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Acute inactivation of the replicative helicase in human cells triggers MCM8–9-dependent DNA synthesis

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DNA replication fork progression can be disrupted at difficult to replicate loci in the human genome, which has the potential to challenge chromosome integrity. To model the events stemming from replisome dissociation during DNA replication perturbation, we used a degron-based system for inducible proteolysis of a subunit of the replicative helicase. We show that MCM2-depleted cells activate a DNA damage response pathway and generate replication-associated DNA double-strand breaks (DSBs). Remarkably, these cells maintain some DNA synthesis in the absence of MCM2, and this requires the MCM8–9 complex, a paralog of the MCM2–7 replicative helicase. We show that MCM8–9 functions in a homologous recombination-based pathway downstream from RAD51, which is promoted by DSB induction. This RAD51/MCM8–9 axis is distinct from the recently described RAD52-dependent DNA synthesis pathway that operates in early mitosis at common fragile sites. We propose that stalled replication forks can be restarted in S phase via homologous recombination using MCM8–9 as an alternative replicative helicase.

[Keywords: DNA replication; homologous recombination; fork restart; MCM proteins; genome maintenance; conditional degron]

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a recent report showed that human cells experience at least one replicon failure per S phase, where two forks are irreversibly arrested without there being an intervening origin to rescue them [Moreno et al. 2016]. This feature of S phase is particularly pertinent to regions of the genome that are origin-poor, meaning that some forks must travel over a considerable distance without breakdown. Therefore, cells must have backup systems to deal with situations in which two or more forks are irreversibly stalled; otherwise, the completion of DNA replication would be compromised.

In prokaryotes, fork restart at nonorigin sites plays a crucial role in the protection against irreversible fork stalling. This is particularly required in these organisms because only a single genomic origin generates a pair of forks that replicate the entire genome before converging at a defined termination region. The main fork restart pathway is dependent on the homologous recombination (HR) machinery and is initiated at a DNA double-strand break (DSB) at the stalled fork [Cox et al. 2000; Michel et al. 2004]. For example, replication forks in *Escherichia coli* can be stalled by inactivation of the replicative DnaB helicase [Michel et al. 1997]. This generates a one-ended DSB at the stalled fork that triggers RecBCD- and RuvABC-dependent recombination between sister chromatids [Seigneur et al. 2000]. Following RecA-mediated displacement loop (D-loop) formation and the action of the PriA–PriB–DnaT “primosome” complex, DnaB is reloaded for reassembly of the replisome [Seigneur et al. 1998; Heller and Marians 2006]. Thus, *E. coli* has an efficient system for reassembly of the replisome via HR triggered by a one-ended DSB.

In eukaryotes, the form of HR repair used to deal with one-ended DSBs is known as break-induced replication (BIR) and plays an important role in both telomere maintenance and replication fork restart [McEachern and Haber 2006; Llorente et al. 2008; Verma and Greenberg 2016]. BIR has been characterized in budding yeast through the analysis of interchromosomal HR induced by one-ended DSB [Morrow et al. 1997; Bosco and Haber 1998]. DNA synthesis during BIR is carried out by DNA polymerase δ (Pol δ), which is coupled to Pif1 helicase-dependent migration of a DNA D-loop structure [Saini et al. 2013; Wilson et al. 2013]. The noncatalytic Pol32 subunit of Pol δ is essential for BIR but not bulk DNA replication [Lyeard et al. 2007]. In mammalian cells, BIR is poorly characterized, partly because of a lack of defined assays. However, it has been shown that the POL32 subunit (Pol32 homolog) of Pol δ is also required for BIR and alternative telomere maintenance in human cells [Costantino et al. 2014; Dilley et al. 2016]. However, in contrast to yeast, mammalian POL32 is essential for cell viability [Murga et al. 2016]. Importantly, the mechanism of BIR in mammalian cells is still unclear, and it remains to be confirmed that it plays a key role in rescuing irreversibly stalled replication forks.

Replication forks in eukaryotes are driven by the hexameric MCM2–7 helicase, which forms the so-called CMG replicative holohelicase along with CDC45 and the GINS complex [Ilves et al. 2010]. MCM2–7 activity is tightly controlled during the cell cycle [Blow and Dutta 2005; Masai et al. 2010]. The loading of MCM2–7 at origins is temporally separated from helicase activation, with the former occurring in late M and G1 phases, and the latter occurring only in S phase. Importantly, the loading of additional MCM2–7 is suppressed in S phase, ensuring that DNA replication takes place only once per cell cycle. This implies that, unlike in *E. coli*, the replisome cannot be reassembled at a stalled fork by the reloading of the MCM2–7 helicase.

Most eukaryotic species, with the exception of yeasts and nematodes, have additional MCM family proteins, known as MCM8 and MCM9 [Liu et al. 2009]. We and others reported that MCM8 and MCM9 form a distinct complex that is involved in HR repair [Lutzmann et al. 2012; Nishimura et al. 2012; Park et al. 2013]. MCM9 was also shown recently to interact with mismatch repair (MMR) proteins and work with MCM8 as a helicase during MMR [Traver et al. 2015]. MCM8 and MCM9 are both required for gametogenesis and tumor suppression in mice [Hartford et al. 2011; Lutzmann et al. 2012], and mutations in the human MCM8 or MCM9 genes are associated with premature onset of menopause [He et al. 2009; Wood-Trager et al. 2014]. Many lines of evidence point to a role for the MCM8–9 complex as a helicase in DNA repair, particularly in HR repair. However, there are conflicting views as to whether MCM8–9 is required for an early process (e.g., DNA end resection) or a later process in HR [Lutzmann et al. 2012; Nishimura et al. 2012; Lee et al. 2015].

In order to define the processes required for rescue of stalled forks in human cells and the possible role of MCM8–9 in these processes, we generated a human cell line in which the MCM2–7 helicase could be inactivated in a controlled manner. For this purpose, we used auxin-inducible degron (AID) technology, whereby a degron-tagged protein can be rapidly degraded by adding the plant hormone auxin [Nishimura et al. 2009; Natsume et al. 2016]. This approach was adopted in order to create a situation in which the rescue of stalled forks by fork convergence is not possible [Supplemental Fig. S1B]. We demonstrate that, in response to MCM2 degradation, stalled forks are converted to DSBs that are rescued in a RAD51-dependent manner. Crucially, this rescue requires MCM8–9 to promote new DNA synthesis. Although this reaction is superficially similar to the recently described mitotic DNA synthesis [MiDAS] at fragile sites [Minocherhomji et al. 2015; Blouwnick et al. 2016], we show that MCM8–9-dependent DNA synthesis is distinct from MiDAS. We propose that MCM8–9 is required for HR-mediated fork restart and acts as an alternative replicative helicase to promote DNA synthesis.

Results

Construction of an MCM2 degron cell line for artificial fork stalling

To characterize how human cells deal with stalled replication forks, we took inspiration from studies in prokaryotes.
in which the replicative DnaB helicase had been conditionally inactivated using a temperature-sensitive mutation of DnaB [Michel et al. 1997; Seigneur et al. 1998, 2000]. In our case, we aimed to induce the rapid degradation of MCM2, a component of the replicative MCM2–7 helicase, by taking advantage of the AID technology [Fig. 1A; Nishimura et al. 2009; Natsume et al. 2016]. To achieve this, we tagged both alleles of the MCM2 gene with mini-AID (mAID) using CRISPR–Cas9 in the HCT116 human colorectal cancer line [Fig. 1B; Supplemental S2A,B; Cong et al. 2013; Mali et al. 2013]. Subsequently, we introduced an AFB2 gene (derived from Arabidopsis thaliana), which encodes a paralog of TIR1 (Supplemental Fig. S2C; Havens et al. 2012). In the resultant cells, the MCM2 fusion protein (MCM2-mAID) was degraded efficiently within 4 h of the addition of auxin (Fig. 1C), leading to an accumulation of cells in S phase, as expected [Supplemental Fig. S2D]. To analyze this more systematically, we synchronized cells in the G1 phase using lovastatin and then treated them with or without auxin before releasing them into S phase [Fig. 1D,E; Javanmoghadam-Kamrani and Keyomarsi 2008]. In control cells not exposed to auxin, DNA replication was generally completed within 19 h of release from G1 [Fig. 1E, control]. In sharp contrast, a profound defect in DNA replication was observed in cells treated with auxin [Fig. 1E, +auxin]. Importantly, this clear S-phase defect could not be seen using siRNA-mediated depletion methods because of the presence of residual MCM2 protein [Supplemental Fig. S2E,F].

**Fork stalling by induced proteolysis of MCM2 during S phase**

In order to induce fork stalling only after the cells had initiated S phase, we synchronized the MCM2-mAID cells in G1 phase as in Figure 1E but then only added auxin 13 h after release, when most of the cells were in early S phase [Fig. 2A,B]. In control cells not exposed to auxin, DNA replication was generally completed by the 19-h time

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**Figure 1.** Construction of MCM2-mAID cells for artificial fork stalling. **(A)** A schematic representation of auxin-mediated proteolysis of MCM2-mAID. An auxin-dependent F-box protein, AFB2 of A. thaliana (AtAFB2), forms an E3 ubiquitin ligase with the endogenous SCF components. In the presence of auxin, MCM2-mAID is targeted by AtAFB2 for polyubiquitylation and subsequent destruction by the proteasome. **(B)** Evidence that clones 1 and 2 express the MCM2-mAID protein. **(C)** Time course of proteolysis of MCM2-mAID. Asynchronously growing MCM2-mAID cells were treated with auxin [indole-3-acetic acid (IAA)] before harvesting at the indicated time points. Ponceau staining shows a loading control. **(D,E)** MCM2-mAID cells were arrested in G1 phase and released into S phase after MCM2-mAID depletion. In control cells, DMSO replaced auxin. MCM2-mAID proteins were detected by immunoblotting using the anti-mAID antibody (D), and DNA content was measured by flow cytometry (E).
point, and the cells then progressed through to mitosis (Fig. 2B, control). In contrast, the auxin-treated cells failed to carry out bulk DNA replication and eventually lost viability (Fig. 2B, +auxin; data not shown). The MCM2-mAID protein was undetectable by 2 h after auxin addition, suggesting that most replication forks would be inactivated at that point or soon afterward (Fig. 2A, +auxin 15 h). We also confirmed that the chromatin-bound fraction of MCM2-mAID, which is associated with forks and origins, was efficiently degraded [Supplemental Fig. S3A]. Intriguingly, we noted that the peak of DNA content drifted toward a 4C DNA content between the 15- and 21-h time points (Fig. 2B, +auxin). More detailed flow cytometric analysis of cells containing a more than 2C DNA content showed that the F1 and F2 populations decreased, while the F3 and F4 populations increased between 15 and 21 h (Fig. 2C). This alternative DNA synthesis that occurs without the MCM2-7 helicase is addressed below.

Because AID technology functions at the protein level, we took advantage of the ability to rapidly replenish the MCM2-depleted cells by removal of auxin from the medium. For this, we induced the rapid degradation of MCM2-mAID from 13 to 17 h after release from G1 phase, when the cells were in early S phase, and then removed auxin to allow MCM2-mAID re-expression (Supplemental Fig. S3B,C). Re-expression of MCM2-mAID was detectable 2 h after auxin removal (Supplemental Fig. S3B; 19 h). However, this failed to rescue the defective replication due to MCM2-mAID depletion (Supplemental Fig. S3C; left), consistent with the concept that the replication licensing system in eukaryotes prevents the replicative helicase from being reloaded to chromosomes in S phase [Blow and Dutta 2005]. This is in contrast to the system operating in bacteria for reloading of the DnaB helicase [Marians 2000].

DNA DSBs are induced following fork stalling

Stalled replication forks can be converted into DSBs [Petermann et al. 2010]. We analyzed whether DSBs were formed after MCM2-mAID depletion. For this, we initially looked at the 53BP1 protein, which forms nuclear foci upon DSB formation (Fig. 3A; Schultz et al. 2000; Anderson et al. 2001; Rappold et al. 2001). We observed that the auxin-treated cells accumulated 53BP1 foci in a manner similar to control cells treated with bleomycin, a known DSB-generating agent (Fig. 3B). To detect DSBs directly in the genomic DNA, we performed pulsed-field gel electrophoresis (PFGE) [Ray Chaudhuri et al. 2012]. This revealed that DSBs started to accumulate 4 h after the cells were treated with auxin (Fig. 3C, 17 h). We then analyzed the localization of RAD51 and γH2AX, which form nuclear foci at the sites of damaged DNA (Fig. 4A). We observed a significant enrichment of RAD51 and γH2AX foci in the MCM2-mAID-depleted cells (Fig. 4A, +auxin), indicating that the cells were accumulating DNA damage. Consistent with this, we detected activation of the CHK1 kinase in cells treated with auxin [Supplemental Fig. S4A]. Taken together, these results indicate that fork stalling induced by degradation of MCM2-mAID generates DNA DSBs.

The presence of γH2AX and 53BP1 nuclear foci after MCM2-mAID depletion is indicative of replication-associated DSBs [Figs. 3A,B, 4A]. However, we were intrigued by the colocalization of RAD51 with these DSBs,
suggestive of the activation of some form of HR at the stalled forks. We therefore analyzed whether the MCM8–9 complex might be required for RAD51 focus formation. To this end, we disrupted the MCM9 gene in the MCM2-mAID background (Supplemental Fig. S4B–D). The MCM9 knockout (MCM9-KO) cells did not show a significant growth defect under normal growth conditions, in line with our previous observation that MCM9 loss is not detrimental to the growth of chicken DT40 cells [Supplemental Fig. S4E; Nishimura et al. 2012]. Moreover, consistent with the observation that MCM8 and MCM9 function together in a complex, the formation of DNA damage-induced MCM8 foci was absent from the MCM9-KO cells [Supplemental Fig. S4F; Lutzmann et al. 2012; Park et al. 2013]. We then analyzed RAD51 focus formation in the MCM9-KO cells [Fig. 4B]. We observed that three independent clones of MCM9-KO cells formed RAD51 foci similarly to wild-type cells (MCM9-WT), indicating that MCM9 is not essential for RAD51 loading. Conversely, we obtained evidence that RAD51 is required for the loading of MCM8–9. In cells treated with the RAD51 inhibitor RI-1 [RAD51i] [Budke et al. 2012], we observed that MCM8 focus formation was significantly reduced [Fig. 4C].
MCM8–9 promotes DNA synthesis as a backup of DNA replication

As noted in Figure 2B, we observed that DNA synthesis continued to some extent for several hours after degradation of MCM2-mAID. Although this effect might reflect incomplete MCM2-mAID degradation, we considered the possibility that removal of this core component of the replisome might activate an alternative mechanism of DNA synthesis. We therefore investigated whether MCM8–9 might contribute to this MCM2–7-independent DNA synthesis. For this, we synchronized MCM9-WT and MCM9-KO cells in G1 and then released them into S phase as in Figure 2. In early S phase, auxin was added to induce MCM2-mAID depletion, and then time course samples were taken [Fig. 5A; Supplemental Fig. S5A]. The MCM9-WT and MCM9-KO cells showed very similar replication profiles in the absence of auxin, indicating that MCM9 is dispensable for normal bulk DNA replication [Fig. 5A, control]. In contrast, in the auxin-treated cells in which MCM2-mAID was degraded [Supplemental Fig. S5A, +auxin], the MCM9-KO cells showed a reduced level of DNA synthesis compared with the MCM9-WT cells between the 17- and 21-h time points [Fig. 5A, +auxin]. This result was confirmed by analysis of the percentage of MCM9-KO cells present in the late stages of S phase [F3 and F4] [Supplemental Fig. S5B,C] and was also observed in an independent MCM9-KO clone [data not shown]. To quantify this DNA synthesis defect in the MCM9-KO cells, we used bromodeoxyuridine [BrdU] to label nascent DNA 4 h after auxin addition. The population of cells labeled with BrdU was reduced in the MCM9-KO cells compared with the MCM9-WT cells [Fig. 5B, dotted circle].

To directly quantify ongoing DNA synthesis, we exposed cells sequentially to two thymidine analogs, iodo-deoxyuridine [IdU] and chlorodeoxyuridine [CldU], and then performed immunodetection of the incorporated DNA.
IdU and CldU on stretched DNA fibers (Fig. 5C). This revealed that the proportion of DNA fibers labeled with both IdU and CldU was reduced in the MCM9-KO cells (Fisher's exact test, \( P = 0.0023 \)). However, we still observed that ∼20% of the DNA fibers incorporated both IdU and CldU in the MCM9-KO cells. This suggests either that there is an MCM8–9-independent mechanism of DNA synthesis in the absence of MCM2-mAID or that degradation of MCM2-mAID was incomplete. We then analyzed the fork speed by measuring the length of CldU tracts (Fig. 5D). The fork speed after MCM2-mAID depletion was significantly slower than that of normal forks. This result supports the notion that MCM2-mAID depletion was efficient and that alternative DNA synthesis with MCM9 is qualitatively different from normal DNA synthesis. To define the location of DNA synthesis in the nucleus, we degraded MCM2-mAID using 4 h of auxin treatment and then labeled the MCM9-WT and MCM9-KO cells with ethynyldeoxyuridine (EdU) for 30 min. We observed that 17% of the large γH2AX foci colocalized with EdU foci (Supplemental Fig. S6A,B). However, colocalization of EdU and γH2AX foci was significantly reduced in the MCM9-KO cells (Supplemental Fig. S6B). Taken together, these results are consistent with the hypothesis that MCM8–9 contributes significantly to DNA synthesis after forks have been stalled due to a lack of the MCM2–7 complex.

It has been shown that a reduction in MCM2–7 expression causes spontaneous fork stalling by an extension in the size of replicons (Moreno et al. 2016). If, as we propose, MCM8–9 functions as a backup replicative helicase, the MCM2–7-depleted cells should have an increased reliance on MCM8–9. To test this hypothesis, we treated the MCM9-WT and MCM9-KO cells with various doses of auxin to reduce, but not eliminate, the cellular level of MCM2-mAID. We observed that the MCM9-WT cells were more resistant to a reduction in the level of MCM2-mAID than the MCM9-KO cells (Fig. 5E), supporting the hypothesis that MCM8–9 functions as a backup replicative helicase.

MCM8–9 promotes DNA synthesis during DSB-induced HR

Considering that MCM8–9 plays a role in HR repair (Lutzmann et al. 2012; Nishimura et al. 2012; Park et al. 2013), we investigated whether DNA synthesis occurring during DSB-induced HR requires MCM8–9. For this, we used an established HR reporter system in chicken DT40 cells, in which a copy of an SCneo substrate is stably introduced at the ovalbumin locus (Nishimura et al. 2012). These cells can repair the neomycin (Neo) resistance gene by HR upon expression of I-SceI, an endonuclease that generates a DSB in S2neo (Fig. 6A). If repaired by HR, two outcomes are possible, depending on whether short-tract gene
conversion (STGC) or long-tract gene conversion (LTGC) is used (Johnson et al. 1999). To generate the STGC product, only 300 base pairs (bp) of DNA synthesis is required. In contrast, >3.3 kb of DNA synthesis is required to generate the LTGC product. A similar large product can be generated by sister chromatid exchange (SCE), but this is a rare event (Johnson and Jasin 2000).

To analyze the putative role of MCM8–9 in DSB-induced HR, we compared wild-type DT40 cells with MCM8-KO or MCM9-KO. Initially, we looked at HR efficiency by quantifying Neo-resistant clones following I-SceI expression. This revealed that the MCM8-KO and MCM9-KO cells showed comparably reduced HR efficiencies, as reported previously (Fig. 6B; Lutzmann et al. 2012; Nishimura et al. 2012). Next, we analyzed the Neo-resistant clones that did arise in the MCM8-KO and MCM9-KO cells to define the outcome of the HR reactions. LTGC products can be distinguished from STGC by the size of a SacI–KpnI restriction fragment (Fig. 6A). In the wild-type background, 17% of clones were generated by LTGC, while this was reduced to 5% in the MCM8-KO and MCM9-KO cells (Fig. 6C,D). Conversely, STGC frequencies were elevated in the MCM8-KO and MCM9-KO cells. Taken together, we conclude that the MCM8–9 complex promotes DNA synthesis during DSB-induced HR.

MCM8–9-dependent DNA synthesis is distinct from RAD52-dependent MiDAS

Recently, it was shown that DNA synthesis could occur at common fragile sites in the human genome in the early stages of mitosis if cells are exposed to replication stress (Minocherhomji et al. 2015). This MiDAS requires RAD52 and is mechanistically distinct from conventional DNA replication (Bhowmick et al. 2016). Therefore, we investigated whether MCM8–9 might function in MiDAS. To this end, we disrupted the MCM8 gene in U2OS cells, in which MiDAS is known to occur very efficiently (Supplemental Fig. S7A–C; Minocherhomji et al. 2015). We then treated MCM8-WT and MCM8-KO U2OS cells with a low dose of the DNA polymerase inhibitor aphidicolin to induce replication stress and with the CDK1 inhibitor RO-3306 to induce a late G2-phase arrest. The cells were then released into mitosis in the presence of EdU to define sites of new DNA synthesis occurring in mitosis. Unexpectedly, we found that two independent clones of MCM8-KO cells showed an increase in EdU incorporation compared with MCM8-WT cells, showing that MiDAS is enhanced in MCM8-KO cells (Fig. 7A,B). Consistent with there being no role for MCM8–9 in MiDAS, the frequency of 53BP1 nuclear bodies in the following G1 phase (a marker of failed MiDAS) was unchanged in the MCM8-KO cells (Supplemental Fig. S7D,E; Minocherhomji et al. 2015). Interestingly, a similar increase in EdU incorporation has been observed previously in RAD51-depleted cells (Bhowmick et al. 2016). Therefore, we analyzed the epistatic relationship between RAD51 and MCM8–9. For this, we depleted RAD51 using siRNA in MCM8-WT and MCM8-KO cells and then quantified MiDAS (Fig. 7C; Supplemental Fig. S7F). We observed that RAD51 depletion enhanced the level of MiDAS in the MCM8-WT cells but not the MCM8-KO cells, consistent with RAD51 and MCM8–9 operating in the same pathway (Fig. 7C). We conclude that the RAD51/MCM8–9 axis promotes a backup form of DNA synthesis that is distinct from MiDAS.

Discussion

Inactivating MCM2–7 as a novel strategy for stalling replication forks

Replication stress is frequently induced in cells by exposure to aphidicolin or hydroxyurea. These agents can
lead to DSB formation, albeit after a long period of drug exposure (Petermann et al. 2010; Toledo et al. 2013). However, it is impossible to study DSB-induced DNA synthesis using these inhibitors, as they inhibit polymerase function. An alternative approach is to use DNA-damaging agents, but these cause the stalling of only a subset of replication forks, making it difficult to avoid fork convergence (Supplemental Fig. S1). Therefore, an efficient experimental system is required to permit the analysis of protective mechanisms involved in dealing with fork stalling.

In prokaryotes, the response to fork stalling has been studied either by using the replication terminator Ter or through inactivation of the DnaB replicative helicase (Horiuchi and Fujimura 1995; Seigneur et al. 2000). Similarly, protein-mediated endogenous barriers have been used for arresting replication forks in yeasts (Ahn et al. 2005; Calzada et al. 2005; Lambert et al. 2005). Recently, the E. coli Tus/Ter system was transplanted into yeast and mouse cells to serve as a heterologous replication barrier (Larsen et al. 2014; Willis et al. 2014). These studies revealed detailed molecular events following fork pausing, in which recombination proteins often played a prominent part. Importantly, Willis et al. (2014) showed that the HR induced following Tus-induced fork pausing was regulated differently from that induced by a DSB.

Fork stalling caused by the inactivation of the replicative MCM2–7 helicase has been achieved using the heat-inducible degron system in budding yeast (Labib et al. 2001). Analogous studies using the same experimental system have not been possible in human cells until now because of the requirement for such a drastic temperature shift to induce protein degradation. We have now overcome this problem and achieved inactivation of MCM2–7 at replication forks for the first time in human cells by using AID technology (Nishimura et al. 2009; Natsume et al. 2016). Although simultaneous stalling of all forks is not expected to occur regularly under physiological conditions [and therefore some caution must be applied to interpretation using the MCM2-mAID cells], we suggest that this system could have broad applicability in the study of fork stalling in the future. Consistent with a previous report in yeast (Labib et al. 2001), we showed that inactivation of MCM2–7 in human cells causes fork stalling. Interestingly, this revealed the presence of residual DNA synthesis that occurred via an MCM8–9-dependent process. This new DNA synthesis was apparently not robust enough for the cells to complete S phase, indicating that MCM8–9-dependent DNA synthesis is either incapable of rescuing all stalled forks or lacks the processivity to cover the full genome. Alternatively, efficient fork restart might not be possible due to a limitation in the cellular level of HR repair factors in cases where multiple forks collapse simultaneously. MCM8–9-dependent synthesis might be designed to operate less efficiently than conventional S-phase replication and might normally be called into action at only a very small number of irreversibly stalled replication forks. Indeed, the MCM8–9 system might represent a double-edged sword for the maintenance of genome integrity. In support of this idea, the expression level of MCM8–9 is >100 times lower than that of MCM2–7 (Beck et al. 2011), and we note that overexpression of MCM8–9 is detrimental to proliferation (our unpublished data).

The MCM8–9 complex promotes HR-mediated DNA synthesis

The MCM8–9 complex has been implicated previously in HR, although there is a lack of consensus concerning the process involved [Nishimura et al. 2012; Park et al. 2013; Lee et al. 2015]. We demonstrated that the formation of MCM8 foci is RAD51-dependent (Fig. 4B,C) and that MCM8–9 is required for both DNA synthesis after fork stalling induced by MCM2-mAID degradation (Fig. 5) and DSB-induced HR (Fig. 6). These results are consistent with previous observations showing that loss of the MCM8 homolog in mice and flies affects meiotic recombination only at a stage after the loading of the meiosis-specific RAD51 homolog DMC1 (Blanton et al. 2005; Lutzmann et al. 2012). Considering that the CMG helicase and the MCM8–9 complex each possess helicase activity [Ivics et al. 2010; Traver et al. 2015], we propose that MCM8–9 promotes DNA synthesis following RAD51-dependent DNA strand invasion between sister chromatids by acting as an alternative replicative helicase (Fig. 7D). In the future, it will be interesting to test whether DNA synthesis driven by MCM8–9 is semiconservative, like in normal DNA replication, or conservative in nature. This is because recent reports indicated that DNA synthesis occurring during BIR in yeast and DNA synthesis occurring during both the alternative lengthening of telomeres process and MiDAS in human cells are conservative events [Saini et al. 2013; Bhowmick et al. 2016; Roumelioti et al. 2016]. It should be noted that, even though we propose that MCM8–9 promotes DNA synthesis in HR, it is possible that MCM8–9 is also involved in the resection of DSB ends (Lee et al. 2015). Both the human HCT116 and the chicken DT40 cells that we studied are recombination-proficient, and therefore any defect in resection might be masked by the availability of redundant resection systems dependent on BLM, DNA2, or EXO1 [Gravel et al. 2008; Mimitou and Symington 2008; Zhu et al. 2008].

HR involved in interstrand cross-link (ICL) repair

We reported previously that the MCM8–9 complex is involved in HR-mediated ICL repair (Nishimura et al. 2012). Studies using Xenopus egg extracts showed that ICL repair in a replication-competent plasmid proceeds only after the convergence of replication forks on either side of the ICL [Raschle et al. 2008; Zhang et al. 2015]. DSBs generated by ICL unhooking are repaired using HR. Indeed, MCM8–9 has been detected at sites of ICLs in the Xenopus system but was found not to be essential for ICL repair [Park et al. 2013]. On the other hand, cells deficient in MCM8 or MCM9 are hypersensitive to ICL-inducing anti-cancer agents but only marginally sensitive to ionizing radiation, which generates two-ended DSBs...
late that, in eukaryotes, the MCM2-mediated DNA replication (Hawkins et al. 2013). We speculate that the helicase can carry out both origin-dependent and HR-mediated DNA synthesis, respectively. This division of labor might have occurred in response to the establishment of the replication licensing system in eukaryotes – a single fork might have to travel over a long distance.

Noncanonical DNA syntheses as a backup of DNA replication

We showed that the RAD51/MCM8–9 axis operates separately from MiDAS. We therefore propose that cells have at least three systems to deal with instances of fork stalling (Fig. 7D): fork convergence, MCM8–9-dependent DNA synthesis, and MiDAS. We hypothesize that MCM8–9-dependent DNA synthesis and MiDAS are mechanistically related by being functionally analogous to yeast BIR but use a different set of HR factors.

Archaeal replication was carried out by an MCM homo-hexameric helicase. It is likely that the archaeal MCM helicase can carry out both origin-dependent and HR-mediated DNA replication [Hawkins et al. 2013]. We speculate that, in eukaryotes, the MCM2–7 and MCM8–9 helicases evolved from a single ancestral MCM in order to catalyze origin-dependent DNA replication and HR-mediated DNA synthesis, respectively. This division of labor might have occurred in response to the establishment of the replication licensing system in eukaryotes [Blow and Dutta 2005]. Evolutional loss of the MCM8 and MCM9 genes in yeast might be due to the fact that the number and distribution of origins on chromosomes evolved for optimal fork convergence so that MCM8–9-dependent DNA synthesis was dispensable in the ancestor of yeast [Liu et al. 2009; Newman et al. 2013]. Further analysis of the fate of stalled forks and the role of MCM8–9 in their repair will hopefully reveal the relationship between conventional DNA replication and HR-mediated DNA synthesis in human cells.

Materials and methods

Cell lines

Genetically engineered HCT116 cell lines used in this study were as follows: HCT116 MCM2-mAID [clone 1: #269; clone 2: #270], HCT116 MCM2-mAID AtAFB2 ([#310]), HCT116 MCM2-mAID AtAFB2 MCM8–mCherry2 [#353], and HCT116 MCM2-mAID AtAFB2 MCM8–mCherry2 MCM9–KO [clone 1: #395; clone 2: #396; clone 3: #398]. Genetically engineered U2OS cell lines used in this study were as follows: U2OS MCM8–9 (clone 1: #513; clone 2: #514).

Cell culture, transfection, and cloning

Human cell culture was undertaken as described previously [Natsume et al. 2016]. To induce the degradation of MCM2-mAID, 500 µM indole-3-acetic acid (IAA; a natural auxin, Nacalai Tesque) was added to the culture medium unless otherwise noted. The RAD51 inhibitor RI-1 [Abcam] and bleomycin [Nippon Kayaku] were used at concentrations of 100 µM and 10 µg/mL, respectively. Transfection was performed using FuGENE HD [Promega]. Transfected cells were selected with 1 µg/mL puromycin, 700 µg/mL G418, or 100 µg/mL HygroGold [InvivoGen]. The detailed procedure for generation of mutant cells was described previously [Natsume et al. 2016].

Cell synchronization

HCT116 cells were synchronized in the G1 phase as described previously [Javanmoghadam-Kamrani and Keyomarsi 2008]. Briefly, asynchronously growing HCT116 cells were treated with 20 µM lovastatin [LKT Laboratories] for 24 h to arrest them in G1 phase. Following that, the cells were washed twice with fresh medium and then grown in medium containing 2 mM mevalonic acid [Sigma-Aldrich].

RNAi

For depletion of the MCM2 and MCM5 proteins, HCT116 cells were transfected with 50 nM Silencer Select siRNAs (Thermo Fisher Scientific) using Lipofectamine RNAiMAX reagent [Thermo Fisher Scientific] following the manufacturer’s instructions. The transfected cells were harvested after 72 h. RAD51 depletion in U2OS cells was performed as described previously [Bhowmick et al. 2016]. We used the following siRNAs: siCONT [negative control; 43908943], MCM2-1 [88586], MCM2-2 [85857], MCM2-3 [88588], MCM5-1 [88595], and Mcm5-1i and Mcm5-2i [Ge et al. 2007].

Plasmid construction

We used pX330-U6-Chimeric_BB-CBh-hSpCas9 (Addgene, 42230) for the construction of CRISPR plasmids following a published protocol [Ram et al. 2013]. To construct a donor plasmid for the expression of A. thaliana AFB2 [AtAFB2] from the AAVS1 locus, pMK232 was modified [Natsume et al. 2016]. The donor plasmid for tagging MCM2 with mAID was constructed by using PCR-amplified homology arms [1 kb each] and pMK286. The donor plasmid for tagging MCM8 with mCherry2 was constructed by using PCR-amplified homology arms [850 bp each] and pMK281. The donor plasmid for generating MCM8–KO in a U2OS background was constructed by using PCR-amplified homology arms [900 bp each] and a puromycin-resistant gene from pMK194.

Genomic PCR

To prepare genomic DNA, cells were lysed in buffer [100 mM Tris-HCl at pH 8.0, 200 mM NaCl, 5 mM EDTA, 1% SDS, and 0.6 mg/mL proteinase K] for 1 h at 55°C. After isopropanol precipitation, DNA pellets were washed with 70% ethanol and resuspended in TE containing 50 µg/mL RNase A overnight at 37°C. Genomic PCR was performed using Takara Gold DNA polymerase [Takara Bio] according to the manufacturer’s instructions (30 cycles of the following protocol: 10 sec at 98°C, 15 sec at 55°C, and 0.5 min at 68°C per kilobase).

Flow cytometry

Cells were collected and fixed in 70% ethanol. Fixed cells were washed once with PBS and then resuspended in PBS containing 1% BSA, 50 µg/mL RNase A, and 40 µg/mL propidium iodide.
After incubation for 30 min at 37°C, the cells were filtered through a nylon mesh filter (42-µm pore size). DNA content was measured using an Accuri C6 flow cytometer (BD Biosciences) and analyzed by FCS Express 4 software [De Novo Software]. For the analysis of DNA synthesis, cells were pulse-labeled with 30 µM BrdU before fixation with 90% ethanol. After washing with PBS, the fixed cells were treated with 2 M HCl and 0.5% Triton X-100 for 30 min for denaturation of genomic DNA. The cells were gently resuspended in 0.1 M Na2B4O7 (pH 8.5) and incubated for 30 min. After washing once with the antibody solution (1% BSA, 0.2% Tween 20 in PBS), cells were treated with the anti-BrdU antibody (BD Biosciences, B44) diluted in the antibody solution for 30 min. After washing once, the cells were treated with the TMC5-mCherry2, RFP (Sigma, 00020911), and anti-phospho-CHK1 (Ser345) antibody (Cell Signaling, 2341). For detection of MCM8-mCherry2, RFP-Booster ATTO 594 (Chromotech, rba594) was used.

**Immunofluorescence staining**

HCT116 cells were cultured in a glass-bottomed dish (MatTek) before fixation with 3.7% formaldehyde/PBS for 15 min. After washing twice with PBS, the cells were permeabilized with 0.5% Triton X-100/PBS for 20 min followed by a blocking treatment with 3% skim milk/PBS for 1 h. After washing twice with PBS, primary antibodies diluted in 1% BSA/PBS were applied before incubation for 1 h at room temperature. After washing three times with 0.05% Tween 20/PBS (PBS-T), secondary antibodies diluted in 1% BSA/PBS were applied before incubation for 1 h at room temperature. The cells were washed twice with PBS-T and once with PBS before DNA staining with 5 µg/mL Hoechst 33342 in PBS for 30 min. The coverslips were overlaid with Vectashield mounting medium (Vector Laboratories). Images were captured using a DeltaVision microscope equipped with deconvolution software, an incubation chamber, and a CO2 supply (GE Healthcare Life Sciences). For live-cell imaging, HCT116 cells were cultured in a glass-bottomed dish (MatTek) at 37°C with 5% CO2. To visualize nuclei in live cells, 10 µg/mL Hoechst 33342 was added to the medium before observation. DNA damage foci were analyzed using the Volocity software (PerkinElmer).

**DNA fiber assays**

DNA fiber assays were performed following a published protocol with minor modifications (Schwab and Niedzwiedz 2011). Cells were pulse-labeled with 25 µM IdU for 30 min followed by a second labeling with 250 µM CldU for 30 min. The labeled cells were collected in ice-cold medium. Two microliters of cell suspension were gently resuspended in 0.1 M Na2B4O7 (pH 8.0, 1% sarkosyl, 0.2% sodium deoxycholate, 1 mg/mL proteinase K) overnight at 37°C. The cell plugs were washed once with the plug wash buffer (20 mM Tris-HCl at pH 8.0, 50 mM EDTA) before insertion into the wells of a 0.9% agarose gel. PFGE was performed using a CHEF Mapper PA PFGE system (Bio-Rad) with 0.5x TBE for 21 h at 14°C as follows: block 1: 9 h, 120° pulse angle, 5.5 V/cm, 30 sec to 18 sec switch time; block 2: 6 h, 117° pulse angle, 4.5 V/cm, 18 sec to 9 sec switch time; and block 3: 6 h, 112° pulse angle, 4.0 V/cm, 9 sec to 5 sec switch time. The gel was stained with GelGed (Biotium), and images were acquired using a ChemiDoc touch imaging system (Bio-Rad) and analyzed using Image Lab software (Bio-Rad).

**PFGE**

PFGE was performed using a published protocol [Zellweger et al. 2015]. To prepare cell plugs, 2 x 10^6 cells were embedded in 1% 2-hydroxyethylagarose (Sigma-Aldrich) using the 50-well plug mold (Bio-Rad). To lyse cells, the cell plugs were incubated in the plug lysis buffer (100 mM EDTA at pH 8.0, 1% sarkosyl, 0.2% sodium deoxycholate, 1 mg/mL proteinase K) overnight at 37°C. The cell plugs were washed once with the plug wash buffer (20 mM Tris-HCl at pH 8.0, 50 mM EDTA) before insertion into the wells of a 0.9% agarose gel. PFGE was performed using a CHEF Mapper PA PFGE system (Bio-Rad) with 0.5x TBE for 21 h at 14°C as follows: block 1: 9 h, 120° pulse angle, 5.5 V/cm, 30 sec to 18 sec switch time; block 2: 6 h, 117° pulse angle, 4.5 V/cm, 18 sec to 9 sec switch time; and block 3: 6 h, 112° pulse angle, 4.0 V/cm, 9 sec to 5 sec switch time. The gel was stained with GelGed (Biotium), and images were acquired using a ChemiDoc touch imaging system (Bio-Rad) and analyzed using Image Lab software (Bio-Rad).

**Protein detection**

To prepare whole-cell extracts, cells were lysed in RIPA buffer (25 mM Tris-HCl at pH 7.6, 150 mM NaCl, 1% Nonidet P-40, 1% sodium deoxycholate, 0.1% SDS) containing a protease inhibitor cocktail (Complete EDTA-free, Roche). Protein concentration was then measured using a Bradford assay kit (Thermo Fisher Scientific) before primary antibody treatment. EdU was visualized using Click-iT Plus Alexa fluor 647 imaging kit (Thermo Fisher Scientific) before primary antibody treatment following the manufacturer’s instruction.

**Antibodies**

Antibodies used for immunoblotting and immunofluorescence were as follows: anti-MCM8 and anti-MCM9 antibodies (raised in rabbits; in-house antibodies), anti-MCM2 antibody (Santa Cruz Biotechnology, sc-9839), anti-MCM5 antibody (Santa Cruz Biotechnology, sc-22780), anti-mAID antibody [MBL, M214-3], anti-RFP antibody [MBL, M204-3], anti-RAD51 antibody [BioAca- demia, 70001], anti-γH2AX antibody [Millipore, 05-636], anti-53BP1 antibody [Santa Cruz Biotechnology, sc-22760], anti-His- tone H3 [Abcam, ab1791] and Active Motif, 39763], anti-a-tubulin [Sigma, 00020911], and anti-phospho-CHK1 (Ser345) antibody (Cell Signaling, 2341). For detection of MCM8-mCherry2, RFP-Booster ATTO 594 (Chromotech, rba594) was used.
I-SceI-induced gene conversion assay

The I-SceI-induced gene conversion assay was performed as described previously [Yamamoto et al. 2005].

Detection of MiDAS

The assay was performed as described previously with minor modifications [Minocherhomji et al. 2015]. Briefly, the assay used Click-iT chemistry according to the manufacturer’s instructions but with a 1x final concentration of the Click-iT EdU buffer additive (Click-iT EdU Alexa fluor 594 imaging kit, Thermo Fisher Scientific). Asynchronously growing cells were treated with low-dose APH (0.4 µM) and RO-3306 (9 µM) (Sigma) for 16 h. Cells synchronized in late G2 were released into early prophase by vigorous washing (three to four times for up to 5 min each with 1x PBS prewarmed to 37°C). Subsequently, cells in early prophase were maintained for 30 min at 37°C in a humidified atmosphere containing 5% CO2 in prewarmed fresh medium supplemented with10µM EdU (Thermo Fisher Scientific). Loosely attached mitotic cells were then shaken off and seeded on polylysine-coated slides and kept for 10 min at room temperature before simultaneous fixation and permeabilization using PTEMP buffer and subsequent EdU detection using Click-iT chemistry.

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References


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