vanderstraeten 1992). Moreover, defects in the Mip1 exonuclease activity combined with a defective mutation in the MSH1 gene, encoding the mitochondrial mutS homolog required in MMR in yeast mitochondria (see below), resulted in a synergistic mitochondrial error catastrophe and the complete loss of rho⁺ mtDNA (and conversion to rho⁻) after a few generations (Vanderstraeten et al. 1998). Thus, the exonuclease proofreading activity of Mip1 and the Msh1-dependent MMR pathway are the main, at least partially independent, guardians of genome stability during mtDNA replication in S. cerevisiae. The 5′-dRP lyase activity of the Mip1 polymerase is important for the elimination of intermediates during BER, which is reviewed below.

Interestingly, despite the fact that Mip1 is a major replicative DNA polymerase in S. cerevisiae mitochondria, it is not the only DNA polymerase in this compartment. Two subunits of Pol α, including the largest catalytic subunit, Pol1, and the accessory subunit Pol12, have been detected in yeast mitochondria (Lasserre et al. 2013). It is not known whether the mitochondrial subunits of Pol α are actually involved in the replication of mtDNA. However, several lines of evidence clearly indicate that the TLS polymerases, DNA Pol ζ (Rev3/Rev7) and Rev1 function in yeast mitochondria in pathways that are not equivalent to the nuclear functions of the polymerases. First, Zhang, Chatterjee and Singh (2005) showed that all three polymerases localize to yeast mitochondria, and inactivations of REV3/REV7 and REV1 differently interact with the MIP1-deficient mutation in an in vivo assay of mtDNA frameshift mutagenesis. Second, in contrast to TLS repair in the nucleus (Friedberg, Lehmann and Fuchs 2005), cells lacking Rev3 or Rev7 show increased spontaneous and UV-induced point mutagenesis in mtDNA (Kalifa and Sia 2007). The mutagenesis of mtDNA is also increased in rev1 null cells, but only upon exposure to UV. Thus, the TLS polymerases in yeast mitochondria protect mtDNA from accumulating base substitutions. Considering that the nuclear mutagenesis mediated through Rev3/Rev7 and Rev1 is dependent on the formation of the four-subunit complex between Rev3, Rev7, Pol31 and Pol32 and its interaction with PCNA (Makarova, Stodola and Burgers 2012) and that the formation of this complex in mitochondria is not likely, it is not so unexpected that Pol ζ and Rev1 are not as mutagenic in mitochondria as observed in the nucleus. However, both Pol ζ and Rev1 are responsible for the majority of spontaneous and UV-induced frameshift mutations associated with mitochondrial microsatellite sequences and UV-induced petite formation (Kalifa and Sia 2007). This result clearly indicates that Pol ζ and Rev1 contribute to the generation of mtDNA rearrangements during mtDNA replication, even in unstressed cells. On the other hand, the instability of rho⁺ mtDNA in certain mip1 mutants is partially suppressed through the overexpression of REV3, but not REV1, whereas increased mitochondrial point mutagenesis in these mutants is suppressed through the overexpression of both REV3 and REV1, but not of each individual gene (Baruffini et al. 2012). These results suggest that Pol ζ and Rev1 function in yeast mitochondria in several, at least partly separate, lesion-dependent and as yet uncharacterized pathways. To complicate the matter even more, Pol η (Rad30) is also localized to yeast mitochondria (Chatterjee, Pabla and Sleede 2013). In the mitochondria of budding yeast, Rad30 acts in...
at least two mtDNA repair pathways: one pathway reduces UV-induced mitochondrial point mutagenesis, together with Pol ε, and the other pathway counteracts UV-induced rearrangements of mtDNA and petite generation, opposing the activity of Pol ε (Chatterjee, Pabla and Siede 2013). Orthologs of the *S. cerevisiae* polymerases described in this section are in *Sc. pombe*; however, there are no reports about the potential mitochondrial functions of these enzymes.

**Initiation of mtDNA replication and transmission of mtDNA during cell division**

The initiation of mtDNA replication in *S. cerevisiae* is a controversial topic. The debate stems from the long-held opinion that the topology of yeast mtDNA should be similar to the topology of the circular mtDNA in mammalian cells and the circular genomes of the endosymbiotic precursors of mitochondria. Moreover, the genetic map of the budding yeast is also circular. The sequence of *S. cerevisiae* mtDNA contains eight GC (guanine and cytosine)-rich cluster elements that resemble the structures in the heavy-strand replication origin of mammalian mtDNA. These clusters were named ori/rep sequences, and analogous to those detected in mammalian mitochondria, three to four of these elements (Lecrenier and Foury 2000) are thought to be active origins of mtDNA replication initiated via transcription mediated through the mitochondrial RNA polymerase, the Rpo41 protein in *S. cerevisiae* (Fig. 8A). However, cells lacking the Rpo41 polymerase, although unable to maintain rho− genomes, maintain rho+ genomes, even those containing only repeated ori sequences (Lorimer, Brewer and Fangman 1995). In *S. cerevisiae* cells, mitochondrial protein synthesis is required for the maintenance of the rho+ mitochondrial genome (Myers, Pape and Tzagoloff 1985). Thus, it remains unclear whether Rpo41 is actually needed for the initiation of mtDNA replication in budding yeast. Furthermore, many lines of evidence in various yeast species (*S. cerevisiae* and *Sc. pombe*) included have demonstrated that only a small fraction of mtDNA in these cells is actually circular; a majority of the mtDNA molecules are linear and have heterogeneous sizes due to tandem (concatameric) arrays of genomic units (Maleszka, Skelly and Clark-Walker 1991; Bendich 1996). Maleszka et al. also detected circular molecules with single- or double-stranded tails (lariats) and proposed a rolling-circle replication (RCR) mechanism as a model for yeast mtDNA replication, featuring uncoupled synthesis of the leading and lagging strand. In this model, circular mtDNA molecules arise only through the recombination-mediated looping out of a single genetic unit from the concatameric mtDNA array. 2D gel studies on *Sc. pombe* mtDNA confirmed this model of mtDNA replication for fission yeast cells as well (Han and Stachow 1994).

There is now prevailing genetic evidence from budding yeast that the RCR model is true for mtDNA replication in this species (Fig. 8B). Ling and Shibata (2002) have proposed this model to explain the finding that, in *S. cerevisiae*, mother cells primarily contain concatameric mtDNA arrays, whereas in buds mtDNA is predominantly in the form of circular monomers. These authors also identified Mhr1 as the protein required for the transmission of nascent concatameric mtDNA into buds, and this protein catalyzes a recombination reaction involving ssDNA pairing with homologous dsDNA, thereby forming a heteroduplex, the basic intermediate of HR. Surprisingly, Mhr1 is not structurally related to the RecA family of recombinases. In the model by Ling and Shibata, Mhr1-mediated pairing generates a primer for mtDNA RCR, producing a concatameric mtDNA array. This concatameric molecule is subsequently transmitted to buds with a concomitant monomerization of mtDNA. In three follow-up studies, the same authors confirmed this model, showing that (1) Mhr1-dependent RCR initiated at the DSB generated at ori5 is responsible for concatamer formation and the biased inheritance of the rho− genome containing this ori element (Ling, Hori and Shibata 2007); (2) ROS exposure leads to increased mtDNA copy numbers by promoting Mhr1-initiated RCR of mtDNA at ROS-generated DSB ends (Hori et al. 2009); (3) the mitochondrial 5′ exonuclease DIn7 participates with Mhr1 in ROS-enhanced mtDNA replication and recombination at ori5 (Ling et al. 2013).

Therefore, it can be concluded that Mhr1 mediates, at least partially, recombination-dependent replication (RDR) in yeast mitochondria. However, strains with the mhr1−1 allele, encoding an Mhr1 variant that is inactive in the ssDNA pairing activity in vitro (Ling and Shibata 2002), exhibit a moderate phenotype (i.e. temperature dependent loss of rho+ DNA), whereas the deletion of the Mhr1 gene results in the unconditional loss of rho+ genomes (conversion to rho−). Consequently, it is highly likely that another, yet unidentified, Mhr1 activity is indispensable for the maintenance of rho+ mtDNA. In addition, there is recent evidence (Fritsch et al. 2014) supporting the notion that there are more recombinases in *S. cerevisiae* mitochondria than only Mhr1 (Ling et al. 1995). Interestingly, the Mhr1 function is conserved in fission yeast. There is a clear ortholog protein, and inactivation of the coding gene is lethal for the petite-negative yeast *Sc. pombe* (PomBase. http://www.pombase.org/). However, in the Mhr1-dependent concatamer formation model, the mechanism of mtDNA partitioning into buds, i.e. how mtDNA monomers are actually looped out from nascent Mhr1-initiated concatamers before being transmitted to buds, has not yet been elucidated. Future studies should identify other nucleoid proteins that are involved in the partitioning process. This will provide a better understanding of yeast mitochondrial genomes. Currently, another protein, conserved in both yeast species, has been implicated in the Mhr1-related RDR pathway. The mitochondrial *S. cerevisiae* Cce1/Mgt1 structure-selective endonuclease resolves HJs (Kleff, Kemper and S terngranz 1992). In *Sc. pombe*, an orthologous nuclease, Ydc2, exhibits the same activity as the budding yeast homolog (O rum, Keeley and Tsaneva 1998). Strains in both yeast species lacking these nuclease display strikingly similar phenotypes with their mtDNA aggregated in an interlinked network joined by unresolved recombination junctions (Lockshon et al. 1995; Doe et al. 2000). A double *S. cerevisiae* mutant lacking Cce1 and harboring the defective allele mhr1−1 (see above) lacks mtDNA (Ling and Shibata 2002), and thus, these two mutations interact synergistically, suggesting that the two proteins, Cce1 and Mhr1, function in two parallel pathways necessary for mtDNA transmission to progeny cells.

**Okazaki fragment processing in mitochondria**

Information on lagging-strand synthesis in yeast mitochondria is limited. Surprisingly, almost the entire apparatus for Okazaki fragment processing is present in budding yeast mitochondria and in human mitochondria (reviewed in Holt 2009). The Okazaki fragment processing apparatus comprises (1) the largest subunit of Pol α, Pol1, with its accessory subunit Pol12 (Lasserre et al. 2013), (2) the Pol1 helicase (Lahaye et al. 1991; Schulz and Zakian 1994) and (3) the Rad27 endonuclease (Kalifa et al. 2009). The essential Dna2 nuclease/helicase is likely also localized to yeast mitochondria based on genetic evidence (Budd et al. 2006), but direct evidence is lacking. Yeast Pol1 and Dna2 demonstrate that it is exceptionally challenging to study the mitochondrial function of an essential protein with dual nuclear and mitochondrial localization. However, at least for the three classical proteins of the well-characterized Okazaki fragment
Figure 8. Current models of initiation of mtDNA replication in *S. cerevisiae*. (A) Transcription-dependent mtDNA replication. In the model, mtDNA replication is primed by the mitochondrial RNA polymerase Rpo41 at ori/rep sequences and the primer is extended afterwards by the Mip1 polymerase. However, the resulting expanding bubble mtDNA structures (θ structures), that are expected according the model, have not been detected (Maleszka et al. 1991; Bendich 1996). (B) Rolling-circle mtDNA replication mechanism proposed by Maleszka et al. (1991) and later expanded by Ling et al. (2007) and Hori et al. (2009) to explain the increases in mtDNA copy number triggered by ROS. In the model, mtDNA RCR is primed by homologous pairing catalyzed by the Mhr1 recombinase between ssDNA derived from the resection of DSB ends generated by the Ntg1-mediated processing of mtDNA lesions caused by DNA-damaging agents, e.g. ROS, as it has been shown for H$_2$O$_2$-induced Ntg1-dependent DSBs at the ori5 sequence that result in the Mhr1-mediated initiation of mtDNA replication. Two strands of the schematic representation reflect the later stages of RCR, when the lagging-strand synthesis (mediated by the putative mitochondrial Okazaki fragment processing apparatus whose factors are listed in the box) has been initiated on the single-stranded tail produced by RCR which ultimately results in the production of concatameric linear molecules that are the prevalent topological form of mtDNA in yeast cells. In this model, rare circular mtDNA, monomeric or oligomeric, are derived from multisemec molecules through intramolecular recombination events.
processing pathway (reviewed in Balakrishnan and Bambara 2011), Pif1, Rad27 and Dna2, the dual nuclear-mitochondrial localization is conserved in evolution, as orthologs of the three proteins display dual localization in human cells (Futami, Shimamoto and Furuichi 2007; Liu et al. 2008; Duxin et al. 2009), indicating that these proteins could function in a fundamental mitochondrial process, likely associated with the lagging-strand synthesis during mtDNA replication. Nevertheless, the proteins do not function in mitochondria exclusively in this process. In this section, one potential mitochondrial Okazaki fragment processing factor, the budding yeast Pif1 helicase and its fission yeast ortholog, Pfh1, are reviewed. Saccharomyces cerevisiae Rad27 and S. pombe orthologous Rad2 (Fen1) are reviewed in the section on BER in mitochondria.

The Pif1 helicase in mitochondria

The Pif1 helicase, which has 5′→3′ DNA helicase activity, has been associated with several mtDNA transactions, from replication and recombination (reviewed in Contamine and Picard 2000) to repair (O’Rourke et al. 2002; Doudican et al. 2005; Cheng, Dunaway and Ivesa 2007) in S. cerevisiae yeast mitochondria. This helicase and its paralog, Rrm3, are nuclear non-overlapping regulators of replication fork passage through hard-to-replicate regions, such as rDNA and G4 (G-quadruplex) structures (reviewed in Chung 2014). The ability of Pif1 to unwind G4 structures is likely essential for mtDNA replication because in budding yeast, there is a 10-fold higher concentration of the G4 motifs in mtDNA than in nuclear DNA (Capra et al. 2010). Furthermore, Pif1 has recently been shown to facilitate DNA synthesis in the D-loop during HR (Wilson et al. 2013). Strains lacking Pif1 exhibit high rho− mtDNA instability, however, depending on genetic background, the final petite pif1Δ phenotype is either rho′ (Merz and Westerman 2009) or rho– (Doudican et al. 2005). In the latter case, a significant portion of pif1Δ cells maintain rho+ mtDNA, but with a marked accumulation of mitochondrial point mutations (O’Rourke et al. 2005), suggesting that the Pif1 helicase is also involved in mtDNA repair pathways. This conclusion is supported by the synergy between PIF1 and SOD2 deletions. The latter gene encodes mitochondrial matrix superoxide dismutase, and loss of this gene leads to increased mtDNA damage and, consequently, increased mtDNA point mutagenesis. However, loss of this gene does not impair the maintenance of rho+ genomes (Doudican et al. 2005; Kaniak et al. 2009). Furthermore, the maintenance of rho+ mtDNA in pif1Δ cells is synergistically destabilized by concomitant inactivation of the Ntg1 gene, which encodes a nuclear and mitochondrial N-glycosylase/lyase specific for certain oxidatively modified bases (O’Rourke et al. 2002). Interestingly, this protein also stimulates DSBs (the enzymatic mechanism of this Ntg1 activity has not yet been elucidated) at ori5 for Mfrr1-dependent RDR, resulting in concatamer formation and biased inheritance of a rho− ori5-containing genome (Ling, Hori and Shibata 2007). Although the lack of Ntg1 in cells maintaining rho+ mtDNA at wild-type levels does not influence mtDNA stability (e.g. O’Rourke et al. 2002; Doudican et al. 2005; Kaniak et al. 2009), this glycosylase is important for the maintenance of rho+ genomes in cells with destabilized rho+ mtDNA (i.e. pif1Δ cells) and in abf2Δ cells (O’Rourke et al. 2002). These results suggest that Ntg1 might be an element of the recombination-dependent system, regulating mtDNA replication and, consequently, mtDNA copy number. The synergy between the inactivation of either PIF1 or ABF2 and the inactivation of NTG1 suggests that Pif1 and Abf2 are important for rho+ maintenance independently of the Ntg1-mediated pathway, underscoring the complexity of mtDNA transactions in yeast cells.

In S. pombe, the unique Pif1 helicase ortholog Pfh1 is essential for the replication of both nuclear and mitochondrial genomes (Pinter, Aubert and Zakian 2008). In fission yeast cells, the helicase from the budding yeast, S. cerevisiae Pif1, and its nuclear paralog, S. cerevisiae Rrm3, failed to replace both essential nuclear and mitochondrial functions of Pfh1. However, Rrm3 partially suppressed the phenotypes of a nuclear Pfh1-depleted strain. These results suggest that the mitochondrial Pif1-like helicases in budding and fission yeast significantly diverged after the evolutionary separation of the two lineages. Nevertheless, it would be interesting to repeat the inactivation of the pif1 gene in a petite-positive Sc. pombe strain to better characterize the Pfh1 functions.

Mitochondrial BER

The exposure of DNA to endogenous threats (e.g. metabolism-derived ROS) and exogenous genotoxic agents produces various modifications to the structure of constituent bases. These lesions, if they do not distort the DNA helix, are repaired through BER (Baute and Depicker 2008; Svilar et al. 2011). Many BER enzymes in budding yeast have dual nuclear-mitochondrial localization (reviewed in Boiteux and Jinks-Robertson 2013). The pathway exists in two basic variants: a short-patch pathway and a long-patch pathway (Fig. 9A).

The short-patch BER pathway (SP BER) employs various enzymes and involves several stages of intermediate processing: (1) recognition and excision of specific bases (in S. cerevisiae mitochondria, Ung1 recognizes and cleaves off uracil (Chatterjee and Singh 2001), Oggl excises 7,8-dihydro-8-oxo-2′-deoxyguanosine, 8-oxog, a common oxidative modification of guanine, opposite cytosine (Singh et al. 2001) and Ntg1 (a homolog of E. coli endonuclease III, mentioned in the section ‘The Pif1 helicase in mitochondria’) excises oxidized pyrimidines (Alseth et al. 1999)); (2) cleavage of the resulting AP site through either the AP lyase activities of bifunctional Oggl or Ntg1 (in contrast to the monofunctional glycosylase Ung1, which is unable to catalyze this reaction), producing phospho-α,β,γ-unsaturated aldehyde (3’-PUA), a DNA polymerase blocking group on the 3′ end of the gap and a phosphate group on the 5′ end, or through the mitochondrial Apn1 AP endonuclease (Vongsamphan, Fortier and Ramotar 2001), which cuts the phosphodiester backbone of the AP site to generate a strand break with 3′-OH and 5′-terminal deoxyribose-phosphate (5′-dRP) sugar moieties; (3) DNA-end processing to generate DNA 3′ ends that are filled in by a polymerase, and 5′ ends that are subsequently ligated. During the AP lyase reaction, the blocking PUA group is further processed by the 3′-phosphodiesterase activity of Apn1 to generate a 3′-OH end. Upon the AP endonuclease reaction, the 5′-dRP moiety is converted through the 5′-dRP lyase activity of Pol γ (mentioned above) to a 5′-phosphate that can be subsequently ligated; and (4) single-nucleotide gap filling and ligation. The actual role of the Apn1 protein goes beyond the simple SP BER model. Apn1 also functions in DNA repair without the need for a preceding glycosylase. The enzyme nicks DNA on the 5′ side of various oxidatively damaged bases, thereby generating 3′-OH and 5′-phosphate termini (Ischenko and Saparbaev 2002). This nuclease incision repair (NIR) activity is also exhibited by AP endonucleases from other organisms (reviewed in Daley, Zakaria and Ramotar 2010).

Single mutants lacking the individual SP BER enzymes listed above under normal growth conditions do not display a
Figure 9. Mitochondrial BER pathways in yeast cells. (A) Pathways of mtBER in *S. cerevisiae* cells. In the box, presumed factors of mt LP BER pathway are listed, with the Dna2 name dimmed, since its presence in mitochondria, though very likely, has not been directly proved yet. (B) BER pathways in *S. pombe* mitochondria. Highlighted in red are names of those proteins, which have been experimentally confirmed to be localized to mitochondria. In the box, presumed factors of mt LP BER pathway are listed.
phenotype associated with mtDNA stability (Apn1: Vongsamphanh, Fortier and Ramotar 2001), or display moderately increased mtDNA point mutagenesis (Apn1: Kaniak et al. 2009) that might differ, however, in the specificity of substitutions fixed in mtDNA depending on some unknown features of a genetic context (Ogg1: Dzierzbicki et al. 2004; Pogozhala, Mookerjee and Sia 2009), or display even moderately decreased mitochondrial point mutagenesis (Ntg1, Apn1: Pogozhala, Mookerjee and Sia 2009; Ntg1: Kaniak et al. 2009). The phenotype of ung1 null mutants was also described as a moderately increased frequency of respiratory-deficient cells with intact mitochondrial genomes; thus, these mutants displayed the properties of mit− mutants (Chatterjee and Singh 2001). Importantly, apn1 null strains exhibit significantly increased mitochondrial point mutagenesis upon exposure to the alkylating agent methyl methanesulfonate, MMS (Vongsamphanh, Fortier and Ramotar 2001), and show delayed kinetics of alkylation-induced damage repair specifically in mtDNA (Acevedo-Torres et al. 2009). Thus, the SP BER enzymes might recognize specific types of environmentally induced mtDNA damage that are rarely observed under normal growth conditions. Consistently, the Ntg1 glycosylase has been implicated, upon treatment of isolated mitochondria with hydrogen peroxide, in the generation of DSBs at ori5 to initiate Mhr1-mediated replication of mitochondrial genome, thereby increasing the mtDNA copy number (Hori et al. 2009) (Fig. 8A).

The long-patch BER pathway (LP BER) assists SP BER enzymes in 5′-blocked intermediate processing. The distinguishing feature of the two BER pathways is the size of the patch being repaired: one nucleotide in the case of SP BER, as described above, and two or more nucleotides in the case of LP BER (reviewed in Fortini and Dogliotti 2007). LP BER takes over the repair when the 5′-blocking group generated in the first steps of BER, typically when 5′-DRP cannot be removed by polymerase, e.g. the end is further oxidized. First, the 5′-blocking group is converted into a 5′-flap through strong strand displacement synthesis mediated by yeast Pol γ (Vík, Váljamäe and Sedman 2011). Then, a 5′-flap endonuclease removes the short flap, the polymerase fills in the gap of several nucleotides and the ligase completes the repair. The identity of the LP BER flap endonuclease in yeast mitochondria is not clear, but in yeast nuclei, this endonuclease has been identified as Rad27 (Ayyagari et al. 2003), an ortholog of mammalian EXOG and END0 nuclease activities (reviewed in Schneiter 2006). In mammalian cells, the EXOG nuclease activity is required for repairing endogenous 5′-blocking single-strand breaks (SSBs) in mtDNA. EXOG depletion results in the accumulation of persistent SSBs in mtDNA, enhances ROS levels and induces apoptosis (Tann et al. 2011). The Nuc1 nuclease, the S. cerevisiae ortholog of mammalian END0 and EXOG nucleases (Büttnner et al. 2007; Cymerman et al. 2008), is localized to the mitochondria where it functions in HR (Zassenhaus and Denninger 1994). Nuc1 produces recombinogenic DNA ends by introducing single-strand gaps into DNA through endonuclease and 5′→3′ exonuclease activities (Dake et al. 1988). Nuc1 has also been implicated in the degradation of damaged mtDNA under conditions of acute oxidative stress accompanied by a shortage of intramitochondrial energy (Dzierzbicki et al. 2012). In response to apoptotic stimuli, mammalian END0 translocates to the nucleus, where it participates with other nucleases in the fragmentation of chromosomal DNA (Li, Luo and Wang 2001). The Nuc1 nuclease has also been described to function in a similar pathway in response to apoptotic signals in budding yeast (Büttnerr et al. 2007). Another nuclease, Din7, a paralog of the Rad27 nuclease, is localized to mitochondria (Fikus et al. 2000). The role of the protein has not been established yet, but its overproduction results in the increase of the mitochondrial HR, elevated petite formation, mitochondrial microsatellite instability and induced point mutagenesis of mtDNA (Koprowski et al. 2003). The Din7 nuclease is also involved in the Mhr1-dependent RCR of mtDNA (Ling et al. 2013), as mentioned in the section ‘Initiation of mtDNA replication and transmission of mtDNA during cell division’. It remains to be established whether Rad27, Din7, Rev3, Nuc1 and Dna2 are indeed involved in the mitochondrial LP BER pathway in budding yeast cells.

The set of SP BER enzymes in Sc. pombe is markedly divergent from the BER enzymes in budding yeast cells (Table 3). There are more mono-functional glycosylases in fission yeast cells, similar to higher eukaryotic cells. Nevertheless, only one bifunctional Nth1 functions in Sc. pombe cells, presumably, providing the major, if not unique, AP lyase activity for processing AP sites (reviewed in Kanamitsu and Ikeda 2010). However, the mitochondrial localization of BER glycosylases in fission yeast cells remains unclear (Fig. 9B). As in S. cerevisiae, there are two AP endonucleases in Sc. pombe: Apn1, the homolog of the bacterial exonuclease III (Xth), and Apn2, the homolog of the bacterial exonuclease IV (Nfo), and these AP endonuclease orthologs evolved differently in the two yeast species (Kanamitsu and Ikeda 2010). Interestingly, Sc. pombe AP endonucleases do not cut AP sites, which are cut by Nth1. Instead, they remove the 3′-blocked ends that remain after Nth1 AP lyase activity. However, another Nfo-like endonuclease in fission yeast cells, Uve1, is localized to both the nucleus and mitochondria. This
Table 3. Comparison of BER and SSB repair enzymes in Sc. pombe, S. cerevisiae and human cells in respect to their subcellular localization (PomBase. http://www.pombase.org/; Saccharomyces Genome Database. http://www.yeastgenome.org/; Online Mendelian Inheritance in Man. http://www.omim.org/).

| Description of the protein | Saccharomyces cerevisiae (subcellular localization) | Schizosaccharomyces pombe (subcellular localization) | Human ortholog (subcellular localization) |
|---------------------------|---------------------------------------------------|-------------------------------------------------|-----------------------------------------|
| **SP BER pathway**        |                                                   |                                                 |                                         |
| Uracil/thymine (in G:T) DNA | –                                                 | Thp1 (n)                                        | hTDG (n)                                |
| N-glycosylase Thp1        |                                                   |                                                 |                                         |
| Uracil DNA N-glycosylase Ung1 | Ung1 (mt/n)                                      | Ung1 (mt?/n)                                   | hUNG (mt/n)                             |
| DNA-3-methyladenine glycosylase | Mag1 (n)                                        | Mag1 (n)                                       | –                                       |
| DNA-3-methyladenine glycosylase | Mag1 (n)                                        | Mag2 (n)                                       | –                                       |
| Adenine (in 8-oxoG:A) DNA glycosylase | –                                              | Myh1 (n)                                       | hMUTYH (n)                              |
| Homolog of endonuclease III from E. coli | Ntg1α (mt/n), Ntg2α (n) | Nth1 (mt?/n)                                   | hNTHL1 (mt/n)                           |
| 8-Oxoguanine glycosylase | Ogg1 (mt/n)                                      | –                                              | hOGG1 (mt/n)                            |
| Homolog of E. coli exonuclease III [Xth]; | Apn2 (n)                                        | Apn2 (n)                                       | hAPEX2 (mt/n)                           |
| AP-endonuclease Apn2      |                                                   |                                                 |                                         |
| Homolog of E. coli endonuclease IV [Nfo]; AP-endonuclease Apn1 | Apn1 (mt/n) | Apn1β (mt/n)                                  | –                                       |
| Endonuclease Uve1         |                                                   |                                                 |                                         |
| Homolog of endonuclease VII of E. coli | –                                              | Uve1 (mt/n)                                    | –                                       |
| Homolog of endonuclease VII of E. coli | –                                              | –                                              | NEIL1 (mt/n)                            |
| ERCC-8 DNA repair homolog | Rad28 (n)                                        | Ckn1 (mt?/n)                                   | ERCC8 (mt/n)                            |
| SNF2 family helicase Rhp26 | Rad26 (n)                                        | Rhp26 (mt?/n)                                  | ERCC6 (mt/n)                            |
| **LP BER pathway**        |                                                   |                                                 |                                         |
| Flap endonuclease, FEN1 ortholog | Rad27 (mt/n)                                   | Rad2 (mt?/n)                                   | FEN1 (mt/n)                             |
| Mitochondrial endo- and exonuclease | Nuc1 (mt/n)                                    | Pnu1 (m)                                       | ENDOG (mt/n), EXOG (mt)                  |
| DNA replication endonuclease-helicase | Dna2 (mt?/n)                                  | Dna2 (mt?/n)                                   | DNA2 (mt/n)                             |
| Exonuclease Exo1          | Exo1 (n)                                         | Exo1 (n)                                       | EXO1 (n)                                |
| Exonuclease Exo1          | Din7 (mt?)                                       | Exo1 (n)                                       | EXO1 (n)                                |
| **SSB repair**            |                                                   |                                                 |                                         |
| Aprataxin                 | Hnt3 (n)                                         | Hnt3 (mt?/n)                                   | APTX (mt/n)                             |
| Tyrosyl-DNA phosphodiesterase Tdp1 | Tdp1 (n)                                      | Tdp1 (n)                                       | TDP1 (mt/n)                             |

mt—mitochondrial localization; n—nuclear localization.

enzyme functions in an alternative AP endonuclease pathway, as mentioned above for S. cerevisiae Apn1, catalyzing the NIR-mediated removal of a variety of UV and non-UV lesions. Consequently, it is presumed that in fission yeast cells, both dual-localization Apn1 and Uve1 engage in an Nth1-independent NIR, and 5'-blocked DNA ends generated in NIR reactions can be further processed through the LP BER pathway. Interestingly, according to a recent report, most laboratory Sc. pombe strains, but not other independent isolates, carry a defective mutation in the apn1 gene (Laerdahl et al. 2011). Nevertheless, even in the context of the functional Apn1 endonuclease, these authors showed that Nth1 remains responsible for the majority of the AP lyase activity in cell extracts. Considering that the mitochondrial localizations of both Nth1 and Apn2 are unclear, the dual localization of its functional variant, Apn1, might significantly contribute to the AP lyase and NIR activities in fission yeast mitochondria. Future studies are needed to confirm this idea. In Sc. pombe cells, there are also orthologs of all the putative S. cerevisiae mitochondrial LP BER proteins described above. However, there are no reports of mitochondrial functions of these proteins, except for the confirmation of the mitochondrial localization of the fission yeast ortholog of Nuc1, Pnu1 (Oda et al. 2007). In pnu1 null cells, the ectopic production of a truncated Pnu1 lacking the mitochondrial targeting signal sequence, i.e. a fully active form of the nuclease generated during Pnu1 import to the mitochondria, results in cell death due to DNA fragmentation (Oda et al. 2007), suggesting that ENDOG orthologs may indeed be involved in a conserved apoptotic pathway in eukaryotic cells.

**Dual role for the Msh1 protein in mutation avoidance and rho− genome stability**

The MSH1 gene encodes a homolog of E. coli MutS protein (Reenan and Kolodner 1992), the critical component of the bacterial MMR pathway. In vitro, the Msh1 protein hydrolyzes ATP and recognizes DNA substrates containing mismatches and unpaired nucleotides (Chi and Kolodner 1994). MutS homologs generally act by initiating the repair of base substitution and insertion–deletion mismatches that arise during either replication or recombination (reviewed in Schofield and Hsieh 2003). Strains lacking this gene exhibit a petite phenotype due to rho− deletions, suggesting that Msh1 plays an important role in the maintenance of rho− mtDNA stability. In addition, Msh1 is required in yeast mitochondria to prevent the accumulation of point mutations in mtDNA as a crucial factor in the proposed, but as yet uncharacterized, pathway of mitochondrial MMR. Strains with partial Msh1 defects, e.g. msh1Δ/ΔMSH1 heterozygous strains, exhibit increased accumulation of point mutations in the mtDNA (Reenan and Kolodner 1992). Thus, the MSH1 gene is haplo insufficient, suggesting that the Msh1 level is
fine-tuned for optimal mitochondrial genome maintenance. Accordingly, the overexpression of the MSH1 gene leads to the instability of rho+ genomes and increases the instability of poly(AT), but not poly(GT) repeats, through an unknown mechanism (Koprowski et al. 2002). However, moderate MSH1 overexpression suppresses the mitochondrial mutator phenotype of the ogg1Δ mutant (Dzierzbicki et al. 2004) and the mitochondrial mutator phenotype of the sod2Δ mutant (Kaniak et al. 2009). Additionally, the increased dosage of wild-type MSH1, but not of mutant alleles encoding deficient variants of the protein, results in enhanced allelic mitochondrial HR (Kaniak et al. 2009). On the other hand, the Msh1 activity suppresses direct repeat-mediated deletions, thereby inhibiting genomic rearrangements that may result in defective mtDNA (Mookerjee and Sia 2006). Therefore, Msh1 promotes allelic HR in mitochondria, but inhibits non-allelic direct-repeat-mediated deletion events and GT-repeat frameshift mutations. We propose that the Msh1 protein scans mtDNA duplexes, generated during either replication or recombination-associated strand invasion or strand annealing, for the presence of mismatches, and it directs the duplexes into either HR combined with mismatch correction or homoduplex recombination and inhibition of the incipient recombination, depending on the quality of the duplex. Thus, on the one hand, Msh1 is a key player in the mitochondrial mutation avoidance system, and on the other hand, it is a chief controller of recombination processes in yeast mitochondria. The two functions of Msh1 are genetically separated (Mookerjee, Lyon and Sia 2005), which should facilitate the discrimination of Msh1 functions. Consistently, Pogorzala, Mookerjee and Sia (2009) also showed that Msh1 plays multiple roles in mitochondrial BER as a part of mitochondrial mutation avoidance, but this function is separate from the Msh1 function required for rho+ genome maintenance. A mitochondrial ortholog of Msh1 has been identified in S. pombe cells (Matsuyama et al. 2006); however, its role in fission yeast mitochondria has not been determined.

HR in yeast mitochondria

The pathways of mitochondrial recombination in yeast are not well understood (reviewed in Chen 2013). HR is highly active in S. cerevisiae mitochondria, e.g. markers separated by only 1 kb are practically unlinked (Dujon 1981). As mentioned in the section ‘Initiation of mtDNA replication and transmission of mtDNA during cell division’, the current model of initiation of mtDNA replication posits that HR-dependent processes generate primers for the mitochondrial RCR (Ling and Shibata 2002). The model explains the ROS-induced increase in mtDNA copy numbers through Mhr1-initiated Din7-dependent RCR of mtDNA at ROS-generated DSB ends (Hori et al. 2009; Ling et al. 2013). However, none of the proteins described in the Mhr1-dependent pathway (the DSB-generating Ntg1 glycosylase, the exonuclease Din7 or the Mhr1 recombinase), or in other recombination-related processes in mitochondria, like the Nuc1 nuclease (section ‘Mitochondrial base excision repair’) or the Cce1 resolvase (section ‘Initiation of mtDNA replication and transmission of mtDNA during cell division’), is essential for mitochondrial HR (Ling et al. 1995; Phadnis, Sia and Sia 2005; Fritsch et al. 2014). As mentioned above, genetic evidence suggests that the nuclease-helicase Dna2, the key end resection nuclease in the nuclear HR (reviewed in Mimitou and Symington 2011), functions in the yeast mitochondria and is necessary for the maintenance of rho+ mtDNA (Budd et al. 2006), though direct evidence is missing. The Pif1 helicase has been implicated for some time in the regulation of mitochondrial recombination (reviewed in Contamine and Picard 2000). The finding that during BIR and crossover recombination in the nucleus Pif1 facilitates DNA synthesis by promoting the establishment of a migrating D-loop structure (Wilson et al. 2013) suggests the possibility that Pif1 is a crucial factor mediating RDR of mitochondrial genomes. Consistently, the Pif1 helicase physically interacts with a mitochondrial single-stranded-binding protein, Rim1 (Ramanagounder-Bhojappa et al. 2013). Nevertheless, there is also evidence that the function of another mitochondrial helicase Hml1, that is required for the maintenance of rho+ genomes, partially overlaps with that of the Pif1 helicase (reviewed in Chen 2013). Clearly, there are several mitochondrial HR pathways operating in mitochondria that remain to be clarified. In addition, the set of mitochondrial proteins involved in DNA transactions has recently increased by components of the MRX complex and Ku proteins, as described in the following section, showing that the DSB repair pathways in the nucleus and the mitochondria share some central components. Last but not least, a new SSA activity has been uncovered for the Mgm101 protein (Mbantenkhu et al. 2011), as described in the section ‘The Mgm101 protein is required for rho+ 2305 genome stability’, offering the alternative to the Mhr1 recombination activity.

MRX complex and Ku proteins function in mitochondria

In a recent report, Kalifa et al. (2012) have provided evidence that the factors involved in HR and NHEJ in the nucleus also function in the mitochondria. These authors showed that the inactivation of crucial DSB repair pathways mediated by the MRX complex and Ku proteins synergistically reduces the rate of deletions mediated by direct repeats in yeast mtDNA. The results suggest that these deletions primarily originate from the processing of DSB lesions in the repeated sequences via two independent pathways: mitochondrial MRX and mitochondrial Ku proteins. Surprisingly, the decrease in the direct repeat deletions in mtDNA in a strain lacking both MRX and Ku complexes was associated with a modest but significant increase in spontaneous petite mutants. These results indicate that the mechanisms of direct repeat-mediated deletions and rearrangements responsible for petite formation are mechanistically different. Under certain conditions, both pathways, mitochondrial HR and NHEJ, contribute to the maintenance of the rho+ genome. However, in another study, Dzierzbicki et al. (2012) showed that mutants deficient in the MRX function exhibit increased susceptibility to oxidative stress-induced rearrangements in mtDNA in antymycin A-treated cells. Under these conditions, the inactivation of RAD50 is not synergistic, but epistatic to YKU70 deletion, suggesting that in this case, the function of Rad50 is engaged in a different process than the function detected during the direct deletion-mediated rearrangements observed by Kalifa et al. (2012). In the nucleus, the MRX complex is involved in both HR and NHEJ pathways, therefore the MRX complex likely also participates in a mitochondrial HR pathway that contributes to the maintenance of rho+ genomes under oxidative stress conditions. Consistently, the elevated petite formation in cultures of antymycin A-treated rad50Δ cells correlates with the decreased potential for sustaining mitochondrial allelic HR. Thus, under certain conditions, the mitochondrial MRX-dependent pathway plays a significant role in the maintenance of rho+ mtDNA. Other factors acting in this mtDNA repair pathway remain unknown. Similarly, other components of mitochondrial NHEJ have not yet been identified.
The Mgm101 protein is required for rho\(^{-}\) genome stability

The mitochondrial nucleoid protein Mgm101 is essential for the maintenance of rho\(^{+}\) mtDNA, but not rho\(^{-}\), genomes (Zuo, Clark-Walker and Chen 2002). However, in certain genetic contexts, strains lacking the Mgm101 are rho\(^{0}\) (Crider et al. 2012). Mgm101-deficient cells are more sensitive to mtDNA damage induced through UV irradiation and are hypersensitive to mtDNA damage induced through \(\gamma\) rays and hydrogen peroxide treatment (Meeusen et al. 1999). Consequently, Mgm101 performs an essential function in the repair of oxidatively damaged mtDNA and is critical for maintaining the mitochondrial genome. Meeusen and Nunnari (2003) showed that Mgm101 forms part of a two-membrane spanning (TMS) protein structure that traverses both the outer and inner mitochondrial membrane and associates with replicating nucleoids. The Mmm1 outer membrane protein and the Mip1 polymerase are also components of this structure. Interestingly, the Pif1 helicase is also a component of an inner mitochondrial membrane complex (Cheng and Ivessa 2010). It is not clear whether the Mgm101-containing TMS structure is the same as the Pif1 complex.

The report by Mbantenku et al. (2011) has shown that Mgm101 shares conserved motifs with the N-terminal ssDNA-anealing domain of the yeast Rad52 protein. Rad52 primarily functions as a mediator of Rad51 recombinase activity, but this protein also has ssDNA-anealing activity nucleus (as mentioned in the section ‘HR subpathways’). Consistently, the authors demonstrated that purified Mgm101 promotes ssDNA-anealing reactions in vitro in the presence of the mitochondrial ssDNA-binding protein Rim1. Moreover, Mgm101-deficient cells exhibit decreased rates of direct repeat-mediated deletions in mtDNA, consistent with Phadnis, Sia and Sia (2005) who proposed that the pathways affecting these rearrangements of mtDNA predominantly lead to non-crossover exchanges. Certainly, new insights about the mitochondrial functions of S. cerevisiae Mgm101, Pif1 and Msh1 will be informative for research on Sc. pombe mitochondria. All of these proteins have orthologs in fission yeast cells and the systems of mtDNA replication and inheritance are comparable between these species. Surprisingly, S. cerevisiae Mgm101 also forms a complex with the FANCM ortholog Mph1 helicase and MutS\(_{\alpha}\) (Msh2–Msh6 complex) in the nucleus and participates in the recombination-dependent repair of interstrand cross-link lesions in nuclear DNA (Ward et al. 2012). However, it is unclear whether similar Mgm101-dependent pathways function in yeast mitochondria.

It should be noted that the Mmm1 protein, a component of the above-mentioned TMS structure, has been demonstrated to be also an endoplasmic reticulum (ER) transmembrane protein and a part of the ER-mitochondria encounter structure (ERMES), a multiprotein complex physically tethering the ER with mitochondria (Kornmann et al. 2009). Strains defective in the ERMES function display an abnormal mitochondrial morphology and a deficiency in the maintenance of rho\(^{-}\) mtDNA (Hobbs et al. 2001; Kornmann et al. 2009). It has been suggested that the ERMES contact sites serve to link the segregation of mtDNA with the mitochondrial division (Murley et al. 2013). Studies on the interplay between mitochondrial nucleoids and ERMES proteins as well as other cellular components interacting with them will be important for understanding the pathways of mtDNA inheritance in yeast cells.

MITOCHONDRIAL INFLUENCE ON THE NUCLEAR GENOME

Mitochondria are crucial for diverse array of cellular processes, including energy metabolism, metabolic pathways involving lipids, amino acids, nucleotides and iron, signaling and programmed cell death (apoptosis). However, one of the mitochondrial pathways has been recognized in budding yeast as essential for cell viability: the generation and maturation of Fe-S prosthetic groups (ISCs) for both mitochondrial and extramitochondrial Fe-S proteins (Kispal et al. 1999; reviewed in Lill and Kispal 2000; Kaniak-Golik and Skoneczna 2015). ISCs facilitate redox enzyme activities: they are also components of various protein complexes, cytosolic ribosomes included (Kispal et al. 2005), and, in addition, are required for various enzymes, including those engaged in DNA replication (Netz et al. 2012; reviewed in White and Dillingham 2012). It is thus not surprising that functional mitochondria are crucial for maintaining the stability of cellular genetic information. This requirement is not limited to the mitochondrial genome; it also extends to the stability of the nuclear genome (reviewed in Kaniak-Golik and Skoneczna 2015). In yeast cells, mitochondrial dysfunction leads to a nuclear mutator phenotype (measured by the frequency of canavanine-resistant colonies), regardless of whether this dysfunction can be attributed to blocking oxidative phosphorylation with antimycin A at mitochondrial complex III, deletions in mtDNA (rho\(^{-}\)) or the complete loss of mtDNA (rho\(^{0}\)) (Rasmussen et al. 2003). The mutator phenotype is not caused by increased intracellular ROS levels because although antimycin A treatment elevates ROS levels, ROS levels are decreased in the rho\(^{-}\) and rho\(^{0}\) strains. Moreover, nuclear mutations arising in rho\(^{0}\) cells depend on the Rev1 protein and DNA Pol \(\zeta\), whereas those resulting from antimycin A treatment do not. In a recent study, Dirick et al. (2014) showed that the genomic instability specific to rho\(^{0}\) diploid cells results from DNA breaks and mitotic recombination, primarily occurs in non-dividing cells, and tends to fluctuate depending on the environmental conditions. Consequently, mitochondrial dysfunction is mutagenic, and multiple pathways are likely responsible for the resulting nuclear mutator phenotype. Similar conclusions can be drawn from other studies showing that ROS elevation due to mitochondrial disorder and subsequent oxidative damage cause the destabilization of the mitochondrial and/or nuclear genome (Malc et al. 2009; Yazgan and Krebs 2012). Other studies have confirmed the notion that the nuclear DNA instability phenotype is not generated by mitochondrial ROS but by defects in other processes (reviewed in Kaniak-Golik and Skoneczna 2015), such as dNTP production (Debler et al. 2007), ISC formation (Veatch et al. 2009), metal ion detoxification/homeostasis (Karthikeyan, Lewis and Resnick 2002) or apoptosis (Gao et al. 2011). When considering mitochondrial dysfunction, it is important to differentiate between a mutation in a single mitochondrial gene encoding a component of the respiratory chain, which is necessary for proper mitochondrial function and rho\(^{-}\) or rho\(^{0}\) mutations. All of these mutations affect cellular energy production through oxidative phosphorylation, but the former preserves the mitochondrial genome and the production of other mitochondrially encoded proteins, whereas the latter disrupts the ability to synthesize those proteins (components of the mitochondrial translation apparatus that are required for the process are also encoded by mtDNA, as described in the section ‘Mitochondria and genome stability’). Respiratory-deficient rho\(^{+}\) mutants do not exhibit as robust a nuclear mutator phenotype as that of the rho\(^{0}\) strains (Veatch
CONCLUDING REMARKS

The long evolutionary history of life has brought mechanisms of genome stability to perfection. Various mechanisms have been employed to maintain constant genome content. In yeast cells, replication is accurate, errors and lesions are effectively removed from DNA, the breaks in DNA are ligated, abnormal DNA structures arising during cell cycle are resolved, cell divisions are causally monitored to assure equal chromosome segregation, and DNA damage and replication blocks are detected and appropriately responded to, all with respect to cell phase, ploidy, metabolic activity and environmental conditions. However, occasionally, the scrupulous protection of the genome can lead directly to death, and only by accepting the risk of mutation can a cell enable itself to survive. This reflects the evolution of mutation tolerance mechanisms in parallel with genome maintenance mechanisms. Optimizing the fidelity of replication and the balance between the efficiency and costs of DNA repair assures success in future generations that must adapt to a changing environment. Thus, genetic variations contribute to survival and evolution.

In this review, we have focused on the mechanisms leading to somatic genome instability in yeast, providing a comprehensive overview of how these pathways are orchestrated in eukaryotic cells and describing how cells pay a high mutagenic price for survival. All processes ensuring genome stability are highly conserved from yeast to humans, so the significant findings in yeast can be extrapolated to vertebrates, greatly facilitating the molecular analysis of these complex regulatory networks.

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