Mechanism of replication origin melting nucleated by CMG helicase assembly

Since the discovery of the double helix, molecular biologists have been asking how the separation of two DNA strands is nucleated after the initiation of chromosome replication. In vitro reconstitution of bacterial1, viral1 and eukaryotic DNA replication1 have started to address this question. By studying these systems, a universal role has been identified for ATP binding by multimeric enzymes that untwist the double helix at the start of replication. Existing atomic models of initiators and helicases bound to single-stranded DNA mimic the structure of origin DNA immediately after melting4-6. However, to understand the mechanism of the ATP-triggered opening of duplex DNA at the molecular level, the structure of an origin duplex caught in the act of nucleating a replication bubble must be obtained. To achieve the opening of origin DNA, bacteria7 and eukaryotic viruses5,6 use one single protein that oligomerizes around the double helix and causes its deformation, but such melting intermediates have not to our knowledge been structurally characterized so far. Origin opening in Saccharomyces cerevisiae requires not one, but thirty-two distinct polypeptides that act sequentially. First, the origin recognition complex (ORC) together with loading factors Cdc6 and Cdt1 recruit a set of two ring-shaped MCM helicases that form an inactive double hexamer around duplex DNA7. Activation requires the recruitment of two firing factors, Cdc45 and GINS and base pairing is broken at this stage in origin activation remains to be determined. After the recruitment of Mcm10, the lagging-strand template is ejected from the MCM ring pore, which leads to the establishment of the replication fork and the ATPase-powered translocation along single-stranded DNA1. How the CMG selects the translocation strand in this context is unknown. Assembly of two CMGs at an origin disrupts the double hexamer interface1. Mapping the relative orientation of the two separated CMGs on the origin DNA is important to understand how replication forks are established bidirectionally and how replisome stability is maintained in the early stages of replication initiation17-22.

**CMG assembly on chromatinized origin DNA**

To understand how ATP-dependent CMG formation leads to double hexamer separation and DNA untwisting, we used electron microscopy to image origin-dependent CMG formation reconstituted in vitro with Cdc45 to the double hexamer11-15. Sld3 is in turn phosphorylated by the Clb5–Cdc28 (CDK) kinase, which also phosphorylates the firing factor Sld2. Phospho-Sld2 and phospho-Sld3 bind Dpb11, which engages Pol ε and GINS to mediate their origin recruitment11,12,16. After ADP release and ATP binding by MCM, GINS and Cdc45 stably engage MCM, forming two distinct CMG assemblies that disrupt the double hexamer interface through an unknown mechanism. Topology footprint assays indicate that CMG formation leads to partial DNA untwisting, but whether base pairing is broken at this stage in origin activation remains to be determined. After the recruitment of Mcm10, the lagging-strand template is ejected from the MCM ring pore, which leads to the establishment of the replication fork and the ATPase-powered translocation along single-stranded DNA1. How the CMG selects the translocation strand in this context is unknown. Assembly of two CMGs at an origin disrupts the double hexamer interface1. Mapping the relative orientation of the two separated CMGs on the origin DNA is important to understand how replication forks are established bidirectionally and how replisome stability is maintained in the early stages of replication initiation17-22.
purified yeast proteins and in a near-native chromatin environment. To this end, we reconstituted CMG on ARS1 origin DNA flanked at both ends by a nucleosome assembled on strong Widom positioning sequences. Nucleosome capping of the naked, AT-rich ARS1 DNA recapitulates the architecture of chromatinized origins that is found in cells and serves to trap double hexamers on duplex DNA, preventing dissociation by sliding. Double hexamers were (i) loaded onto origin DNA using MCM–Cdt1, ORC, Cdc6 and ATP; (ii) phosphorylated with DDK in solution; and (iii) isolated using Strep-TactinXT-coated paramagnetic beads that capture a twin-strep tag on histone H3 of the nucleosome. After a high-salt wash that removes helicase loading intermediates and beads that capture a twin-strep tag on histone H3 of the nucleosome, we reconstituted CMG on ARS1 flanked at both ends by a nucleosome assembled on strong Widom positioning sequences. To find that on average 32% of double hexamers were converted to CMG. Of these, 70% were homo-dimeric and Pol ε-engaged (dCMGE) and 19% were single CMGs. In most of the dCMGE particles, GINS–Cdc45 and Pol ε mapped on opposite sides around the MCM ring, giving rise to a trans configuration (Fig. 1b and Extended Data Fig. 1c). The remaining dCMGE particles (11%) were in a cis configuration, with GINS–Cdc45–Pol ε located on the opposite sides around the MCM ring, giving rise to a trans configuration (Fig. 1b and Extended Data Fig. 1c). The remaining dCMGE particles (11%) were in a cis configuration, with GINS–Cdc45–Pol ε located on the same side (Fig. 1b and Extended Data Fig. 1c). In silico reconstitution (ReconSil), performed by overlaying nucleosome and MCM-containing 2D averages onto the corresponding particles in the raw micrographs, revealed that dCMGEs were tightly packed between the two flanking nucleosomes (Fig. 1b). A measured inter-nucleosome distance of 136 bp ± 23 bp (s.d.) (Fig. 1c) matches the expected 136 bp, supporting the notion that these represent bona fide reconstituted origins and not neighbouring particles bound to different DNA molecules. Similar results were obtained for chromatinized origins trapping a double hexamer (133 bp ± 18 bp (s.d.); Fig. 1c). We did not observe single CMGs trapped in between nucleosomes, suggesting that these may represent helicases that fell off the DNA (Fig. 1b). It is established that ARS1 DNA substrates capped by nucleosomes, or by covalently linked HpaII methyltransferase (MH) roadblocks, support double hexamer loading. Given the tighter protein packing that is caused by the formation of dCMGEs on the ARS1 origin, we asked whether...
enough space is available between nucleosomes for two activated CMGE particles to cross paths during the establishment of the replication fork. To address this question, we reconstituted roadblocked origin replication in vitro using a minimal set of replisome factors (Fig. 1d,e), matching established conditions that support the replication of an ARS-containing 10.6-kb supercoiled plasmid11 (Fig. 1f). DNA products separated by alkaline agarose gel electrophoresis showed that the nucleosome-flanked ARS1 substrate is copied in full (Fig. 1e). This is evident from the size of duplicated DNA, which is longer for the Widom-flanked ARS1, compared to a shorter construct in which nucleosomes are swapped for MH caps (Extended Data Fig. 1b and Fig. 1e). The tight packing of a dCMGE particle between two flanking nucleosomes raises the question of whether the CMGE dimer is a stable complex or whether it is formed as a result of the spatial constraints imposed by the two roadblocks that prevent dissociation. We reasoned that CMG assembly on longer, less crowded DNA substrates might allow enough space for the dCMGE complex to dissociate into two discrete CMGE particles. To test this hypothesis, we performed origin-dependent CMG assembly reactions on MH-capped 864-bp DNA substrates that contain an array of 6 consecutive ARS1 sequences separated by 40 bp of linker DNA. We only observed dCMGE particles and not separated CMGs on the array substrate, irrespective of the efficiency of double hexamer loading (Fig. 1g and Extended Data Fig. 1d–g). Thus, stability of the dCMGE complex assembled during ATP-dependent double hexamer activation is independent of nucleosomes and independent of the position of flanking roadblocks, as well as the level of protein saturation of DNA.

**Cryo-EM structure of the CMGE dimer**

Our previous NS-EM work on origin-dependent CMG reconstitution in vitro involved high-salt treatment of protein–DNA tethered to...
strepavidin-coated paramagnetic beads, followed by elution using DNA digestion. This procedure disrupted CMGE dimers but not the double hexamer, which suggests that the conversion of double hexamer to dCMGE reconfigures and weakens the MCM dimerization interface. To understand the conformational transitions that occur upon CMG formation, we determined the cryo-electron microscopy (cryo-EM) structure of a dCMGE complex assembled on chromatinized ARS1 DNA (Extended Data Fig. 2a–c). Both C2 symmetry and asymmetric refinement yielded a structure with limited resolution (Extended Data Figs. 2d–f and 3). However, symmetry expansion approaches revealed that the two rings in the dimer are identical. Combined with three-dimensional (3D) classification, variability analysis and refinement, followed by iterative cycles of contrast transfer function (CTF) refinement and Bayesian polishing, this process yielded a structure at 3.5 Å resolution (or 3.4 Å after density modification; Fig. 2a, Supplementary Video 1, Extended Data Figs. 2g–j, 3 and 4 and Extended Data Table 1). By overlaying two copies of the CMGE monomer to the lower-resolution dimer, we can therefore obtain a high-resolution view of the entire dCMGE assembly. While in the double hexamer the tight homo-dimerization interface is formed by the packing between six MCM zinc-finger domains (ZnfS; degenerate in Mcm3), the transition to dCMGE involves the loss of several trans-ring interactions and a one-subunit register shift for the remaining tethering elements. The residual inter-ring contacts involve the Mcm6 Znf that transitions from an Mcm6–Mcm6–2 to a Mcm6–Mcm4–6 interaction (Fig. 2b and Supplementary Video 2). What role this register shift might have during origin unwinding is addressed in the Supplementary Discussion. As the Mcm5–7 trans contact is disrupted after double-hexamer-to-dCMGE conversion, a helical insertion in the A domain of Mcm7 disengages from the Mcm5 A domain on the opposed ring, releasing a steric impediment that would prevent stable CMG assembly by hindering the association of GINS with Mcm5 (ref. 3) (Fig. 2c and Supplementary Video 2). As a result, the two CMG particles in the dCMGE complex become splayed open, pivoting around Mcm6 and creating a large cavity that exposes 1.5 turns of duplex DNA. The intervening DNA appears to stabilize the dCMGE dimer interface (Fig. 2d). In fact, partial digestion with the restriction enzyme Msel promotes the disassembly of the dCMGE complex into separated CMGs (Fig. 2e and Extended Data Fig. 1g).

Nucleotide and DNA binding in the dCMGE complex

Structural analysis of the double-hexamer-to-dCMGE transition indicates that three sites (Mcm2–5, 5–3 and 3–7) exchange ADP for ATP, whereas Mcm7–4 remains ADP-bound. Mcm4–6, which is nucleotide-free in the double hexamer, is ADP-bound in the dCMGE complex, suggesting consecutive ATP binding and hydrolysis on the path to dCMGE formation. The Mcm6–2 interface instead transitions from an ATP-bound state in which the bipartite catalytic site is open and hydrolysis-incompetent, to a more compact, hydrolysis-competent state that contains a mixture of ADP and ATP in the nucleotide-binding pocket (Fig. 3a,b and Supplementary Video 3). Overall, four of the six ATPase sites of MCM release ADP and bind ATP, in agreement with an analysis of bound nucleotides by thin-layer chromatography. This nucleotide exchange in turn alters the way in which duplex DNA is gripped by ATPase pore loops of the MCM. Although in the double hexamer, MCM engages in the same number of contacts with both the leading and the lagging strands, the vast majority of MCM interactions in the dCMGE involve the leading-strand template (Extended Data Fig. 5a). The ATPase domains of Mcm3, Mcm5, Mcm2 and Mcm6 provide most of the leading-strand contacts, which explains how the double-hexamer-to-dCMGE transition leads to selection of the translocation strand (Fig. 3c). This leading-strand binding mode in dCMGE is consistent with previously reported structures of single- and duplex-DNA-engaged recombinant CMG (refs. 13–14), which indicates that interaction with the leading strand is conserved from initiation to termination (Fig. 3d). Given the sparse pore-loop contacts, the lagging-strand template appears to be in turn poised for ejection from the MCM ring pore, which is required to achieve the establishment of the replication fork. How nucleotide-triggered conformational changes in MCM affect the structure of duplex DNA will be discussed in the next paragraph.

Mechanism of DNA-bubble nucleation

Captured between the ATPase and the N-terminal pore loops in the dCMGE complex, a stretch of seven base pairs is unwound. This observation agrees with previous topology footprint measurements that indicate that 0.7 turns of DNA become untwisted per CMG complex, after the double-hexamer-to-CMG transition in the absence of Mcm10 (ref. 3). On the basis of the cryo-EM density of the untwisted
lagging strand, three flipped-out bases can be confidently built (Fig. 4a), which are stabilized by two conserved residues (T423 and R424) located within the Mcm6-specific insertion of the N-terminal hairpin insertion. c, Within the Mcm2 ATPase domain, only residues K587 contacts duplex DNA in the double hexamer. In dCMGE, five additional Mcm2 ATPase residues contact the DNA, which promotes widening of the minor groove, untwisting of duplex DNA and disruption of base pairing. d, Topology footprint assay for DNA unwinding. Complete reactions contained all firing factors after MCM loading plus TopoI; omission of DDK blocks all untwisting. Omission of Mcm10 captures the initial untwisted state1. This initially untwisted state generates topoisomers of −2 and −3 (cyan arrowheads) as previously observed1. Additional negatively supercoiled topoisomers can be detected when Mcm10 is present, indicating further untwisting after ejection of the lagging strand from CMG. No topoisomers were observed with the Mcm2 6A mutant. For gel source data, see Supplementary Fig. 1. This experiment was performed twice. e, NS-EM CMG averages derived from the CMG assembly reaction using wild-type or Mcm2 6A MCMs. CMG assembly reactions were performed on a 2 × MH DNA template (Extended Data Fig. 1c) to reduce background single CMG particles that are present owing to incomplete roadblocking of DNA. dCMGE is the product of CMG assembly using wild-type proteins, whereas Mcm2 6A primarily forms single CMGs. A minority of proteins are compatible with dCMG or dCMGE formation (although in the latter, Pol ε occupancy is only partial (dCMGE)). f, Mcm2 6A MCMs are converted to CMG at wild-type levels. P values were determined by two-tailed Welch’s t test; NS, not significant. This experiment was performed three times. Error bars, mean ± s.d. g, Mcm2 6A disrupts the dCMGE dimer mostly into single isolated CMGs. P values were determined by two-tailed Welch’s t test; **P = 0.0030. This experiment was performed three times. Error bars, mean ± s.d.

Fig. 4 | dCMGE formation leads to the untwisting of duplex DNA and breaks at least three base pairs. a, Cryo-EM density of origin DNA. b, 0.7 turns of the double helix become untwisted after dCMGE formation. Three orphan bases become stabilized by residues T423 and R424 from the Mcm6-specific N-terminal hairpin insertion. c, Within the Mcm2 ATPase domain, only residue K587 contacts duplex DNA in the double hexamer. In dCMGE, five additional Mcm2 ATPase residues contact the DNA, which promotes widening of the minor groove, untwisting of duplex DNA and disruption of base pairing. d, Topology footprint assay for DNA unwinding. Complete reactions contained all firing factors after MCM loading plus TopoI; omission of DDK blocks all untwisting. Omission of Mcm10 captures the initial untwisted state1. This initially untwisted state generates topoisomers of −2 and −3 (cyan arrowheads) as previously observed1. Additional negatively supercoiled topoisomers can be detected when Mcm10 is present, indicating further untwisting after ejection of the lagging strand from CMG. No topoisomers were observed with the Mcm2 6A mutant. For gel source data, see Supplementary Fig. 1. This experiment was performed twice. e, NS-EM CMG averages derived from the CMG assembly reaction using wild-type or Mcm2 6A MCMs. CMG assembly reactions were performed on a 2 × MH DNA template (Extended Data Fig. 1c) to reduce background single CMG particles that are present owing to incomplete roadblocking of DNA. dCMGE is the product of CMG assembly using wild-type proteins, whereas Mcm2 6A primarily forms single CMGs. A minority of proteins are compatible with dCMG or dCMGE formation (although in the latter, Pol ε occupancy is only partial (dCMGE)). f, Mcm2 6A MCMs are converted to CMG at wild-type levels. P values were determined by two-tailed Welch’s t test; NS, not significant. This experiment was performed three times. Error bars, mean ± s.d. g, Mcm2 6A disrupts the dCMGE dimer mostly into single isolated CMGs. P values were determined by two-tailed Welch’s t test; **P = 0.0030. This experiment was performed three times. Error bars, mean ± s.d.
or selected dropout controls, in the presence of TopoI (Fig. 4d and Extended Data Fig. 6b,c). As previously described, three topoisomers, α, α−1 and α+1, were visible in the absence of DDK, indicating that there was no MCM-dependent change in DNA topology. Omission of Mcm10 with wild-type MCM led to an additional accumulation of negatively supercoiled topoisomers −2 and −3, which are indicative of initial DNA untwisting consistent with that seen in the dCMGE structure. When all firing factors were present with wild-type MCM, a robust accumulation of −2 through to −6 negative supercoils was detected (lane 3), indicating additional DNA untwisting that arises after the ejection of the lagging strand from CMG. None of the topoisomers that are associated with either initial untwisting or full activation appeared when MCM containing the Mcm2 6A variant was assayed. This indicates that DNA engagement by Mcm2—as observed in our dCMGE structure—is essential for the initial untwisting of DNA and the subsequent ejection of the lagging strand from CMG. NS-EM analysis of the same Mcm2 6A mutant revealed that double hexamers can efficiently be converted to CMG (Fig. 4e–f); however, most fail to homo-dimerize and form complete dCMGE complexes (Fig. 4g). Hence, DNA binding by CMG is important for the stability of the dCMGE structure, in agreement with our observation that partial DNA digestion disrupts CMGE dimerization (Fig. 2e). The discovery that a MCM mutant that is unable to untwist DNA and support origin-dependent replication is competent in single leads to the stabilization of three orphan bases in the untwisted origin DNA duplex. d, In the double hexamer, the Mcm4 and Mcm7 h2i pore loops face downwards, with Mcm4 pushing against the Mcm6 N-terminal β-hairpin. This functions as a latch that maintains the Mcm6 wedge packed against the double hexamer dimerization interface. After CMG assembly, global changes in the ATPase tier of MCM cause the Mcm4 and Mcm7 h2i pore loops to move upwards, which releases the safety latch of the Mcm6 wedge insertion. This change promotes the Mcm6 wedge to swing upwards, with the R423 and T424 elements entering the MCM lumen to stabilize three orphan bases.

**Open DNA stabilized as the MCM dimer splits**

CMG assembly leads to the disruption of the double hexamer interface, but how this is linked to the ATPase state of MCM is unclear. When comparing the double hexamer and the dCMGE structures, we observed that the Mcm6 wedge insertion, which stabilizes the lagging-strand orphan bases in the dCMGE complex, is retracted and contributes to stabilizing the dimerization interface in the double hexamer (Fig. 5a). Residues T423 and R424 in the wedge insertion are surface-exposed and face on the outer perimeter of MCM in the double hexamer. As the DNA becomes untwisted, the Mcm6 wedge insertion disengages from the double hexamer interface and enters the helicase ring lumen in the dCMGE complex (Fig. 5b,c and Supplementary Video 5). In agreement with this observation, a combined T423E/R424E mutation (Mcm6 2E) supports CMG loading onto origin DNA to wild-type levels but negatively affects replication (Fig. 1d,f). Specific interactions between the ATPase and the N-terminal tiers in MCM reveal long-range allosteric changes that couple the ATPase state with the movement of the Mcm6 wedge. For example, the Mcm4 h2i ATPase element appears to act as a safety latch that keeps the Mcm6 wedge retracted in the double...
hexamer. With the double-hexamer-to-dCMGE transition, a rigid-body rotation in the ATPase domains of Mcm4 and Mcm7 releases this latch, which creates enough space for the Mcm6 wedge insertion to disengage from the double hexamer interface and invade the helicase central channel (Fig. 5d and Supplementary Video S). By compromising the Mcm4–6 latch interaction through the addition of five glutamate point mutations in the Mcm6 pore loop (T408E, Q409E, L410E, G411E and L412E), we generated an MCM variant (Mcm6 SE) that can be loaded to wild-type levels but is significantly defective for replication, possibly because it uncouples the ATPase state from the reconfiguration of the Mcm6 wedge (Fig. 1d,f). By combining our comparative structural and mutagenesis analyses, we propose a model whereby changes in the ATPase state promote dCMGE complex formation, which in turn couples melting of duplex DNA and splitting of the double hexamer. ADP release and ATP binding in the ATPase tier promotes the concerted movement of the h2i pore loops. Amongst these, Mcm4 h2i pulls on the ADP release and ATP binding in the ATPase tier promotes the concerted movement of the h2i pore loops. Amongst these, Mcm4 h2i releases a latch that pins the N-terminal Mcm6 wedge insertion that is packed at the double hexamer homo-dimerization interface. As the latch is released, the Mcm6 wedge can swing inside of the MCM pore and stabilize the orphan bases that become exposed after the disruption of DNA base pairing in the untwisted DNA duplex. Supplementary Video 5 describes these structural transitions.

**Discussion**

To ensure that replication occurs bidirectionally, several steps along the origin activation pathway in eukaryotic cells occur symmetrically. Symmetry is first established in G1 with the concerted and sequential loading of an inactive double hexamer15,16. During this process, origin loading of a first MCM hexamer creates a binding site for the symmetric loading of the second hexamer17. After entry into S phase, recruitment loading of an inactive double hexamer7,8. During this process, origin activation pathway in eukaryotic cells occur symmetrically. Symmetry is first established in G1 with the concerted and sequential loading of an inactive double hexamer15,16. During this process, origin loading of a first MCM hexamer creates a binding site for the symmetric loading of the second hexamer17. After entry into S phase, recruitment loading of an inactive double hexamer7,8.

The dCMGE nucleates two DNA bubbles inside each MCM ring, separated by 1.5 turns of duplex DNA, which might serve for the concerted recruitment of fork establishment factors, including Mcm10, RPA and Pol α. Mcm10 is known to trigger the ejection of the lagging strand and the ATPase-powered unwinding of the replication fork9. Although the mechanism of origin activation remains unknown, we note that Mcm10 engages the same N-terminal MCM elements11,14,14, that mediate CDMGE dimerization in our structural intermediate. A model for origin activation is presented in Extended Data Fig. 8 and further discussed in the Supplementary Discussion. Studies will be needed to establish whether Mcm10 engagement further disrupts the CDMGE dimer interface, thereby releasing the inhibitory interaction that impairs ATPase-powered DNA translocation and allowing helicase bypass.

DNA replication, transcription and recombination all require the untwisting and opening of the double helix. Recent studies have described these processes in the transcription pre-initiation complex that supports RNA synthesis9,10 and in the recombinases that promote strand exchange19,40. By contrast, the mechanism for the nucleation of DNA melting at an origin of replication has remained—to our knowledge—unknown for decades. Our work fills this gap. We describe the structure of the CDMGE replicative helicase assembled sequentially onto the ARS1 origin, by reconstituting a multistep cellular process that involves 32 polypeptides1. Base-pair disruption involves ATP-triggered changes in CDMGE that promote pulling of the leading-strand and pushing of the lagging-strand template DNA. Our findings provide a framework in which to study replication initiation.

**Online content**

Any methods, additional references, Nature Research sum- maries, source data, extended data, supplementary information, acknowledgements, peer review information; details of author contributions and competing interests; and statements of data and code availability are available at https://doi.org/10.1038/s41586-022-04829-4.

1. Douglas, M. E., Ali, F. A., Costa, A. & Diffley, J. F. X. The mechanism of eukaryotic CMG helicase activation. Nature 555, 265–268 (2018).
2. Duderstadt, K. E., Chuang, K. & Berger, J. M. DNA stretching by bacterial initiators promotes replication origin opening. Nature 478, 209–213 (2011).
3. Dean, D. F. & Hurwitz, J. Simian virus 40 large T antigen untwists DNA at the origin of DNA replication. J. Biol. Chem. 266, 5062–5071 (1991).
4. Enemark, E. J. & Joshua-Tor, L. Mechanism of DNA translocation in a replicative hexameric helicase. Nature 442, 270–275 (2006).
5. Ali, F. A. et al. Cryo-EM structures of the eukaryotic replicative helicase bound to a translocation substrate. Nat. Commun. 7, 10708 (2016).
6. Schuck, S. & Stenlund, A. Mechanistic analysis of local ori melting and helicase assembly by the papillomavirus E1 protein. Mol. Cell. 43, 776–787 (2011).
7. Remus, D. et al. Concerted loading of Mcm2–7 double hexamers around DNA during DNA replication origin licensing. Cell 139, 789–799 (2009).
8. Evrin, C. et al. A double-hexameric MCM-2 complex is loaded onto origin DNA during licensing of eukaryotic DNA replication. Proc. Natl Acad. Sci. USA 106, 20240–20245 (2009).
9. Moyer, S. E., Lewis, P. W. & Botchan, M. R. Isolation of the Cdc45/Mcm2-7/GINS (CMG) complex, a candidate for the eukaryotic DNA replication fork helicase. Proc. Natl Acad. Sci. USA 103, 10236–10241 (2006).
10. Ilves, I., Petoevíc, T., Pesavento, J. J. & Botchan, M. R. Activation of the MCM2-7 helicase by the Cdc45-GINS association promotes DNA melting at ARS1 origins. Mol. Cell. 37, 247–258 (2010).
11. Yeates, T. J., Deegan, T. D., Janska, A., Early, A. & Diffley, J. F. X. regulated by 1.5 turns of duplex DNA, which might serve for the concerted recruitment of fork establishment factors, including Mcm10, RPA and Pol α. Mcm10 is known to trigger the ejection of the lagging strand and the ATPase-powered unwinding of the replication fork9. Although the mechanism of origin activation remains unknown, we note that Mcm10 engages the same N-terminal MCM elements11,14,14, that mediate CDMGE dimerization in our structural intermediate. A model for origin activation is presented in Extended Data Fig. 8 and further discussed in the Supplementary Discussion. Studies will be needed to establish whether Mcm10 engagement further disrupts the CDMGE dimer interface, thereby releasing the inhibitory interaction that impairs ATPase-powered DNA translocation and allowing helicase bypass.

In particular, when CMG–SCFαβ2 is superposed to our dCMGE structure, an extensive steric clash can be identified between the E3 ligase engaged to one ring and the Mcm3 subunit from the opposed ring in the CMG dimer (Extended Data Fig. 7). The dCMGE nucleates two DNA bubbles inside each MCM ring, separated by 1.5 turns of duplex DNA, which might serve for the concerted recruitment of fork establishment factors, including Mcm10, RPA and Pol α. Mcm10 is known to trigger the ejection of the lagging strand and the ATPase-powered unwinding of the replication fork9. Although the mechanism of origin activation remains unknown, we note that Mcm10 engages the same N-terminal MCM elements11,14,14, that mediate CDMGE dimerization in our structural intermediate. A model for origin activation is presented in Extended Data Fig. 8 and further discussed in the Supplementary Discussion. Studies will be needed to establish whether Mcm10 engagement further disrupts the CDMGE dimer interface, thereby releasing the inhibitory interaction that impairs ATPase-powered DNA translocation and allowing helicase bypass.

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42. Sonnevile, R. et al. CUL-2LRR-1 and UBXN-3 drive replisome disassembly during DNA replication termination and mitosis. Nat. Cell Biol. 19, 488–479 (2017).

43. Mayle, R. et al. Mcm10 has potent strand-annealing activity and limits translocase-mediated fork regression. Proc. Natl Acad. Sci. USA 116, 798–803 (2018).

44. Lökke, M., Maloney, M. F. & Bell, S. P. Mcm10 regulates DNA replication elongation by stimulating the CMG replicative helicase. Genes Dev. 31, 291–305 (2017).

45. Schibbach, S., Aubara, S., Diennemann, C., Grabbe, F. & Cramer, P. Structure of RNA polymerase II pre-initiation complex at 2.9 Å defines initial DNA opening. Cell 184, 4064–4072 (2021).

46. Albar, S., Schibbach, S. & Cramer, P. Structures of mammalian RNA polymerase II pre-initiation complexes. Nature 594, 124–128 (2021).

47. Chen, Z., Yang, H. & Pavletich, N. P. Mechanism of homologous recombination from the RecA–ssDNA/dsDNA structures. Nature 453, 489–494 (2008).

48. Frigola, J., Remus, D., Mehanna, A. & Diffley, J. F. X. ATPase-dependent quality control of DNA replication origin licensing. Nature 493, 339–343 (2013).

49. Posse, V., Johansson, E. & Diffley, J. F. X. Eukaryotic DNA replication with purified budding yeast proteins. Methods Enzymol. 661, 1–33 (2021).

50. Coster, G. & Diffley, J. F. X. Bidirectional eukaryotic DNA replication is established by quasi-symmetrical helicase loading. Science 357, 314–318 (2017).

51. Kingston, I. J., Yang, J. S. Y. & Singleton, M. R. Biophysical characterization of the centromere-specific nucleosome from budding yeast. J. Biol. Chem. 286, 4021–4026 (2011).

52. Yueles, J. P. T., Jansa, A., Early, A. & Diffley, J. F. X. How the Eukaryotic replisome achieves rapid and efficient DNA replication. Mol. Cell 65, 105–116 (2017).

53. Bepler, T. et al. Positive-unlabeled convolutional neural networks for particle picking in cryo-electron micrographs. Nat. Methods 16, 1533–1630 (2019).

54. Zhang, K. Gctf: real-time CTF determination and correction. J. Struct. Biol. 193, 1–12 (2016).

55. Schindelin, J. et al. Fiji: an open-source platform for biological-image analysis. Nat. Methods 9, 676–682 (2012).

56. Punjani, A., Rubinstein, J. L., Fleet, D. J. & Brubaker, M. A. cryoSPARC: algorithms for rapid unsupervised cryo-EM structure determination. Nat. Methods 14, 290–296 (2017).

57. Zheng, S. Q. et al. MotionCor2: anisotropic correction of beam-induced motion for improved cryo-electron microscopy. Nat. Methods 14, 331–332 (2017).

58. Wagner, T. et al. Sphire-ocrYLO is a fast and accurate fully automated particle picker for cryo-EM. Commun. Biol. 2, 218 (2019).

59. Terswilliger, T. C., Sobolev, O. V., Afonine, P. V., Adams, P. D. & Read, R. J. Density modification of cryo-EM maps. Acta Crystallogr. D 76, 912–925 (2020).

60. Asar, D., Palovack, E. & Cheng, Y. UCSF pyem v0.5. Zenodo https://doi.org/10.5281/ zenodo.3576630 (2019).

61. Kelley, L. A., Mezulis, S., Yates, C. M., Wann, M. N. & Sternberg, M. J. E. The FlyWeb2 web portal for protein modeling, prediction and analysis. Nat. Protoc. 10, 845–858 (2015).
Methods

Cloning, expression and purification

ORC, Cdc6, Mcm2–7–Cdt1, DDK, CDK, Sld2, Sld3–Sld7, Cdc45, Dpb11, Polε, Polɛ exo-, Polα, Topo1, Mcm10 and yeast histone octamers were purified on the basis of previously established protocols13,24,33,48–51.

Cloning, expression and purification of MCM2–7–Cdt1 mutants.

Designed DNA fragments (Supplementary Table 1) were subcloned from pMA vectors (Supplementary Table 2) to pRS shuttle vectors (Supplementary Table 2), which were used to generate yeast strains (Supplementary Table 3) used to overexpress MCM2–7–Cdt1 mutants. The oMG25 DNA fragment was subcloned from pMG39 to pAM38 using MluI and XbaI restriction sites to obtain PMG69, which was integrated into the yJF21 yeast strain, thus generating the yAE64 strain that was used to overexpress the MCM26A mutant (Mcm6 T423E/R424E). The oMG28 DNA fragment was subcloned from plasmid pMG44 to pJF4 using BsiWI and SpH1 restriction sites to obtain PMG53, followed by the integration of PMG35 into the yAM20 strain, yielding the yAE610 strain, which was used for overexpression of the MCM62E mutant (Mcm6 T423E/R424E). The oMG28 DNA fragment was subcloned from plasmid pMG44 to pJF4 using BsiWI and SpH1 restriction sites, thus obtaining plasmid pMG54. The pMG35 plasmid was integrated into the yAM20 strain, yielding the yAE611 strain that was used to overexpress the MCM62E mutant (Mcm6 T408E/Q409E/L410E/K587A/W589A/K633A). The oMG27 DNA fragment was subcloned from pMG43 to pJF4 using SfiI and SpH1 restriction sites to obtain PMG53, followed by the integration of PMG35 into the yAM20 strain, yielding the yAE610 strain, which was used for overexpression of the MCM62E mutant (Mcm6 T423E/R424E). The oMG28 DNA fragment was subcloned from plasmid pMG44 to pJF4 using BsiWI and SpH1 restriction sites, thus obtaining plasmid pMG54. The pMG35 plasmid was integrated into the yAM20 strain, yielding the yAE611 strain that was used to overexpress the MCM62E mutant (Mcm6 T408E/Q409E/L410E/K587A/W589A/K633A). All MCM2–7–Cdt1 mutants were purified essentially as wild type50.

Cloning, expression and purification of GINS.

A gene block encoding a twin-strep tag and the first three codons of PSF3 was amplified and cloned into pJFDS by restriction-free cloning techniques. A list of primers and gene blocks used is included in Supplementary Table 1. BL21(DE3)-CodonPlus-RIL cells (Agilent) were transformed with GINS expression plasmid (pJL003). Transformant colonies were inoculated into a 250-ml LB culture containing kanamycin (50 µg ml⁻¹) and chloramphenicol (35 µg ml⁻¹), which was grown overnight at 37 °C with shaking at 200 rpm. The following morning, the culture was diluted 100-fold into 6× 11 of LB with kanamycin (100 µg ml⁻¹) and chloramphenicol (35 µg ml⁻¹). The cultures were left to grow at 37 °C until an OD₆₀₀ of 0.5 was reached; 0.5 mM IPTG was added to induce expression and cells were left shaking for 3 h. Cells were collected by centrifugation at 4,000 rpm for 20 min in a JS.4.2 rotor (Beckman). For lysis, cell pellets were resuspended in 80 ml of lysis buffer (20 mM Tris-HCl pH 8.5, 10% glycerol 0.5 mM EDTA, 500 mM KCl, Roche protease inhibitor tablets and 2 mM tris(2-carboxyethyl) phosphine (TCEP)) + 0.7 mM PMSF. The lysate was sonicated for 120 s (5 s on, 5 s off) at 40% on a Sonics Vibra-Cell sonicator. Insoluble material was removed by centrifugation at 20,000 rpm for 30 min in a JS.25.50 rotor (Beckman). The supernatant was loaded by gravity onto a 5-ml Strep-TactinXT column (IBA). The resin was washed extensively with lysis buffer, MH was eluted by the addition of 12 ml of 1× BXT (IBA) supplemented with 10% glycerol and 1 mM DTT. The MH-containing fractions were pooled and loaded onto a HiLoad 16/600 Superdex 75 column equilibrated in the same buffer. Fractions containing MH were pooled and dialysed overnight in 1× buffer BXT (IBA) supplemented with 10% glycerol and 1 mM DTT. The sample was concentrated and loaded onto a HiLoad 16/600 Superdex 200 equilibrated in the same buffer. GINS-containing fractions were pooled, aliquoted and snap-frozen in liquid N₂. About 36 mg MH was purified from a 6-litre culture.

Preparation and purification of chromatinized origin DNA.

Soluble yeast nucleosomes were reconstituted from octamers and DNA by salt gradient dialysis in several steps from 2 to 0.2 M NaCl as previously described34. Following nucleosome refolding, a final dialysis step was performed into loading buffer (25 mM HEPES-KOH pH 7.6, 80 mM KCl, 100 µM sodium acetate, 0.5 mM TCEP) and loaded onto a Superose 6 Increase 3.2/300 column equilibrated in the same buffer. Fractions containing ARS1 origin DNA were collected by centrifugation and pooled, concentrated, and stored at 4 °C. Reconstitution conditions were optimized by small-scale titration and nucleosomes checked by 6% native PAGE.

Preparation and purification of MH-flanked origin DNA. Short 168-bp MH-flanked origins. The conjugation of MH with origin substrates was performed in 50 mM Tris-HCl pH 8.0, 1 mM EDTA and 0.5 mM 2-mercaptoethanol supplemented with 100 µM des-adenosylmethionine (NEB). The reaction was carried out overnight at 30 °C, with a 10:1 molar ratio of MH:DNA. After conjugation, reactions were centrifuged at 14,680 rpm for 5 min and loaded onto a 1 ml RESOURCE-Q column equilibrated into DNA buffer (50 mM Tris-HCl pH 8.0 and 5 mM 2-mercaptoethanol). MH-conjugated DNA was eluted in a linear gradient of DNA buffer B (50 mM Tris-HCl pH 8.0, 5 mM 2-mercaptoethanol and 2 mM NaCl) over 24 column volumes. Fractions containing MH-conjugated DNA were pooled, concentrated and stored at ~80 °C. Conjugations were checked by 6% native PAGE.

6× ARS1-MH-flanked array. The conjugation of MH with origin substrates was performed in 25 mM Tris-HCl pH 7.5, 10 mM magnesium acetate, 50 mM potassium acetate and 1 mg ml⁻¹ BSA supplemented with 150 µM 5-adenosylmethionine (NEB). The reaction was carried out at 32 °C for 1 h then overnight at 4 °C, with a 20:1 molar ratio of MH:DNA. After
In vitro CMG assembly on short chromatinized origins

ARS1 nucleosome-flanked origin DNA (20 nM) was incubated with 52 nM ORC, 52 nM Cdc6 and 110 nM Mcm2–7–Cdt1 for 30 min at 24 °C in loading buffer (25 mM HEPES-KOH pH 7.6, 100 mM potassium glutamate, 10 mM magnesium acetate, 0.02% NP-40 and 0.5 mM TCEP) + 5 mM ATP. The reaction was supplemented with 80 nM DDK, and incubation continued for a further 10 min at 24 °C. Nucleoprotein complexes were isolated by incubation with 5 µl MagStrep type3 XT beads (IBA) pre-washed in 1× loading buffer for 30 min at 24 °C. The beads were washed three times with 100 µl wash buffer (25 mM HEPES-KOH pH 7.6, 105 mM potassium glutamate, 5 mM magnesium acetate, 0.02% NP-40 and 500 mM NaCl) and once with 100 µl loading buffer. Loaded, phosphorylated double hexamers were eluted in 20 µl elution buffer (25 mM HEPES-KOH pH 7.6, 105 mM potassium glutamate, 10 mM magnesium acetate, 0.02% NP-40 and 0.5 mM TCEP) + 5 mM ATP. The reaction was supplemented with 80 nM DDK, and incubation continued for a further 10 min at 24 °C. A mix of firing factors was then added to a final concentration of 30 nM Dpb11, 300 nM GINS, 240 nM Cdc45, 60 nM Pol ε, 30 nM Dpb11, 20 nM GINS, 50 nM Cdc45, 20 nM CDK, 50 nM RPA, 10 nM Topol, 100 nM Pol α, 25 nM Sl3–Sl7, 10 nM Mcm10 and 50 nM Sl2d. After 30 min of incubation, the reaction was applied directly to grids or diluted fivefold in 1× loading buffer for ReconSil experiments.

In vitro CMG assembly on 6× ARS1 MH-capped array

MH-capped ARS1 array DNA (5 nM) was incubated with 52 nM ORC, 52 nM Cdc6 and 110 nM Mcm2–7–Cdt1 for 30 min at 24 °C in loading buffer (25 mM HEPES-KOH pH 7.6, 100 mM potassium glutamate, 10 mM magnesium acetate, 0.02% NP-40 and 0.5 mM TCEP) + 5 mM ATP. The reaction was supplemented with 80 nM DDK, and incubation continued for a further 10 min at 24 °C. A mix of firing factors was then added to a final concentration of 30 nM Dpb11, 100 nM GINS, 80 nM Cdc45, 20 nM Pol ε, 30 nM Sl3–Sl7 and 50 nM Sl2d. After 30 min of incubation, the reaction was applied directly to grids or diluted fivefold in 1× loading buffer for ReconSil experiments.

DNA topology assay

The experiment was performed as described previously. The concentrations of proteins were as follows: 10 nM ORC, 50 nM Cdc6, 100 nM Mcm2–7–Cdt1 (or Mcm mutants), 80 nM DDK for the helicase loading step (5 µl) and 20 nM Pol ε, 30 nM Dpb11, 40 nM GINS, 50 nM Cdc45, 30 nM Cdc6, 10 nM Topol, 25 nM Sl3–Sl7, 5 nM Mcm10, 50 nM Sl2d for the helicase activation step (10 µl). Radiolabelled 616-bp circular DNA (25 fmol) was used. After processing the reactions as described previously, Ficoll 400 (final concentration was 2.5%) and Orange G were used to load the sample onto a native 3.5% bis-polyacrylamide gel (1× TBE) and separation was carried out for 21 h at 90 V using Protean II XL Cell apparatus (Bio-Rad) at room temperature. The 0.7-mm gel was dried (without fixation) at 80 °C for 10 min, exposed to a phosphor screen and scanned with the use of Typhoon phosphor imager.

Sample preparation and data collection for NS-EM

NS-EM sample preparation was performed on 400-mesh copper grids with carbon film (Agar Scientific). Grids were glow-discharged for 30 s at 45 mA using a K100X glow discharge unit (Electron Microscopy Sciences) before a 4-µl sample was applied to the grids and incubated for 2 min. Grids were stained by two successive applications of 4 µl 2% (w/v) uranyl acetate with blotting between the first and second application. Stained grids were blotted after 20 s to remove excess stain. Unless described otherwise, data collection was carried out on a Tecnai LaB6 G2 Spirit transmission electron microscope (FEI) operating at 120 kV. A 2K × 2K GATAN Ultrascan 100 camera was used to collect micrographs at a nominal magnification of 30,000 (with a physical pixel size of 3.45 Å per pixel) within a −0.5 to −2.0 µm defocus range.

NS-EM image processing

A subset of particles was manually picked using RELION-3.1 and used as a training dataset for Topaz training. Subsequent image processing was performed using RELION-3.1. The CTF of each micrograph was estimated using Gctf and particles...
were extracted and subjected to reference-free 2D classification in RELION-3.1.

ReconSil image processing
For ReconSil experiments, image processing was carried out as detailed above. Reference-free 2D classification in RELION generates both 2D class averages and star files detailing the class assignment, particle coordinates and transformations (translations and rotations) applied to the raw particles for alignment. 2D averages are superposed on the raw micrographs, overlaid on the particles that contributed to their generation. This yielded signal-enhanced ReconSiled micrographs reconstituting the context of complete origins of replication. ReconSiled micrographs were used for the selection and rejection of origin nucleoproteins for further analysis.

ReconSil data analysis and statistics
ReconSiled origins were analysed as previously described24. In brief, ReconSiled micrographs were used to re-extract particles of interest in RELION. Selected particles were manually classified for statistical analysis. Measurements of ReconSiled origins were performed manually using Fiji25 and plotted in GraphPad Prism v.9.2.0.

Sample preparation and data collection for cryo-EM
CMG assembly reactions (reconstituted as described in 'In vitro CMG assembly on short chromatinized origins') were frozen on 400-mesh lacey grids with a layer of ultra-thin carbon (Agar Scientific). All grids were freshly glow-discharged for 1 min at 45 mA using a K100X glow discharge unit (Electron Microscopy Sciences) before plunge-freezing. Samples were prepared by applying 4 μl of undiluted CMG assembly reactions for 2 min on a grid equilibrated to 25 °C in 90% humidity. The grid was blotted for 4.5 s and plunged into liquid ethane. Data collection was performed on an in-house Thermo Fisher Scientific Titan Krios transmission electron microscope operated at 300 kV, equipped with a Gatan K2 direct electron detector camera (Gatan) and a GIF Quantum energy filter (Gatan). Images were collected automatically using the EPU software (Thermo Fisher Scientific) in counting mode with a physical pixel size of 1.08 Å per pixel, with a total electron dose of 51.4 electrons per Å² during a total exposure time of 10 s dose-fractionated into 32 movie frames (Extended Data Table 1). We used a slit width of 20 eV on the energy filter and a defocus range of −2.0 to −4.4 μm. A grid was blotted for 4.5 s and plunged into liquid ethane. Data collection was performed on an in-house Thermo Fisher Scientific Titan Krios transmission electron microscope operated at 300 kV, equipped with a Gatan K2 direct electron detector camera (Gatan) and a GIF Quantum energy filter (Gatan). Images were collected automatically using the EPU software (Thermo Fisher Scientific) in counting mode with a physical pixel size of 1.08 Å per pixel, with a total electron dose of 51.4 electrons per Å² during a total exposure time of 10 s dose-fractionated into 32 movie frames (Extended Data Table 1). We used a slit width of 20 eV on the energy filter and a defocus range of −2.0 to −4.4 μm. A total of 65,286 micrographs were collected from two separate sessions.

Cryo-EM image processing
Data processing was performed using RELION-3.1 (ref. 26) and cryoSPARC v.3.2 (ref. 27) (Extended Data Fig. 3). The movies for each micrograph were first corrected for drift and dose-weighted using MotionCorr2 (ref. 28). CTF parameters were estimated for the drift-corrected micrographs using Gctf within RELION-3.1 (ref. 26). Dataset one was first processed separately and combined with dataset two at a later stage.

For the first dataset, particles were picked using a manually curated particle set as a template in cryLOLO v.1.7.5 (ref. 53). All grids were freshly glow-discharged for 1 min at 45 mA using a K100X glow discharge unit (Electron Microscopy Sciences) before plunge-freezing. Samples were prepared by applying 4 μl of undiluted CMG assembly reactions for 2 min on a grid equilibrated to 25 °C in 90% humidity. The grid was blotted for 4.5 s and plunged into liquid ethane. Data collection was performed on an in-house Thermo Fisher Scientific Titan Krios transmission electron microscope operated at 300 kV, equipped with a Gatan K2 direct electron detector camera (Gatan) and a GIF Quantum energy filter (Gatan). Images were collected automatically using the EPU software (Thermo Fisher Scientific) in counting mode with a physical pixel size of 1.08 Å per pixel, with a total electron dose of 51.4 electrons per Å² during a total exposure time of 10 s dose-fractionated into 32 movie frames (Extended Data Table 1). We used a slit width of 20 eV on the energy filter and a defocus range of −2.0 to −4.4 μm. A total of 65,286 micrographs were collected from two separate sessions.

Model building and refinement
CMG (from PDB 6SKL)31, Pol2 subunit (from PDB 6HV9)33 and a homology model of the N-terminal domain of Dpb2 obtained from the Phyre2 server44 were docked initially into the cryo-EM map produced from Resolve CryoEM, using UCSF Chimera, and refined against the map using Coot v.0.9.1 (ref. 45). The DNA and the MCM5 winged helix domain were built de novo. The register of origin DNA engagement was performed using Gctf within RELION-3.1 of the centred single CMGE (consisting of 358,622 particles) was refined to 4.2-Å resolution. These particles were subjected to several rounds of CTF refinement and two rounds of Bayesian polishing. After this, CTF-refined and polished particles were refined with local searches in C1 with a mask encompassing the entire CMGE density to 3.6-Å resolution. To better resolve the DNA inside the MCM central channel, densities corresponding to Cdc45, GINS and Pol e were subtracted in RELION. Signal-subtracted particles were analysed by 3D variability analysis in cryoSPARC (ref. 56). A subset of 71,348 particles was selected based on the quality of DNA density. These signal-subtracted particles were subsequently reverted to the original particles and refined using local searches in C1 using local searches to 3.5-Å resolution.

All refinements were performed using fully independent data half-sets and resolutions are reported based on the Fourier shell correlation (FSC) = 0.143 criterion (Extended Data Fig. 2). FSCs were calculated with a soft mask. Maps were corrected for the modulation transfer function of the detector and sharpened by applying a negative B-factor as determined by the post-processing function of RELION or in cryoSPARC. The final RELION half-maps were used to produce a density modified map using the PHENIX Resolve CryoEM (refs. 28,59). This 3.4-Å map showed significant improvements for side chain and DNA density as well as for overall interpretability. Local-resolution estimates were determined using PHENIX or cryoSPARC (Extended Data Fig. 2f,j). The conversions between cryoSPARC and RELION files were performed using the UCSF pyem v.0.5 package60.

Map and model visualization
Maps were visualized in UCSF Chimera66 and ChimeraX67 and all model illustrations and morphs were prepared using ChimeraX or PyMOL.
Statistics and reproducibility
Statistical analysis was performed using a two-tailed Welch’s t-test in GraphPad Prism v.9.2.0. No statistical methods were used to pre-determine sample size. The experiments were not randomized, and investigators were not blinded to allocation during experiments and outcome assessment.

Reporting summary
Further information on research design is available in the Nature Research Reporting Summary linked to this paper.

Data availability
Data supporting the findings of this study are available within the paper and its Supplementary Information files. Cryo-EM density maps of the CMGE dimer complex have been deposited in the Electron Microscopy Data Bank (EMDB) under the accession code EMD-13988. The cryo-EM density map of the symmetry-expanded CMGE monomer has been deposited in the EMDB under the accession code EMD-13978. Atomic coordinates have been deposited in the PDB with the accession codes 7QHS (symmetry-expanded CMGE monomer) and 7Z13 (monomer docked into the CMGE dimer map).

62. Kidmose, R. T. et al. Namdinator – automatic molecular dynamics flexible fitting of structural models into cryo-EM and crystallography experimental maps. IUCrJ 6, 526–531 (2018).
63. Emsley, P., Lohkamp, B., Scott, W. G. & Cowtan, K. Features and development of Coot. Acta Crystallogr. D Biol. Crystallogr. 66, 486–501 (2010).
64. Adams, P. D. et al. PHENIX: a comprehensive Python-based system for macromolecular structure solution. Acta Crystallogr. D Biol. Crystallogr. 66, 213–221 (2010).
65. Chen, V. B. et al. MolProbity: all-atom structure validation for macromolecular crystallography. Acta Crystallogr. D Biol. Crystallogr. 66, 12–21 (2010).
66. Pettersen, E. F. et al. UCSF Chimera—a visualization system for exploratory research and analysis. J. Comput. Chem. 25, 1605–1612 (2004).
67. Goddard, T. D. et al. UCSF ChimeraX: meeting modern challenges in visualization and analysis. Protein Sci. 27, 14–25 (2018).

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Author contributions
J.S.L. and A.C. conceived the study. J.S.L. and M.H.G. designed biochemistry experiments. J.S.L., S.S.H. and J.F.G. prepared biochemical reagents and developed all assays, with the exception of the DNA replication and topology experiments that were performed by M.H.G., who also produced all MCM mutants under the supervision of J.F.X.D. J.S.L. conducted all image processing. J.S.L. performed the atomic model building and validation with J.S. J.F.X.D. contributed critical reagents and protocols. A.C. supervised the study. J.S.L. and A.C. wrote the manuscript with input from M.H.G., J.F.X.D. and the other authors.

Competing interests
The authors declare no competing interests.

Additional information
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Extended Data Fig. 1 | Origin-dependent CMG assembly with purified proteins visualized by electron microscopy. a. Purified MCM loading and firing factors (left), and additional factors required for DNA substrate preparation (right) analysed by SDS–PAGE with Coomassie staining. For gel source data, see Supplementary Fig. 1. Similar results were observed for at least two independent sample preparations. b. 6% PAGE gel of capped, origin DNA substrates used in this study. (Below) Cartoon representation of ARS1 origins of replication, containing the two inverted ORC-binding sites, ACS (high affinity, red arrow) and B2 (low affinity, orange arrow). ARS1 is flanked by nucleosomes (NCP) or covalently linked methyltransferases (MH). For gel source data, see Supplementary Fig. 1. Similar results were observed for three independent sample preparations. c. Representative NS micrograph and 2D averages of entire CMG assembly reactions used to generate the ReconSiled origins shown in Fig. 1b. 70% of CMG particles exist in a dimeric (dCMGE) trans configuration (light orange), with GINS positioned on opposed sides of MCM. 11% of dCMGE particles exist in a cis configuration (dark orange) that might derive from trans-dCMGE disengagement and rotation. This experiment was performed more than three times. d. Cartoon representation of 6x ARS1 array, containing the two inverted ORC-binding sites, ACS (high affinity, red arrow) and B2 (low affinity, orange arrow). Each ARS1 origin is separated by 40 bp linker DNA. Array is flanked by covalently attached MH. Blue arrows indicate MseI cut sites. e. Reaction scheme for CMG assembly reactions on DNA substrates containing a 6x ARS1 array. f. Representative NS micrograph and representative double hexamer and dCMGE 2D averages obtained from CMG assembly reaction on 6x ARS1 array. This experiment was performed three times. g. 6% PAGE gel of partial DNA digestion of 6x ARS1 array by MseI carried out under the same conditions as NS-EM experiments. Lane 1 contains unmodified 6x ARS1 array. Lane 2 contains MH-conjugated 6x ARS1 array DNA. Lane 3 contains MseI digested 6x ARS1 array DNA. For gel source data, see Supplementary Fig. 1. This experiment was performed twice.
Extended Data Fig. 2 | Sample preparation and validation of dCMGE cryo-EM reconstructions. a. Schematic of biochemical reconstitution used for cryo-EM samples. b. Representative cryo-EM micrograph of entire dCMGE assembly reaction with particles highlighted with white circles. Cryo-EM sample preparation was performed once; similar results were observed in at least three independent NS sample preparations. 
b-c. Representative 2D class averages from NS-EM (left panel) and cryo-EM imaging (right panel). 
d. Fourier shell correlation plot for the C1-refined CMGE dimer map. e. Angular distribution plot for the C1-refined CMGE dimer map. f. CryoSPARC local-resolution estimate for the C1-refined CMGE dimer map. g. Fourier shell correlation plot for symmetry-expanded CMGE map used in model building. h. Angular distribution plot for the symmetry-expanded CMGE map used in model building. i. Model-to-map correlation graph. j. PHENIX local-resolution estimate for the symmetry-expanded CMGE map. Box widths represent 500 Å.
Extended Data Fig. 3 | Cryo-EM data processing pipeline for dCMGE assembled on nucleosome-capped origin DNA. a. Schematic shows the classification and refinement steps taken to achieve the symmetry-expanded refined CMGE cryo-EM structure. The software used during each processing step is listed. b. Schematic of the classification and refinement pipeline for the C1-refined CMGE dimer. c. Symmetry-expanded refined CMGE structure (coloured) docked into C1-refined CMGE dimer map (grey). Reported resolutions in all schematics are calculated based on the FSC = 0.143 criterion.
Extended Data Fig. 4 | Quality of cryo-EM densities. Example cryo-EM density of Mcm2 (a), Mcm4 (b), Mcm7 (c), Pol2 (d), Sld5 (e), Dpb2 (f), and Cdc45 (g). For each subunit, side chain density features are shown in the inset.
Extended Data Fig. 5 | Analysis of protein–DNA interactions within the dCMGE complex. a. MCM contacts with DNA in dCMGE. Phosphate backbone interactions are indicated with black arrows and base interactions are highlighted with blue arrows. Residues are coloured based on individual MCM subunits. b. Comparison of N-terminal domains of Mcm2–7. The N-terminal pore loops are highlighted in red. Mcm6 contains a unique insertion (‘wedge’) with residues that stabilize the orphan bases exposed upon DNA untwisting. c. Cryo-EM density of flexible Mcm6 wedge stabilizing three lagging-strand bases. d. Representative cryo-EM densities of DNA contacts for Mcm3 and Mcm6. e. Sequence alignment of N-terminal region of Mcm6. Conserved T423 and R424 are outlined with black boxes. f. Sequence alignment between the h2i and PS1BH motifs of yeast Mcm2 compared to selected higher eukaryotes. Mcm2 6A mutations are outlined with black boxes. All alignments are coloured using the ClustalX colouring scheme.
Extended Data Fig. 6 | Supporting information for the mechanism of DNA-bubble nucleation. a. Overview of the Mcm2 6A mutant in the context of ATPase domain. Residues that push the lagging-strand template are coloured gold and residues that pull the leading-strand template are coloured in grey.

b. Reaction scheme for topology footprint assay for DNA unwinding. c. Over-exposed gel as seen in Fig. 4d. For gel source data, see Supplementary Fig. 1. This experiment was performed twice.
Extended Data Fig. 7 | dCMGE sterically impedes the docking of the E3 ligase onto MCM. When CMGE–SCF\textsuperscript{D\alpha} (7PMK) is superposed onto the double hexamer (7P30) and the dCMGE structure (this study), major clashes can be identified between SCF\textsuperscript{D\alpha} engaged to one ring and the Mcm3 subunit from the opposed ring in the CMG dimer. This clash explains why the CMG assembled around the origin DNA duplex during initiation is protected from disassembly before lagging-strand ejection\textsuperscript{22}. 
Extended Data Fig. 8 | Schematic representation of the steps that lead to replication origin firing. From top to bottom: double hexamer is loaded onto duplex origin DNA in an ATP-hydrolysis dependent manner. ADP formed during double hexamer assembly remains bound to MCM. The Mcm6 wedge insertion forms part of the double hexamer dimerization interface (inset). After loading the double hexamer makes limited contacts with duplex DNA. DDK phosphorylation triggers the recruitment of firing factors that deposit Cdc45 (C in CMGE) and GINS (G) onto the MCM (M), in the context of the so-called pre-initiation complex, the formation of which requires Pol ε (E) and CDK kinase activity. ADP release and binding of new ATP by MCM triggers CMGE assembly. CMGE assembly leads to the reconfiguration of the double hexamer interface, resulting in hexamer separation and a 1 subunit register shift pivoting around the Mcm6 N-terminal domain. This movement results in the exposure of 1.5 turns of duplex DNA between two MCM rings, and nucleation of DNA melting within each MCM ring. DNA melting is promoted upon ATP-triggered untwisting of 0.7 turns of the DNA, through the action of Mcm2 (pink in the inset), which pushes on the lagging strand while simultaneously pulling on the leading strand. As DNA is untwisted, Watson and Crick base pairs are broken, and three orphan bases become stabilized by the Mcm6 wedge (orange in the inset), which moves from the double hexamer interface and invades the MCM central channel. Action by Mcm10 triggers ATP hydrolysis by CMG and ejection of the lagging strand through an unknown mechanism, resulting in helicase bypass and establishment of replication forks.
### Extended Data Table 1 | Cryo-EM data collection, refinement and validation statistics

|                        | Symmetry expanded CMGE (EMDB-13978) | CMGE dimer (EMD-13988) |
|------------------------|-------------------------------------|------------------------|
| **Data collection and processing** |                                     |                        |
| Magnification          | 130,000x                            | 130,000x               |
| Voltage (kV)           | 300                                 | 300                    |
| Electron exposure (e-/Å²) | 51.4                               | 51.4                   |
| Defocus range (µm)     | 2.0–4.4                             | 2.0–4.4                |
| Pixel size (Å)         | 1.08                                | 1.08 (collection) 1.296 (processing) |
| Symmetry imposed       | C1                                  | C1                     |
| Initial particle images (no.) | 927,109                        | 133,262                |
| Final particle images (no.) | 71,348                      | 49,766                 |
| Map resolution (Å)     | RELION 3.50                        | CryoSPARC 8.20         |
|                        | FSC threshold 0.143                 | 0.143                  |
|                        | Resolve Cryo-EM 3.40                |                         |
|                        | FSC threshold 0.5                   | 0.5                    |
| Map resolution range (Å) | 2.6–6.6                           | 8.0–12.1               |
| **Refinement**         |                                     |                        |
| Initial model used (PDB code) | 6SKL                              | 6HV9                   |
| Model resolution (Å)   | 3.30                                |                         |
| FSC threshold = 0.5    | Resolve Cryo-EM (−10)              |                         |
| **Model composition**  |                                     |                        |
| Non-hydrogen atoms     | 53,679                              |                         |
| Protein residues       | 6,564                               |                         |
| Nucleotides            | 52                                  |                         |
| Ligands                | ZN: 7                               |                         |
|                        | MG: 3                               |                         |
|                        | ATP: 4                              |                         |
|                        | ADP: 2                              |                         |
| **B factors (Å²)**     |                                     |                        |
| Protein                | 38.81                               |                         |
| Nucleotide             | 128.08                              |                         |
| Ligand                 | 40.25                               |                         |
| **R.m.s. deviations**  |                                     |                        |
| Bond lengths (Å)       | 0.003                               |                         |
| Bond angles (°)        | 0.645                               |                         |
| **Validation**         |                                     |                        |
| MolProbity score       | 1.65                                |                         |
| Clashscore             | 7.14                                |                         |
| Poor rotamers (%)      | 0.03                                |                         |
| **Ramachandran plot**  |                                     |                        |
| Favored (%)            | 96.20                               |                         |
| Allowed (%)            | 3.80                                |                         |
| Disallowed (%)         | 0.00                                |                         |
Reporting Summary

Nature Portfolio wishes to improve the reproducibility of the work that we publish. This form provides structure for consistency and transparency in reporting. For further information on Nature Portfolio policies, see our Editorial Policies and the Editorial Policy Checklist.

Statistics

For all statistical analyses, confirm that the following items are present in the figure legend, table legend, main text, or Methods section.

- The exact sample size (n) for each experimental group/condition, given as a discrete number and unit of measurement
- A statement on whether measurements were taken from distinct samples or whether the same sample was measured repeatedly
- The statistical test(s) used AND whether they are one- or two-sided
- Only common tests should be described solely by name; describe more complex techniques in the Methods section.
- A description of all covariates tested
- A description of any assumptions or corrections, such as tests of normality and adjustment for multiple comparisons
- A full description of the statistical parameters including central tendency (e.g. means) or other basic estimates (e.g. regression coefficient) AND variation (e.g. standard deviation) or associated estimates of uncertainty (e.g. confidence intervals)
- For null hypothesis testing, the test statistic (e.g. f, t, r) with confidence intervals, effect sizes, degrees of freedom and P value noted. Give P values as exact values whenever suitable.
- For Bayesian analysis, information on the choice of priors and Markov chain Monte Carlo settings
- For hierarchical and complex designs, identification of the appropriate level for tests and full reporting of outcomes
- Estimates of effect sizes (e.g. Cohen’s d, Pearson’s r), indicating how they were calculated

Our web collection on statistics for biologists contains articles on many of the points above.

Software and code

Policy information about availability of computer code

Data collection: Gatan Digital Micrograph and ThermoFisher EPU v2.9

Data analysis: cryOLO v1.7.5, Topaz v2.0.5, MotionCor2, Gctf v1.06, RELION v3.1, cryoSPARC 3.2, UCSF Chimera v1.14, ChimeraX 1.3, COOT v0.9-pre, Phenix v1.19.2, MolProbity web server, ImageJ v2.0.0, GraphPad Prism v9.2.0, Phyre2 web server, pymol v0.5, PyMOL v2.4.1

For manuscripts utilizing custom algorithms or software that are central to the research but not yet described in published literature, software must be made available to editors and reviewers. We strongly encourage code deposition in a community repository (e.g. GitHub). See the Nature Portfolio guidelines for submitting code & software for further information.

Data

Policy information about availability of data

All manuscripts must include a data availability statement. This statement should provide the following information, where applicable:

- Accession codes, unique identifiers, or web links for publicly available datasets
- A description of any restrictions on data availability
- For clinical datasets or third party data, please ensure that the statement adheres to our policy

Cryo-EM density maps of the CMGE dimer complex has been deposited in the Electron Microscopy Data Bank (EMDB) under the accession number EMD-13988. Cryo-EM density map of the symmetry expanded CMGE monomer has been deposited in the EMDB under the accession number EMD-13978. Atomic coordinates have been deposited in the Protein Data Bank (PDB) with the accession number 7QHS (symmetry expanded CMGE monomer) and 7Z13 (monomer docked into the CMGE dimer map).
Field-specific reporting

Please select the one below that is the best fit for your research. If you are not sure, read the appropriate sections before making your selection.

- [x] Life sciences
- [ ] Behavioural & social sciences
- [ ] Ecological, evolutionary & environmental sciences

For a reference copy of the document with all sections, see nature.com/documents/nr-reporting-summary-flat.pdf

Life sciences study design

All studies must disclose on these points even when the disclosure is negative.

Sample size

In our negative stain EM experiments, we imaged ATP-dependent CMG assembly, yielding different reaction intermediates. To isolate CMG dimers, we usually collected 100-300 micrographs per condition, these numbers of micrographs were sufficient to either allow 2D classification or comparative analysis between MCM mutants.

To obtain high-resolution structure of the CMGE nucleating origin DNA melting from the mixed population of reaction intermediates in the cryo-EM experiment, ~65.3 K micrographs were collected from two independent grids made from the same CMG assembly reaction. This number of micrographs was sufficient to either allow model building or comparative analysis.

No statistical methods were used to predetermine sample size.

Data exclusions

For our ReconstI experiments, samples were prepared with reduced concentrations to limit particle crowding and allow the clear identification of single origins of replication. Particles were picked and multiple rounds of 2D classification were performed to isolate particles contributing to the distinct molecular species in our samples. Picked particles that could not be aligned and classified were discarded and therefore were not reconstructed in silico. ReconSted origins were evaluated and rejected if confident assignment of co-localisation to the same origin could not be made because either, i. the origin was in a region of clustered/aggregated particles or ii. If the origin contained additional particles that had not been 2D classified, and were therefore not overlaid with a 2D class average that would permit confident assignment of the molecular species.

Negative stain and cryo-EM micrographs with poor staining, ice contamination or entirely lacy carbon, respectively, were excluded. Picked particles that did not align to a distinct class in 2D and 3D (cryo-EM only) were excluded from further analysis. CMGs that were not engaged with epsilon were removed from the cryo-EM dataset to yield the best reconstruction of the CMGE dimer complex.

Replication

The cryo-EM dataset of ATP-dependent CMG assembly reaction consisted of a single reaction and two datasets, collected on two independent grids. CMG dimer complex formation in negative stain EM experiments was found to be reproducible across multiple independent sample preparations using different DNA substrates. Details of the number of experimental repeats have been acknowledged in the relevant figure legends. All attempts at data replication were successful. Details of the number of experimental repeats have been acknowledged in the relevant figure legends. All attempts at data replication were successful.

Randomization

For calculation of the resolution of the cryo-EM reconstructions, Fourier shell correlations were calculated using independent halves of the complete datasets, into which the component particles were segregated randomly.

Blinding

Blinding is not relevant for a single particle electron microscopy study such as this.

Reporting for specific materials, systems and methods

We require information from authors about some types of materials, experimental systems and methods used in many studies. Here, indicate whether each material, system or method listed is relevant to your study. If you are not sure if a list item applies to your research, read the appropriate section before selecting a response.

**Materials & experimental systems**

| n/a | Involved in the study |
|-----|-----------------------|
| [x] | Antibodies            |
| [ ] | Eukaryotic cell lines |
| [ ] | Palaeontology and archaeology |
| [ ] | Animals and other organisms |
| [ ] | Human research participants |
| [ ] | Clinical data         |
| [ ] | Dual use research of concern |

**Methods**

| n/a | Involved in the study |
|-----|-----------------------|
| [x] | ChIP-seq               |
| [ ] | Flow cytometry         |
| [ ] | MRI-based neuroimaging |

**Eukaryotic cell lines**

Policy information about [cell lines](#)

**Cell line source(s)**

S. cerevisiae overexpression strains for CMG assembly and DNA replication proteins have previously been described in [in]
| Cell line source(s)                                                                 | multiple studies across several publications. For clarity to the potential readers and reviewers we have included extensive details in extended data table 4. |
|-----------------------------------------------------------------------------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| Authentication                                                                    | *S. cerevisiae* overexpression strains were checked for correct plasmid integration by PCR amplification from extracted genomic DNA.                                                                 |
| Mycoplasma contamination                                                           | *S. cerevisiae* overexpression strains were not tested for mycoplasma contamination.                                                                                                             |
| Commonly misidentified lines (See [ICLC register](https://www.iclc.org))          | No commonly misidentified cell lines were used in this study.                                                                                                                                   |