Knotty Problems during Mitosis: Mechanistic Insight into the Processing of Ultrafine DNA Bridges in Anaphase

KATA SARLÖS,1 ANDREAS BIEBRICHER,2 ERWIN J.G. PETERMANN,2 GIJS J.L. WUITE,2 AND IAN D. HICKSON1

1Center for Chromosome Stability and Center for Healthy Aging, Department of Cellular and Molecular Medicine, University of Copenhagen, 2200 Copenhagen N, Denmark
2Department of Physics and Astronomy and LaserLab, Vrije Universiteit Amsterdam, 1081 HV Amsterdam, The Netherlands
Correspondence: iandh@sund.ku.dk

To survive and proliferate, cells have to faithfully segregate their newly replicated genomic DNA to the two daughter cells. However, the sister chromatids of mitotic chromosomes are frequently interlinked by so-called ultrafine DNA bridges (UFBs) that are visible in the anaphase of mitosis. UFBs can only be detected by the proteins bound to them and not by staining with conventional DNA dyes. These DNA bridges are presumed to represent entangled sister chromatids and hence pose a threat to faithful segregation. A failure to accurately unlink UFB DNA results in chromosome segregation errors and binucleation. This, in turn, compromises genome integrity, which is a hallmark of cancer. UFBs are actively removed during anaphase, and most known UFB-associated proteins are enzymes involved in DNA repair in interphase. However, little is known about the mitotic activities of these enzymes or the exact DNA structures present on UFBs. We focus on the biology of UFBs, with special emphasis on their underlying DNA structure and the decatenation machineries that process UFBs.

Visible evidence of mitotic chromosome segregation problems, such as lagging chromatin or bulky (chromatinized) DNA bridges, has long been used as a marker of genomic instability (McClintock 1938, 1942; Gisselsson et al. 2000, 2002; Hoffelder et al. 2004; Thompson and Compton 2011). These structures are generally revealed by staining with DNA dyes such as DAPI. This explains why ultrafine bridges (UFBs) had escaped detection for decades because they cannot be visualized using any of the commonly used dyes (Fig. 1A). Furthermore, because they are dechromatinized, they also cannot be detected by staining for histones. Instead, they were originally revealed through studies of the mitotic localization of DNA processing enzymes, such as the BLM helicase defective in Bloom's syndrome (Chan et al. 2007) or the Polo-like kinase 1 interacting checkpoint helicase (PICH) (Fig. 1A; Baumann et al. 2007). One curious feature of UFBs is the fact that they are generally coated along their length with PICH/BLM even when they are several microns in length in late anaphase.

A number of studies have investigated the mechanisms by which UFBs are generated and resolved (Wang et al. 2008, 2010b; Chan and Hickson 2009; Naim and Rosselli 2009; Nielsen et al. 2015). It is known that UFBs can be induced by exposure to a range of stressors, and that they often arise from defined genomic loci (centromeres, common fragile sites [CFSs], telomeres, and ribosomal DNA [rDNA]). Moreover, interfering with the functions of UFB-binding proteins has serious consequences for mitosis and genome integrity, such as the generation of aneuploidy, binucleation, and micronucleus formation (Lukas et al. 2011; Nielsen et al. 2015). In our laboratory, we are developing tools to reconstitute anaphase chromosome segregation in vitro. For this, we are investigating the action of recombinant enzymes present at UFBs by combining ensemble biochemical methods with single-molecule optical tweezers combined with fluorescence microscopy. Here, we summarize our current knowledge on UFBs based on cellular observations and introduce our in vitro approaches to build a mechanistic model of sister chromatid disjunction.

THE ORIGINS OF UFBs

Replication causes the newly replicated strands to be interlinked/catenated (Schvartzman and Stasiak 2004; Vos et al. 2011). In parallel to this, the cohesin complex is deposited along the chromosomes to encircle and hold the sister chromatids together (Tanaka et al. 2001; Nasmith 2011). Most of the DNA catenanes are removed by topoisomerase IIα (TopIIα); either during S phase or during early mitosis when DNA condensation occurs (Hirano 2015). In the prophase of mitosis, most of the cohesin located on chromosome arms (but not at centromeres) is released in a condensation-dependent manner (Hirano 2015). The activities of TopIIα, condensin I and II, and cohesin are tightly coordinated and give rise to the classical, X-shaped, chromosome structure, where the arms are devoid of both DNA cohesion and DNA cate-
nanes (Hirano 2015). Interfering with either cohesion or condensation gives rise to UFBs connecting the chromosome arms, suggesting that these processes give directionality to decatenation by TopII (Baxter and Aragón 2012; Minocherhomji et al. 2015; Piskadlo et al. 2017).

**UFBs at Centromeres**

UFBs arising from the centromeres (C-UFBs) can be identified by the presence of centromeric markers such as CENP-A at their termini (Fig. 1B). C-UFBs are by far the most prevalent of all UFBs (Chan et al. 2007, 2009) and exist in every mitosis. Their number is minimized by an active removal process that occurs at anaphase onset (Wang et al. 2008). Importantly, inhibition of TopIIα by specific drugs, such as ICRF-193, induces the persistence and number of centromeric UFBs (Baumann et al. 2007; Chan et al. 2007; Wang et al. 2008). Moreover, TopIIα colocalizes with PICH on ICRF-193-induced UFBs (Nielsen et al. 2015), indicating that TopIIα is required for their decatenation. Because centromeric cohesion is protected from release by Sgo1-PP2A (Kitajima et al. 2006), and is only cleaved at anaphase onset by Separase (Uhlmann et al. 2000), it is thought that TopIIα only has a brief period in which to decatenate any C-UFBs after cohesin cleavage (Wang et al. 2010b). However, based on their frequency, it is also conceivable that the persistence of centromeric UFBs until anaphase is not simply an unwanted side effect of the masking of DNA catenation by cohesion, but rather has a physiological role in maintaining DNA-based sister chromatid “cohesion” until the metaphase-to-anaphase transition.

**UFBs at Common Fragile Sites**

CFSs are viewed as an Achilles’ heel of the genome. They are frequently deleted or rearranged in cancer cells and can appear as gaps or breaks on mitotic chromosomes following replication perturbation (termed CFS expression) (Glover et al. 1984; Durkin and Glover 2007). CFSs are regions where replication is problematic and delayed (Debatisse et al. 2012). According to recent modeling studies, cells with large genomes enter mitosis with, on average, three underreplicated sites per cell, even in unperturbed growth conditions. This problem is exacerbated by conditions that induce replication stress or by reducing the number of origins (Al Mamun et al. 2016; Moreno et al. 2016). The Fanconi anemia (FA) DNA repair proteins, FANCD2 and FANCI (Sims et al. 2007; Smogorzewska et al. 2007), associate with CFSs after replication stress and serve as surrogate markers for these loci (Chan et al. 2009; Naim and Rosselli 2009). In contrast to centromeric UFBs, CFS–UFBs (Fig. 1B) rarely appear spontaneously and cannot be induced by inhibiting TopIIα (Chan et al. 2009). Rather, CFS–UFBs accumulate after perturbation of DNA replication by the DNA polymerase inhibitor aphidicolin (Chan and Hickson 2009; Naim and Rosselli 2009). This suggests that CFS–UFBs are composed of underreplicated DNA.

**UFBs at Telomeres**

The ends of linear eukaryotic chromosomes are organized into well-defined structures called telomeres (Dok-sani and de Lange 2014; Arnould and Karlsseder 2015). The telomeric DNA is looped back in a DNA structure called a T-loop that prevents the DNA end from being exposed (Griffith et al. 1999; Dok-sani and de Lange 2014; Arnould and Karlseder 2015). The shelterin complex, which comprises several proteins including telomeric repeat-binding factors 1 and 2 (TRF1 and TRF2) (Palm and de Lange 2008). T-loops are stabilized by the shelterin complex, which comprises several proteins including telomeric repeat-binding factors 1 and 2 (TRF1 and TRF2) (Palm and de Lange 2008). T-loops are stabilized by the shelterin complex, which comprises several proteins including telomeric repeat-binding factors 1 and 2 (TRF1 and TRF2) (Palm and de Lange 2008). Telomeres show similarities to CFSs in that replication stress induces the so-called “fragile-telomere” phenotype, where the chromosomes appear to be broken at the very end (Sfeir et al. 2009), indicating that these loci are also inherently difficult to replicate (Martinez and Blasco 2015; Higa et al. 2017). Telomeres give rise to BLM-coated UFBs (T-UFBs) (Fig. 1B) following exposure to aphidicolin (Chan and Hickson 2009; Barefield and Karlsseder 2012; d’Alcontres et al. 2014). Interfering with the integrity of the shelterin complex via changing the levels of TRF1 or TRF2 also induces telomere fragility (Martinez et al. 2009; Sfeir et al. 2009) and gives rise to T-UFBs (d’Alcontres et al. 2014; Nera et al. 2015). TRF1 was shown to protect against fragility by recruiting BLM to the telomeres (Sfeir et al. 2009), suggesting that BLM facilitates replication (Drosopoulos et al. 2015) or disentangles late-replicating...
structures at these regions (Chan et al. 2009; Barefield and Karlseeder 2012). In contrast to CFS-UFBs, inhibition of TopIIα by ICRF-193 induces T-UFBs, and TRF1 has been shown to recruit TopIIα to telomeres. These findings suggest that at least a subset of T-UFBs are likely to be completely replicated, double-stranded DNA (dsDNA) catenanes (d’Alcontres et al. 2014).

**UFBs at the rDNA**

PICH is present at rDNA loci in chicken and human cells in early mitosis (Nielsen et al. 2015). The rDNA loci also occasionally give rise to PICH- and TopIIα-decorated R-UFBs (Fig. 1B; Nielsen and Hickson 2016). The number of R-UFBs increases following inhibition of TopIIα by ICRF-193. This suggests that the structure and decatenation mechanism of R-UFBs are similar to those of C-UFBs. The rDNA locus is known to be segregated late during mitosis in yeast (Sullivan et al. 2004; Wang et al. 2004; Clemente-Blanco et al. 2009) and has been shown to be transcriptionally active even in early mitosis in humans (Gebrane-Younes et al. 1997; Sirri et al. 1999; Voit et al. 2015). Because active transcription interferes with condensation, which, in turn, is required for decatenation by TopIIα (Lukas et al. 2011; Baxter and Aragón 2012), this would leave cells only a short time window in which to decatenate the rDNA during mitosis, thus potentially explaining the appearance of UFBs from these loci.

**UBF RECOGNITION AND PROCESSING MACHINERIES**

**PICH**

PICH was first identified as an interacting factor of the mitotic kinase Plk1 (Baumann et al. 2007). PICH is excluded from the nucleus during interphase and is only recruited to chromatin after nuclear envelope breakdown, whereupon it accumulates at centromeric loci. PICH seems to be the main recognition and recruitment factor for UFBs, as several other UFB-processing factors fail to localize to UFBs in the absence of PICH, such as members of the Bloom syndrome complex (Chan et al. 2007) and RIF1 (Hengedal et al. 2015). This makes it difficult to detect or analyze UFBs in the absence of PICH. PICH was reported to influence chromosome condensation, as chromosome structure is abnormal in the absence of PICH (Leng et al. 2008; Kurasawa and Yu-Lee 2010; Rouzeau et al. 2012; Nielsen et al. 2015). PICH belongs to the SNF2 family of translocases (Singleton et al. 2007) and contains a motor domain typical in this enzyme family (Fig. 2A). Consistent with this, PICH possesses ATP-dependent dsDNA translocase activity (Biebricher et al. 2013). In addition to the SNF2 region, PICH has accessory domains, including the PICH-family domain, and two TPR motifs reported to be involved in protein interactions (Hengedal et al. 2015; Pitchai et al. 2017). PICH appears to have a high affinity for stretched dsDNA, which is consistent with the idea that UFBs must be under considerable tension created by the mitotic spindle (Baumann et al. 2007; Biebricher et al. 2013). Indeed, this property of PICH may be the main mechanism for how cells normally sense UFBs. Somewhat surprisingly, a PICH mutant lacking ATPase activity does not increase the number of UFBs, although it does prolong their persistence (Nielsen et al. 2015) and also increases the number of chromatin bridges, indicating that UFBs and chromatin bridges have different origins (Kaulich et al. 2012). It should be noted, however, that the ATPase-dead PICH shows altered localization on metaphase chromosomes (Kaulich et al. 2012).

**The Bloom Syndrome Protein Complex**

BLM is the helicase mutated in Bloom syndrome (BS), a severe autosomal hereditary disorder causing genetic instability and cancer (Ellis et al. 1995; German et al. 2007; Cunniff et al. 2017). BLM belongs to the RecQ family, a group of evolutionary conserved genome caretaking enzymes (Chu and Hickson 2009; Croteau et al. 2014), and comprises a helicase core, flanked by long amino- and carboxy-terminal domains responsible for protein–protein interactions (Fig. 2B; Wu et al. 2000, 2001; Meetei et al. 2003; Doherty et al. 2005; Wang et al. 2013; Blackford et al. 2015). BLM efficiently unwinds various DNA structures such as replication forks (Karow et al. 1997), four-way junctions (Karow et al. 2000), D-loops (Bachrati et al. 2009) and, G4 quadruplexes (Sun et al. 1998). Way junctions (Karow et al. 2000), D-loops (Bachrati et al. 2009) and, G4 quadruplexes (Sun et al. 1998).
2000), a Type 1A topoisomerase that can catalyze only single-stranded DNA (ssDNA) strand passage (Wallis et al. 1989; Vos et al. 2011). TopIIIα is composed of a conserved type 1A topoisomerase domain and multiple zinc-finger motifs that are located in the predominantly disordered carboxyl terminus (Fig. 2C). BLM and TopIIIα together disentangle complex DNA structures, such as the double Holliday junction (dHJ) (Wu and Hickson 2003), a key intermediate in homologous recombination–based DNA repair (Yin et al. 2005; Bizard and Hickson 2014). In higher eukaryotes, the complex is augmented by the RecQ-mediated instability (RMI) 1 (Meetei et al. 2003; Yin et al. 2005) and 2 (Singh et al. 2008; Xu et al. 2008) proteins, forming direct physical interactions with both BLM (Raynard et al. 2006) and TopIIIα (Raynard et al. 2006; Bocquet et al. 2014). RMI1 and RMI2 are both OB-fold-containing proteins (Fig. 2D,E), with no inherent enzymatic activity. However, importantly, RMI1 stimulates the dHJ dissolution by BLM and TopIIIα (Raynard et al. 2006; Wu et al. 2006), whereas RMI2 has a very modest effect on this activity (Singh et al. 2008; Xu et al. 2008). RMI1 and RMI2 form a complex (Hoadley et al. 2010; Wang et al. 2010a) that is required to stabilize TopIIIα. As a result, they form a constitutive heterotrimer (termed the “TRR complex”) in vivo.

BLM has been used as a key marker of UFBs in many studies (Chan and Hickson 2009; Chan et al. 2009; Vinciguerra et al. 2010; Ke et al. 2011; Lukas et al. 2011; Barefield and Karlseder 2012; Broderick et al. 2015; Hengeveld et al. 2015). Considering that the BTRR complex has evolved to disentangle complex DNA structures, it is conceivable that this complex is responsible for UFB processing. This is supported by the observation that BS cells, and cells depleted of BLM by short interfering RNAs (siRNAs), display increased levels of all types of UFBs, and that these UFBs often persist into late telophase in these cells (Chan et al. 2007; Barefield and Karlseder 2012).

The recruitment of the BTRR complex to UFBs depends on PICH, and they always seem to coat the same stretch of DNA (Chan et al. 2007). This localization is somewhat curious, considering the fact that PICH binds exclusively to dsDNA (Biebricher et al. 2013), whereas the BTRR prefers ssDNA. The observation that BLM interacts with the carboxyl terminus of PICH suggests that PICH recruits the BTRR complex via direct interactions (Ke et al. 2011).

RPA

Replication protein A is an essential ssDNA binding protein required for most DNA transactions (Wold 1997; Zou et al. 2006). It is composed of three subunits (Fig. 2F) and interacts with the BTRR complex, both functionally and directly via BLM (Brosh et al. 2000; Meetei et al. 2003; Doherty et al. 2005) and RMI1 (Xue et al. 2013). RPA is detectable on a subset of UFBs in anaphase in response to DNA replication stress induced by aphidicolin (Chan and Hickson 2009; Burrell et al. 2013), indicating the presence of ssDNA on some CFS-UFBs (Chan et al. 2009). BLM and RPA show a nonoverlapping pattern of localization to UFBs, which suggests that the recruitment of BLM to PICH-coated double-stranded UFBs is independent of its interaction with RPA (Porter and Farr 2004; Chan and Hickson 2009). Interestingly, the appearance of RPA-coated UFBs has been shown to be BLM-dependent, implicating BLM in unwinding some structure to create ssDNA (Hengeveld et al. 2015).

Topoisomerase IIα

TopIIα is the enzyme responsible for the majority of decatenation of chromosomes in early mitosis (Porter and Farr 2004). TopIIα is a homodimeric Type IIA topoisomerase (Fig. 2G) that catalyzes the passage of one piece of dsDNA through another in an ATP-dependent manner (Schoeffler and Berger 2008). Even though TopIIα does not seem to directly interact with PICH, it is present at PICH-coated UFBs, and PICH is able to stimulate decatenation by TopIIα in vivo and in vitro (Nielsen et al. 2015). It has been suggested that TopIIα is recruited to a subset of UFBs via direct interaction with TOPBP1 (Broderick et al. 2015).

Other UFB-Associated Proteins

RIF1, TOPBP1, and FANCM were also reported to coat some UFBs (Meetei et al. 2003; Deans and West 2009; Xu et al. 2010; Hoadley et al. 2012; Wang et al. 2013; Blackford et al. 2015). We will not discuss these factors further here, but instead refer readers to relevant publications (German et al. 2007; Vinciguerra et al. 2010; Broderick et al. 2015; Hengeveld et al. 2015; Pedersen et al. 2015).

MODELING UFBs IN A TEST TUBE

Mechanistic insight into the mode of UFB resolution is lacking. To gain a comprehensive understanding of this process, our laboratory is using interdisciplinary approaches to reconstitute mitotic DNA decatenation in vitro. To achieve this, we combine ensemble biochemistry on model DNA substrates, with single-molecule optical tweezers coupled to fluorescence microscopy (Heller et al. 2014).

Modeling UFBs Using Ensemble Biochemistry

CFS and telomeres are both difficult-to-replicate regions, and both CFS-UFBs and T-UFBs are induced by replication stress (Chan and Hickson 2009; Barefield and Karlseder 2012). Therefore, it is thought that these UFBs are composed of underreplicated DNA (Fig. 3A). To study such a UFB in vitro, we created a substrate termed a “late replication intermediate” (LRI), which comprises two interlinked DNA circles mimicking two converging replication forks (A Sarlós, A Biebricher, and AH Bizard, unpubl.). We hypothesized that an LRI would be an ideal substrate for the BTRR complex. Indeed, a similar substrate was shown previously to be processed by the Escherichia coli homologs of the BTRR complex (Suski and Marians 2008).
As discussed above, TopII\(\alpha\) inhibition dramatically increases the number of C-UFBs (Chan et al. 2009). R-UFBs are also induced by ICRF-193 and are thought to arise because of the late condensation of the rDNA locus hindering decatenation (Nielsen and Hickson 2016). Considering that TopII\(\alpha\) is a dsDNA-specific enzyme, this implies that the majority of C-UFBs and R-UFBs are completely double-stranded catenanes (Fig. 3B). To model this, we used a single-catenane substrate comprised of two interlinked dsDNA circles (Stark et al. 1989; Nielsen et al. 2015). Considering that the BTRR complex prefers ss/dsDNA junctions, it is not clear what the function of BTRR might be on C-UFBs (Chan et al. 2007). Because studies involving the yeast and \textit{E. coli} homologs of BTRR reported some dsDNA catenation activity (Harmon et al. 2003; Cejka et al. 2012), it is conceivable that the BTRR can decatenate ds-UFBs if TopII\(\alpha\) is prohibited.

Most of the enzymes implicated in UFB processing are also involved in DNA repair. Therefore, it cannot be excluded that some UFBs are composed of HR intermediates, such as a dHJ or a D-loop (Fig. 3C,D). Such DNA structures would be expected to be “dissolved” efficiently by the BTRR complex (Wu and Hickson 2003; Bachrati et al. 2006).

**Modeling of UFBs in Single-Molecule Experiments**

Single-molecule techniques can provide information on enzymatic mechanisms that would be inaccessible by ensemble methods because of the averaging of the activity of thousands of molecules at the same time (Neuman and Nagy 2008). One of the most widely used single-molecule techniques is optical tweezers (Fig. 4A). Using optical tweezers, a single piece of biotinylated DNA can be stretched between two streptavidin-coated beads (gray) entrapped by laser beams (orange). The exonuclease activity of T7 DNA polymerase (blue) is shown creating ssDNA. A series of DNA force-extension curves, ranging from full-length dsDNA (black) to full-length ssDNA (blue). The orange, green, and cyan curves represent increasing incubation time with T7 DNA polymerase (as shown in the inset), which results in increasing lengths of ssDNA tracts.

![Figure 3. Hypothetical DNA structures that could be present at UFBs. (A) A late replication intermediate, (B) a complete double-stranded catenane, (C) a double Holliday junction, and (D) a D-loop.](image)

![Figure 4. Scheme of optical tweezers experiments. (A) A biotin-labeled dsDNA molecule stretched between two streptavidin-coated beads (gray) entrapped by laser beams (orange). The exonuclease activity of T7 DNA polymerase (blue) is shown creating ssDNA. (B) A series of DNA force-extension curves, ranging from full-length dsDNA (black) to full-length ssDNA (blue). The orange, green, and cyan curves represent increasing incubation time with T7 DNA polymerase (as shown in the inset), which results in increasing lengths of ssDNA tracts. (C) Outline of an experimental scheme for use of a flow cell with five channels. First beads are trapped, and then a dsDNA molecule is captured; this can subsequently be converted into an ss/dsDNA hybrid. This is followed by incubation with fluorescently labeled proteins, such as GFP-PICH (green) or RFP-RPA (red). (D,E) Schemes (D) and fluorescence snapshots (E) of a GFP-PICH-coated dsDNA molecule, a RFP-RPA/GFP-PICH-coated ss/dsDNA hybrid molecule, and a RFP-RPA-coated ssDNA stretch. Note that the extension of the DNA molecule is increasing by the introduction of ssDNA, which has a longer contour length than dsDNA.](image)
be tethered between streptavidin-coated polystyrene microspheres and manipulated in the flow cell of a fluorescence microscope (Heller et al. 2014). The effect of tension applied to a model UFB can thus be addressed, and fluorescently tagged proteins can be directly visualized on the stretched DNA.

The simplest UFB model one can imagine is a piece of dsDNA stretched between two beads. Indeed, we have showed using this approach that PICH is a DNA “tension sensor” that binds to stretched dsDNA with high affinity and translocates along the DNA in an ATP-dependent manner (Biebricher et al. 2013). However, as discussed above, presumably not all UFBs are completely double-stranded, and there are also multiple players involved in their processing. It is possible to generate a section of ssDNA with optical tweezers by stretching a dsDNA molecule to forces beyond 65–70 pN, which induces base pair melting. Occasionally, in the presence of a nick (a break in one of the strands), the melted strand dissociates, yielding a permanent stretch of ssDNA (Candelli et al. 2013). A more controlled way to generate a ssDNA/dsDNA hybrid is by using the exonuclease activity of the T7 DNA polymerase induced by putting the DNA under tension (Wuite et al. 2000; Hoekstra et al. 2017). Because reducing the tension can stop the exonuclease activity, a single DNA molecule can be generated containing the desired amount of ssDNA (Fig. 4B). The use of a multichannel flow cell system allows combination of different channels, containing various buffers and proteins, and the same piece of DNA can be freely manipulated between them with minimal contamination (Fig. 4C; Heller et al. 2014). UFBs can be modeled by incubating the DNA with different fluorescently labeled UFB-binding factors. Examples of a full-length dsDNA, a mixed ss/dsDNA, and a full-length ssDNA molecule coated with either GFP-PICH and/or strawberry RPA are shown in Figure 4D,E. By using other fluorescently labeled UFB-factors, such as BLM, the hierarchical recruitment of UFB-processing enzymes to PICH- versus RPA-coated UFBs could in principle be analyzed.

As described above, UFBs are likely to comprise interlinked dsDNA catenanes, and therefore a single piece of DNA, even if it is a mixture of ss/dsDNA, is not suitable for addressing details of the decatenation process. Braiding two DNA molecules together using quadruple-trap optical tweezers would provide a way to achieve this (Brouwer et al. 2017). Furthermore, implementation of confocal microscopy, or recent advances such as super-resolution imaging, allows the monitoring of real-time dynamics of individual molecules on model UFBs even in high fluorescent protein background (Heller et al. 2013).

WHAT HAPPENS IF THE CONVENTIONAL DNA DECATENATION PATHWAYS FAIL?

Cells seem to wait until the last moment to achieve correct segregation and can delay the completion of anaphase/telophase to accomplish this. This is reflected in BS cells, where UFBs persist even in very late telophase, spanning several microns between the two forming daughter cells (Chan et al. 2007). However, in some cases, when TopIIα or the BTRR complex is overwhelmed or inhibited, endonucleases might ensure that the DNA be cleaved in an apparently less controlled way, but nevertheless in a manner that avoids rupture of the bridge during abscission (Sarbajna et al. 2014; Maciejowski et al. 2015). Unresolved UFBs can also cause cytokinesis delay or the abandonment of cytokinesis, resulting in binucleation (Vinciguerra et al. 2010; Wang et al. 2010b; Germann et al. 2014; Nielsen et al. 2015).

Oncogene-induced replication stress generates UFBs, which is exacerbated in cancer cells (Burrell et al. 2013). Unresolved CFS–UFBs induced by replication stress can lead to the formation of 53BP1 nuclear bodies in the next G1 phase (Harrigan et al. 2011; Lukas et al. 2011). Interfering with CFS processing mechanisms also induces chromosome missegregation and aneuploidy (Naim and Rosselli 2009; Ying et al. 2013; Minocherhomji et al. 2015). In line with this, depletion of UFB-associated enzymes such as BLM also generates micronuclei (Rosin and German 1985). Micronuclei are a source of chromothripsis, one of the drivers of genomic rearrangements in cancer (Crasta et al. 2012; Zhang et al. 2015).

CONCLUSION

Since their discovery 10 years ago, UFBs have become one of the key markers of genomic instability. It has also become more obvious that cells have to cope with DNA entanglements during mitosis much more frequently than was thought previously. As UFBs arise systematically from specific loci (centromeres, fragile sites, telomeres, and rDNA), specialized recognition and processing machineries have evolved to maintain the stability of these important regions. The significance of understanding fundamental mechanisms ensuring correct chromosome segregation cannot be overestimated, especially in the context of cancer. All essential cellular processes can only be understood properly by combining in vivo observations with in vitro model building studies. In the field of mitosis, comprehensive studies where key aspects of mitosis are reconstituted properly by combining in vivo observations with in vitro model building studies. In the field of mitosis, comprehensive studies where key aspects of mitosis are reconstituted are still in a very early phase. Clearly, interdisciplinary studies are required to construct a mechanistic model of chromosome segregation. In this review, we have highlighted how the combination of biochemical and single-molecule modeling of DNA structures is a useful tool to study UFBs, impediments of faithful segregation.

ACKNOWLEDGMENTS

We thank all members of the Hickson laboratory for helpful discussions and Hocine W. Mankouri and Anna H. Bizard for valuable comments on the manuscript. Work in the authors’ laboratory is supported by the Danish National Research Foundation (DNRF115), the European Research Council, the Nordea Foundation, and a Future and Emerging Technologies grant from the European Union H2020 fund.
REFERENCES

Al Mamun M, Albergante L, Moreno A, Carrington JT, Blow JJ, Newman TJ. 2016. Inevitability and containment of replication errors for euchromatin genome lengths spanning megabase to gigabase. *Proc Natl Acad Sci* 113: E5765–E5774.

Arnoult N, Karlseder J. 2015. Complex interactions between the DNA-damage response and mammalian telomeres. *Nat Struct Mol Biol* 22: 859–866.

Bachurin C, Bortolussi G, Hickson ID. 2006. Mobile D-loops are a preferred substrate for the Bloom’s syndrome helicase. *Nucleic Acids Res* 34: 2269–2279.

Barefield C, Karlseder J. 2012. The BLM helicase contributes to telomere maintenance through processing of late-replicating intermediate structures. *Nucleic Acids Res* 40: 7358–7367.

Baumann C, Körner R, Hofmann K, Nigg EA. 2007. PICH, a centromere-associated SNF2 family ATPase, is regulated by Plk1 and required for the spindle checkpoint. *Cell* 128: 101–114.

Baxter J, Aragón L. 2012. A model for chromosome condensation based on the interplay between condensin and topoisomerase II. *Trends Genet* 28: 110–117.

Biebricher A, Hirano S, Enzlin JH, Wiedensch N, Streicher WW, Huttner D, Wang LHC, Nigg EA, Owen-Hughes T, Liu Y, et al. 2013. Bloom syndrome DNA translocase adapted for processing anaphase bridge DNA. *Mol Cell* 51: 691–701.

Bizard AH, Hickson ID. 2014. The dissolution of double Holliday junctions. *Curr Opin Genet Dev* 26: 105–111.

Blackford AN, Nieminuszczy J, Blackford AN, Niedzwiedz W. 2015. Top-Q helicases in DNA metabolism but is dispensable for preventing DNA degradation. *Mol Cell* 57: 1133–1141.

Bocquet N, Bizard AH, Abdulrahman W, Larsen NB, Fatty M, Cavadini S, Bunker RD, Kowalczykowski SC, Cejka P, Hickson ID, et al. 2014. Structural and mechanistic insight into Holliday-junction dissolution by Topoisomerase IIIz and RMI1. *Nat Struct Mol Biol* 21: 261–268.

Broderick R, Nieminuszczy J, Blackford AN, Wncyzna A, Niedzwiedz W. 2015. TopBP1 recruits TOP2A to ultra-fine anaphase bridges to aid in their resolution. *Nat Commun* 6: 6572.

Brosh RM, Li JL, Kenny MK, Karow JK, Cooper MP, Kurosky AM, Brosh RM. 2005. Physical and functional interactions of the replication protein A interaction domain of the Werner and Bloom syndrome helicases. *J Biol Chem* 280: 29494–29505.

Deksiński J, de Lange T. 2014. The role of double-strand break repair pathways at functional and dysfunctional telomeres. *Cold Spring Harb Perspect Biol* 6: a016576.

Deksiński J, Wu YJ, de Lange T, Zhuang XW. 2013. Super-resolution fluorescence imaging of telomeres reveals TRF2-dependent T-loop formation. *Cell* 155: 345–356.

Drosopoulos WC, Kociyatratkul ST, Schildkraut CL. 2015. BLM helicase facilitates telomere replication during leading strand synthesis of telomeres. *J Cell Biol* 210: 191–208.

Durkin SG, Glover TW. 2007. Chromosome fragile sites. *Annu Rev Genet* 41: 169–192.

Ellis NA, Lennon DJ, Protycheva M, Alhadef B, Henderson EE, Germain J. 1995. Somatic intragenic recombination within the mutated locus BLM can correct the high sister-chromatid exchange frequency of Bloom syndrome cells. *Am J Hum Genet* 57: 1019–1027.

Debatisse M, Le Tallec B, Letessier A, Dutrillaux B, Brison O. 2012. Common fragile sites: Mechanisms of instability revisited. *Trends Genet* 28: 22–32.

Doherty KM, Sommers JA, Gray MD, Lee JW, von Kobbe C, Thorin NH, Kureeckatil RP, Kenny MK, Brosh RM. 2005. Physical and functional interactions of the replication protein A interaction domain of the Werner and Bloom syndrome helicases. *J Biol Chem* 280: 29494–29505.

Debatisse M, Le Tallec B, Letessier A, Dutrillaux B, Brison O. 2012. Common fragile sites: Mechanisms of instability revisited. *Trends Genet* 28: 22–32.

Doherty KM, Sommers JA, Gray MD, Lee JW, von Kobbe C, Thorin NH, Kureeckatil RP, Kenny MK, Brosh RM. 2005. Physical and functional interactions of the replication protein A interaction domain of the Werner and Bloom syndrome helicases. *J Biol Chem* 280: 29494–29505.

Debatisse M, Le Tallec B, Letessier A, Dutrillaux B, Brison O. 2012. Common fragile sites: Mechanisms of instability revisited. *Trends Genet* 28: 22–32.

Doherty KM, Sommers JA, Gray MD, Lee JW, von Kobbe C, Thorin NH, Kureeckatil RP, Kenny MK, Brosh RM. 2005. Physical and functional interactions of the replication protein A interaction domain of the Werner and Bloom syndrome helicases. *J Biol Chem* 280: 29494–29505.

Debatisse M, Le Tallec B, Letessier A, Dutrillaux B, Brison O. 2012. Common fragile sites: Mechanisms of instability revisited. *Trends Genet* 28: 22–32.

Doherty KM, Sommers JA, Gray MD, Lee JW, von Kobbe C, Thorin NH, Kureeckatil RP, Kenny MK, Brosh RM. 2005. Physical and functional interactions of the replication protein A interaction domain of the Werner and Bloom syndrome helicases. *J Biol Chem* 280: 29494–29505.

Debatisse M, Le Tallec B, Letessier A, Dutrillaux B, Brison O. 2012. Common fragile sites: Mechanisms of instability revisited. *Trends Genet* 28: 22–32.

Doherty KM, Sommers JA, Gray MD, Lee JW, von Kobbe C, Thorin NH, Kureeckatil RP, Kenny MK, Brosh RM. 2005. Physical and functional interactions of the replication protein A interaction domain of the Werner and Bloom syndrome helicases. *J Biol Chem* 280: 29494–29505.

Debatisse M, Le Tallec B, Letessier A, Dutrillaux B, Brison O. 2012. Common fragile sites: Mechanisms of instability revisited. *Trends Genet* 28: 22–32.

Doherty KM, Sommers JA, Gray MD, Lee JW, von Kobbe C, Thorin NH, Kureeckatil RP, Kenny MK, Brosh RM. 2005. Physical and functional interactions of the replication protein A interaction domain of the Werner and Bloom syndrome helicases. *J Biol Chem* 280: 29494–29505.

Debatisse M, Le Tallec B, Letessier A, Dutrillaux B, Brison O. 2012. Common fragile sites: Mechanisms of instability revisited. *Trends Genet* 28: 22–32.

Doherty KM, Sommers JA, Gray MD, Lee JW, von Kobbe C, Thorin NH, Kureeckatil RP, Kenny MK, Brosh RM. 2005. Physical and functional interactions of the replication protein A interaction domain of the Werner and Bloom syndrome helicases. *J Biol Chem* 280: 29494–29505.

Debatisse M, Le Tallec B, Letessier A, Dutrillaux B, Brison O. 2012. Common fragile sites: Mechanisms of instability revisited. *Trends Genet* 28: 22–32.

Doherty KM, Sommers JA, Gray MD, Lee JW, von Kobbe C, Thorin NH, Kureeckatil RP, Kenny MK, Brosh RM. 2005. Physical and functional interactions of the replication protein A interaction domain of the Werner and Bloom syndrome helicases. *J Biol Chem* 280: 29494–29505.
Harmon FG, Brockman JP, Kowalczykowski SC. 2003. RecQ helicase stimulates both DNA catenation and changes in DNA topology by topoisomerase III. *J Biol Chem* 278: 42668–42678.

Harrigan JA, Belotserkovskaya R, Coates J, Dimitrova DS, Polo SE, Bradshaw CR, Fraser P, Jackson SP. 2011. Replication stress induces 53BP1-containing OPT domains in G1 cells. *J Cell Biol* 193: 97–108.

Heller I, Sitters G, Brockmans OD, Farge G, Mengers C, Wende W, Hell SW, Pederman EJG, Wuite GJL. 2013. STED nanoscopy combined with optical tweezers reveals protein dynamics on densely covered DNA. *Nature Methods* 10: 910–916.

Heller I, Hoekstra TP, King GA, Pederman EJ, Wuite GJ. 2014. Optical tweezers analysis of DNA–protein complexes. *Chem Rev* 114: 3087–3119.

Hengeveld RC, de Boer HR, Schoono MN, de Vries EG, Lens SM, van Vugt MA. 2015. Rif1 is required for resolution of ultrafine DNA bridges in anaphase to ensure genomic stability. *Dev Cell* 34: 466–474.

Higa M, Fujita M, Yoshida K. 2017. DNA replication origins and fork progression at mammalian telomeres. *Genes (Basel)* 8: E112.

Hirano T. 2015. Chromosome dynamics during mitosis. *Cold Spring Harb Perspect Biol* 7: a021592.

Hoadley KA, Xu DY, Xue YT, Satyshur KA, Wang WD, Keck JL. 2010. Structure and cellular roles of the RMI core complex from the Bloom syndrome dissolvasome. *Nature Struct Mol Biol* 17: 1149–1158.

Hoadley KA, Xue YT, Ling C, Takata M, Wang WD, Keck JL. 2012. Defining the molecular interface that connects the Fanconi anemia protein FANCM to the Bloom syndrome dissolvasome. *Proc Natl Acad Sci* 109: 4437–4442.

Hoekstra TP, Depken M, Lin SN, Cabanas-Danès J, Gross P, Dame RT, Pederman EJ, Wuite GJ. 2017. Switching between exonucleaseysis and replication by T7 DNA polymerase ensures high fidelity. *Biophys J* 112: 575–583.

Hoffelder DR, Luo L, Burke NA, Watkins SC, Gollin SM, Saunders WS. 2004. Resolution of anaphase bridges in cancer cells. *Chromosoma* 112: 359–397.

Karow JK, Chakravarty RK, Hickson ID. 1997. The Bloom’s syndrome gene product is a 3·5′ DNA helicase. *J Biol Chem* 272: 30611–30614.

Karow JK, Constantinou A, Li JL, West SC, Hickson ID. 2000. The Bloom’s syndrome gene product promotes branch migration of Holliday junctions. *Proc Natl Acad Sci* 97: 6504–6508.

Kaurich J, Cubizolles F, Nigg EA. 2012. On the regulation, function, and localization of the DNA-dependent ATPase PICH. *Chromosoma* 121: 395–408.

Ke Y, Huh JW, Warrington R, Li B, Wu N, Leng M, Zhang J, Ball HL, Li B, Yu H. 2011. PICH and BLM limit histone association with anaphase centromeric DNA threads and promote their resolution. *EMBO J* 30: 3309–3321.

Kijitima TS, Sakuno T, Ishiguro K, Jimura S, Natsune T, Kawashima S, Watanabe Y. 2006. Shugoshin collaborates with protein phosphatase 2A to protect cohesin. *Nature* 441: 54–52.

Kunnsawa Y, Yu–Lee LY. 2010. PICH and cotargeted Plk1 coordinate maintain prometaphase chromosome arm architecture. *Mol Biol Cell* 21: 1188–1199.

Leng M, Besusso D, Jung SY, Wang Y, Qin J. 2008. Targeting Plk1 to chromosome arms and regulating chromosome compaction by the PICH ATPase. *Cell Cycle* 7: 1480–1489.

Lucas C, Savic V, Bekker-Jensen S, Doil C, Neumann B, Pedersen RS, Grufte M, Chan KL, Hickson ID, Bartek J, et al. 2011. 53BP1 nuclear bodies form around DNA lesions generated by mitotic transmission of chromosomes under replication stress. *Nature Cell Biol* 13: 243–253.

Maciejewski JI, Li YL, Bosco N, Campbell PJ, de Lange T. 2015. Chromatin scope and kategias induced by telomere crisis. *Cell* 163: 1641–1654.

Martinez P, Blasco MA. 2015. Replicating through telomeres: A means to an end. *Trends Biochem Sci* 40: 504–515.

Martinez P, Thanasoula M, Munoz P, Liao CY, Tejera A, McNees C, Flores JM, Fernandez-Capetillo O, Tarsounas M, Blasco MA. 2009. Increased telomere fragility and fusions resulting from TRF1 deficiency lead to degenerative pathologies and increased cancer in mice. *Genes Dev* 23: 2060–2075.

McClelland B. 1938. The production of homozygous deficient tissues with mutant characteristics by means of the aberrant mitotic behavior of ring-shaped chromosomes. *Genetics* 23: 315–376.

McClelland B. 1942. The fusion of broken ends of chromosomes following nuclear fusion. *Proc Natl Acad Sci* 28: 458–463.

Meetei AR, Sechi S, Wallisch M, Yang DF, Young MK, Joenje H, Hoatlin ME, Wang WD. 2003. A multiprotein nuclear complex connects Fanconi anemia and Bloom syndrome. *Mol Cell Biol* 23: 3417–3426.

Menochromijii S, Ying SM, BjERRragna VA, Bursomanno S, Alelaimate A, Wu W, Mankouri HW, Shen HH, Liu Y, Hickson ID. 2015. Replication stress activates DNA repair synthesis in mitosis. *Nature* 528: 286–290.

Moreno A, Carrington JT, Albergante L, Al Mamun M, Haagenesen EJ, Komseli ES, Gorgoulis VG, Newman TJ, Blow JJ. 2016. Unreplicated DNA remaining from unperturbed S phases passes through mitosis for resolution in daughter cells. *Proc Natl Acad Sci* 113: E5757–E5764.

Naim V, Rosselli F. 2009. The FANC pathway and BLM collaborate during mitosis to prevent micro-nucleation and chromosome abnormalities. *Nat Cell Biol* 11: 761–768.

Nasmyth K. 2011. Cohesion: A catenase with separate entry and exit gates? *Nat Cell Biol* 13: 1170–1177.

Ner A, Huang HS, Lai T, Xu L. 2015. Elevated levels of TRF2 induce telomeric ultrafine anaphase bridges and rapid telomere deletions. *Nat Commun* 6: 10132.

Neuman KC, Nagy A. 2008. Single-molecule force spectroscopy: Optical tweezers, magnetic tweezers and atomic force microscopy. *Nat Methods* 5: 491–505.

Nielsen CF, Hickson ID. 2016. PICH promotes mitotic chromosome segregation: Identification of a novel role in rDNA disjunction. *Cell Cycle* 15: 2704–2711.

Nielsen CF, Huttner D, Bizert AH, Hirano S, Li TN, Palmai-Pallag T, Bjerrregaard VA, Liu Y, Nigg EA, Wang LHC, et al. 2015. PICH promotes sister chromatid disjunction and cooperates with topoisomerase II in mitosis. *Nat Commun* 6: 8962.

Palm W, de Lange T. 2008. How shelterin protects mammalian telomeres. *Annu Rev Genet* 42: 301–334.

Pedersen RT, Kruse T, Nilsson J, Oestergaard VH, Lisby M. 2015. TopBP1 is required at mitosis to reduce transmission of DNA damage to G1 daughter cells. *J Cell Biol* 210: 565–582.

Piskadlo E, Tavares A, Oliveira KA. 2017. Metaphase chromosome structure is dynamically maintained by condensin I-directed DNA (de)catenation. *Elife* 6: 1328.

Pitchai GP, Kaulich M, Bizert AH, Piskadlo E, Nigg EA, Wang WD, Wong CC, Beyermann M, Cirillo ML, Zhou D, Blazquez ME, et al. 2013. A novel TPR-BEND domain interaction mediates PICH-BEND3 association. *Nucleic Acids Res* (in press).

Porter AC, Farr CJ. 2004. Topoisomerase II: Untangling its contribution at the centromere. *Chromosomes* 12: 569–583.

Raynard S, Bussen W, Sung P. 2006. A double Holliday junction dissolves comprising BLM, topoisomerase III, and BLAP75. *J Biol Chem* 281: 13861–13864.

Rosin MP, German J. 1985. Evidence for chromosome instability in vivo in Bloom syndrome: Increased numbers of micronuclei in exfoliated cells. *Hum Genet* 71: 187–191.

Rouzeau S, Cordelieres FP, Buhagiar-Labarchede G, Hurbain B, Onclerc-Delic R, Gembal S, Magnaghi-Jaulin L, Jaulin C, Amor-Gueret M. 2012. Bloom’s syndrome and PICH helicases cooperate with topoisomerase II in centromere disjunction before anaphase. *PLoS One* 7: e33905.

Sarbajna S, Davies D, West SC. 2014.Roles of SLX1-SLX4, MUS81-EME1, and GEN1 in avoiding genome instability and mitotic catastrophe. *Genes Dev* 28: 1124–1136.

Schoeffler AJ, Berger JM. 2008. DNA topoisomerases: Harnessing and constraining energy to govern chromosome topology. *Q Rev Biophys* 41: 41–101.

Schwartzman JA, Stasiak A. 2004. A topological view of the replication. *EMBO Rep* 5: 256–261.
Disentangling mitotic chromosomes

Sfeir A, Kosiyantrakul ST, Hockemeyer D, MacRae SL, Karhede J, Schildkraut CL, de Lange T. 2009. Mammalian telomeres resemble fragile sites and require TRF1 for efficient replication. Cell 138: 90–103.

Sims AE, Spiteri E, Sims RJ III, Arita AG, Lach FP, Landers T, Wang BD, Yong-Gonzalez V, Strunnikov AV. 2004. Cdc14p-FAER pathway controls segregation of nucleolus in S. cerevisiae by facilitating condensin targeting to rDNA chromatin in anaphase. Cell Cycle 3: 960–967.

Wang LHC, Schwarzenbraun T, Speicher MR, Nigg EA. 2008. Persistence of DNA threads in human anaphase cells suggests later onset of sister chromatid deconcatenation. Chromosoma 117: 123–135.

Wang F, Yang YT, Singh TR, Bussygina V, Guo R, Wan K, Wang WD, Sung P, Meetee AR, Lei M. 2010a. Crystal structures of RM11 and RM12, two OB-fold regulatory subunits of the BLM complex. Structure 18: 1159–1170.

Wang LHC, Mayer B, Stemmann O, Nigg EA. 2010b. Centromere DNA deconcatenation depends on cohesion removal and is required for mammalian cell division. J Cell Sci 123: 806–813.

Wang JD, Chen JI, Gong ZH. 2013. TopBP1 controls BLM protein level to maintain genome stability. Mol Cell 52: 667–678.

Wold MS. 1997. Replication protein A: A heterotrimeric, single-stranded DNA-binding protein required for eukaryotic DNA metabolism. Annu Rev Biochem 66: 61–92.

Wu L, Hickson ID. 2003. The Bloom’s syndrome helicase suppresses crossing over during homologous recombination. Nature 426: 870–874.

Wu L, Davies SL, North PS, Goulauouic H, Riou JF, Turley H, Gatter KC, Hickson ID. 2000. The Bloom’s syndrome gene product interacts with topoisomerase III. J Biol Chem 275: 9636–9644.

Wu L, Davies SL, Levitt NC, Hickson ID. 2001. Potential role for the BLM helicase in recombinational repair via a conserved interaction with RAD51. J Biol Chem 276: 19375–19381.

Wu L, Bachrati CZ, Xu C, Xu C, Yin J, Chang M, Wang W, Li L, Brown GW, Hickson ID. 2006. BLAP75-RMI1 promotes the BLM-dependent dissolution of homologous recombination intermediates. Proc Natl Acad Sci U S A 103: 4066–4071.

Wuite GJL, Smith SB, Young M, Keller D, Bustamante C. 2000. Single-molecule studies of the effect of template tension on T7 DNA polymerase activity. Nature 404: 103–106.

Xu DY, Guo R, Sobeck A, Bachrati CZ, Yang J, Enomoto T, Brown GW, Holatin ME, Hickson ID, Wang WD. 2008. RMI1, a new OB-fold complex essential for Bloom syndrome protein to maintain genome stability. Genes Dev 22: 2843–2855.

Xu D, Muniaindi P, Leo E, Yin J, Thangavel S, Shen X, Li M, Agama K, Guo R, Fox D III, et al. 2010. Rif1 provides a new DNA-binding interface for the Bloom syndrome complex to maintain normal replication. EMBO J 29: 3140–3155.

Xue XY, Raynard S, Busygina V, Singh AK, Sung P. 2013. Role of replication protein A in double Holliday junction dissolution mediated by the BLM-Topo III-RM11-RM12 protein complex. J Biol Chem 288: 14221–14227.

Yin JH, Sobeck A, Xu C, Meetee AR, Holatin M, Li L, Wang WD. 2005. BLAP75, an essential component of Bloom’s syndrome protein complexes that maintain genome integrity. EMBO J 24: 1465–1476.

Yin SM, Minocherhomjii S, Chan KL, Palmai-Pallag T, Chu WK, Wass T, Mankouri HW, Liu Y, Hickson ID. 2013. MUS81 promotes common fragile site expression. Nat Cell Biol 15: 1001–1007.

Zhang CZ, Spektor A, Cornils H, Francis JM, Jackson EK, Liu SW, Meyerson M, Pellman D. 2015. Chromothripsis from DNA damage in micronuclei. Nature 522: 179–184.

Zou Y, Liu YY, Wu XM, Shell SM. 2006. Functions of human replication protein A (RPA): From DNA replication to DNA damage and stress responses. J Cell Physiol 208: 267–273.