

Structural and mechanistic insights into the MCM8/9 helicase complex

Zhuangfeng Weng, Jiefu Zheng, Yiyi Zhou, Zuer Lu, Yixi Wu, Dongyi Xu, Huanhuan Li, Huanhuan Liang , Yingfang Liu 

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Shenzhen Key Laboratory for Systems Medicine in Inflammatory Diseases, School of Medicine, Shenzhen Campus of Sun Yat-sen University, Shenzhen, Guangdong 518107, China • State Key Laboratory of Protein and Plant Gene Research, School of Life Sciences, Peking University, Beijing, China • Department of Colorectal Surgery, The Sixth Affiliated Hospital, Sun Yat-sen University, Guangdong Institute of Gastroenterology, Guangdong Provincial Key Laboratory of Colorectal and Pelvic Floor Diseases, Guangzhou, Guangdong 510655, China • Pharmaceutical Sciences (Shenzhen), Sun Yat-sen University, Shenzhen, Guangdong 518107, China

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Abstract

MCM8 and MCM9 form a functional helicase complex (MCM8/9) that plays an essential role in DNA homologous recombination repair for DNA double-strand break. However, the structural characterization of MCM8/9 for DNA binding/unwinding remains unclear. Here, we report structures of the MCM8/9 complex using cryo-electron microscopy single particle analysis. The structures reveal that MCM8/9 is arranged into a heterohexamers through a three-fold symmetry axis, creating a central channel that accommodates DNA. Multiple characteristic hairpins from the N-terminal oligosaccharide/oligonucleotide (OB) domains of MCM8/9 protrude into the central channel and serve to unwind the duplex DNA. When activated by HROB, the structure of MCM8/9's N-tier ring converts its symmetry from C_3 to C_1 with a conformational change that expands the MCM8/9's trimer interface. Moreover, our structural dynamic analyses revealed that the flexible C-tier ring exhibited rotary motions relative to the N-tier ring, which is required for the unwinding ability of MCM8/9. In summary, our structural and biochemistry study provide a basis for understanding the DNA unwinding mechanism of MCM8/9 helicase in homologous recombination.

eLife assessment

This paper presents **important** findings on the hexameric structure of MCM8/9, which potentially explain its role as a DNA helicase in homologous recombination. This **solid** work will be of interest to biologists studying DNA transactions.

Introduction

DNA helicases play critical roles in multiple cellular processes including DNA replication, transcription, recombination and repair. Dysfunction of multiple DNA helicases have been correlated with various human diseases including cancers, Bloom Syndrome, Werner Syndrome, Fanconi Anemia, reproductive deficiencies and infertility, *etc* (1 , 2 ). One of the most

extensively studied subfamilies of DNA helicases is the minichromosome maintenance (MCM) proteins. In the human genome, the MCM family contains two subgroups of AAA+ ATPase/helicase complexes, consisting of eight MCM members (MCM2-9) (3 [↗](#)). Structural studies of these complexes are critical for elucidating their mechanisms.

The hexameric MCM2-7 complex, also known as a replicative helicase, plays a vital role in the initiation and elongation of eukaryotic chromosome replication. Previous structural and biochemical studies have revealed that the MCM2-7 complex exhibits high structural flexibility during DNA replication (4 [↗](#)). It can adopt multiple conformations by interacting with its loaders and activators to form different intermediates, including the helicase-loading intermediate Orc-Cdc6-Cdt1-MCM2-7 (OCCM), the inactive MCM2-7 Double Hexamer (DH) and the active replicative helicase complex Cdc45-MCM2-7-GINS (CMG) (5 [↗](#)–8 [↗](#)). For loading onto DNA, the apo MCM2-7 hexamer first binds to the licensing factor Cdt1. In the presence of ATP, the initiators ORC and Cdc6 assemble on the origin DNA to form an active loading platform and load the Cdt1-bound MCM2-7 onto DNA to form the OCCM complex (4 [↗](#)). The origin DNA is encircled into the central channel of the MCM2-7 hexamer via the MCM2-MCM5 entry “gate” (9 [↗](#)). While ATP hydrolysis occurs in OCCM, it seems that Cdt1, Cdc6 and ORC are released in sequential disassembly steps, and then the first loaded MCM2-7 encircling DNA, recruits a second ORC-Cdc6 complex, which in turn loads the second Cdt1-bound MCM2-7 to form a head-to-head MCM2-7 DH (4 [↗](#)). During double-hexamer assembly, the N-tier ring of MCM2-7 rotates by nearly 30° when aligned with the MCM2-7 hexamer in its loading intermediate OCCM (10 [↗](#)). As the MCM2-7 double hexamer on DNA is inactive, lots of regulators along with the helicase activators Cdc45 and GINS complex, are required to convert the double hexamer into two activated CMG replicative helicases that translocate along the leading strands with a bi-directional replication mechanism (11 [↗](#)–13 [↗](#)). The aforementioned structural studies have also revealed that layers of characteristic hairpin loops inside the chamber of MCM2-7 hexamer, including the helix-2 insertion loops (H2I), the presensor-1 (PS1) as well as β -turn motifs of oligosaccharide/oligonucleotide (OB) domains, contribute to DNA binding and separation (14 [↗](#)). Additionally, another allosteric communication loop (ACL) conserved in all MCM helicases was also found to be important for helicase activity. The ACL mediates inter-subunit interactions between the N-terminal DNA processing domain and the C-terminal AAA+ motor domain (15 [↗](#), 16 [↗](#)).

MCM8 and MCM9 make up the other helicase complex, which is the homolog of MCM2-7 (17 [↗](#)). In contrast to the replicative helicase MCM2-7, MCM8 and MCM9 function as a DNA helicase complex (MCM8/9) in HR-mediated DNA repair for DNA double-strand breaks (DSBs) and DNA interstrand crosslinks (ICLs) (18 [↗](#)). Several studies have suggested that cells lacking *MCM8* and *MCM9* are highly sensitive to DNA cross-linking reagents (19 [↗](#), 20 [↗](#)). The *MCM8* and *MCM9* knock-out mice were found to be sterile and revealed gametogenesis deficiency as well as chromosomal damage due to impaired HR (18 [↗](#)). During HR repair, MCM8/9 was rapidly recruited to the DNA damage sites and colocalized with the recombinase Rad51 (21 [↗](#)). It also interacted with the nuclease complex MRN (MRE11-RAD50-NBS1), which was required for DNA resection at DSBs to facilitate the HR repair (22 [↗](#)). Recently, HROB (also known as C17orf53 or MCM8IP) has been identified as an essential factor in loading the MCM8/9 complex to the sites of DNA damage and stimulating its helicase activity to promote replication fork progression during DNA recombination and replication. Loss of HROB led to HR defects because it is necessary for the recruitment and activation of MCM8/9 (23 [↗](#), 24 [↗](#)). The HROB harbors a flexible N-terminal domain (NTD), a central proline-rich region (PRR) and an OB fold domain at the C-terminus. It was reported that the MCM8/9 binding motif (MBM) is embedded in the PRR region and binds to the NTD domains of MCM8/9 (24 [↗](#)). Although progress has been made in understanding the role of MCM8/9 HR during the past decade, the structure/function relationship, especially the structural characterization of MCM8/9 helicase activity remains unknown. Notably, MCM8/9 is composed of two subunits, MCM8 and MCM9. It is challenging to extrapolate its role through a simple comparison with MCM2-7, which consists of six subunits.

MCM8 and MCM9 proteins are structurally similar, with a MCM domain at the N-terminus (NTD) and an AAA+ ATPase domain at the C-terminus (CTD). The NTD and CTD are connected by flexible linkers (N-C linkers), which allow the movement of the AAA+ domain during unwinding (25 [↗](#)). The NTDs of MCM8 and MCM9 share an analogous domain organization and can be divided into three domains: the zinc finger domain (ZF), the helical domain (HD) and the OB domain. In addition to the AAA+ ATPase domain, MCM8 possesses an additional WHD domain in the CTD while MCM9 contains a putative helix-turn-helix (HTH) domain and an extensional C-tail at the C-terminus (26 [↗](#)).

In the previous study, we reported a 6.6 Å cryo-EM structure of human MCM8/9 (hMCM8/9) NTD ring. We also analyzed the crystal structures of MCM8 and MCM9 NTDs separately, which exhibited conformational changes when assembled into the MCM8/9 hexamer (27 [↗](#)). However, the low-resolution structure of the MCM8/9 NTD ring is insufficient to fully illustrate the assembly and activation mechanisms of the MCM8/9 hexamer. Here, we present a near-atomic resolution structure of the chicken MCM8/9 (gMCM8/9) complex and a 3.95 Å cryo-EM structure of the human MCM8/9 NTD ring under HROB induction. Based on these structures, we conducted further investigation on the structural features of the DNA-binding region of MCM8/9 and the conformational changes required for its activation by HROB. Our structural and biochemical studies shed light on the structural and functional relationship of the MCM8/9 helicase.

Results

Overall structure of the gMCM8/9 helicase complex

As our previous study shows the hMCM8/9 helicases complex is unstable for high-resolution structure determination, we first solved the cryo-EM structure of MCM8/9 complex from *Gallus gallus*. (*Gallus gallus*, gMCM8/9; amino acid, gMCM8: 50-830; gMCM9:1-691) (**Figure 1A** [↗](#), **Table 1** [↗](#), Materials and Methods). The gMCM8/9 shares up to 80% sequence homology with hMCM8/9. Based on the electron density map, the structures of the NTD and CTD were reconstructed at the resolutions of 3.66 Å and 5.21 Å, respectively, and then aligned into a composite structure (**Figure 1B**, **1C** [↗](#) and S1).

The structure reveals that gMCM8 and gMCM9 alternately assemble into a pinwheel-like hexamer with a central channel. The trimerization of the gMCM8/9 heterodimer is in line with our previously reported hMCM8/9 structure at 6.6 Å resolution (27 [↗](#)). When we performed structural superposition, the NTD ring of gMCM8/9 superimposes well onto that of hMCM8/9, indicating that eukaryotic MCM8 and MCM9 form a conserved hexamer complex during evolution (Figure S2, S3 and S4). The diameter of the NTD ring is ~132 Å and that of the inner channel is ~28 Å which is large enough to accommodate dsDNA to pass through (**Figure 1C** [↗](#) and S5). The assembly pattern of gMCM8/9 complex with a three-fold (C3) symmetry axis is unique among MCM helicases. It arranges the three MCM8 and three MCM9 in the same planes, enclosing a channel with multiple symmetrical hairpins. This assembly pattern differs from that of the eukaryotic MCM2-7 heterohexamer with non-symmetry and the archaeal MCM homohexamer with 6-fold symmetry (7 [↗](#), 28 [↗](#)). In addition, a distinguishing feature of the gMCM8/9 is that the helical domain of gMCM8 contains a long α5 helix that protrudes around the hexameric ring (**Figure 1B** [↗](#)).

The domain structures of gMCM8/9's, including the MCM domains, the AAA+ motor domains and the WH domain, fit well into their electron density maps (Figure S5). For each gMCM8/9 heterodimer, the positions of characteristic hairpin loops of MCM helicase, such as H2I, PS1, OB hairpins (referred to OB-hps) and the ACL were well defined. Among them, the OB-hps extend to the central channel, while the ACLs are close to the H2I and PS1 hairpins within the AAA+ domains (**Figure 1B** [↗](#)).

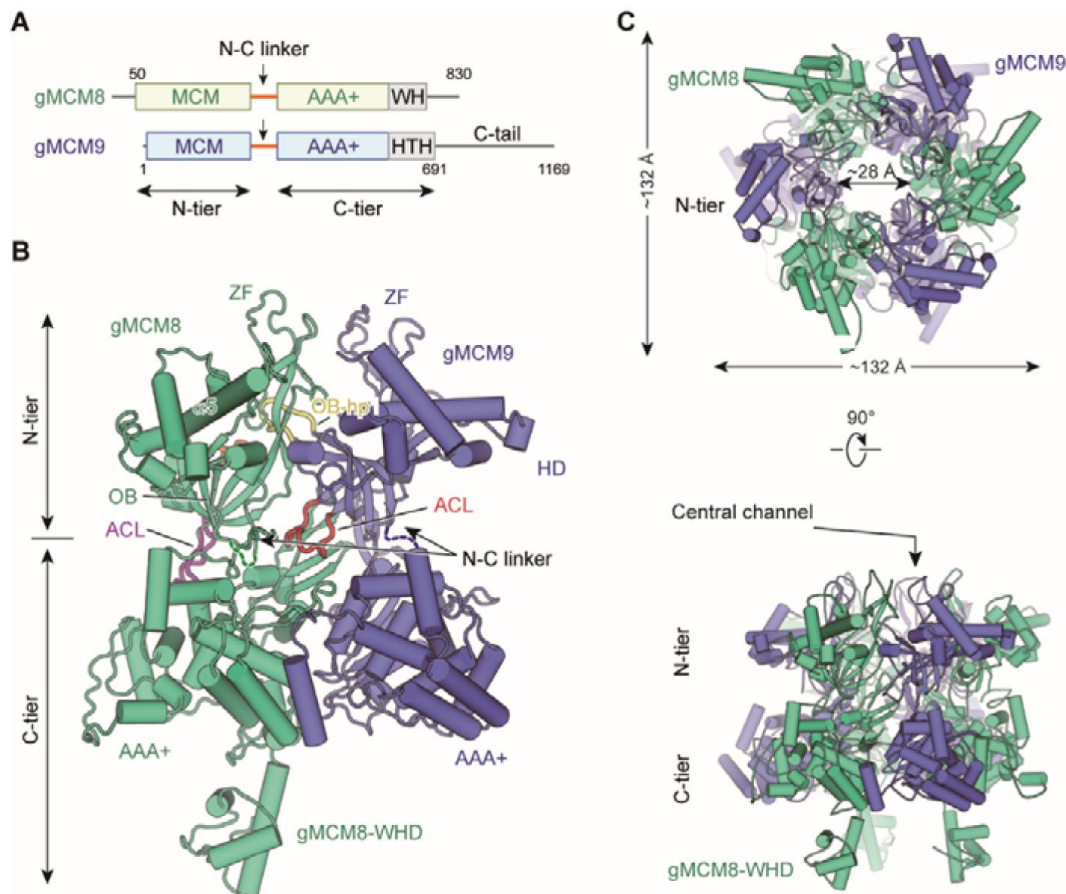


Figure 1.

Overall structural features of the gMCM8/9 hexamer.

(A) Domain organization of chicken MCM8 and MCM9. (MCM8, greencyan; MCM9, slate), N-C linker, linkers connecting the NTD and CTD of MCM8 or MCM9; WH, winged helix domain; HTH, helix-turn-helix; the C-tail domain is not included in this study. (B) Fold of the gMCM8/9 dimer. The N-C linkers are shown as dotted lines and the OB hairpins OB-hp are highlighted in orange (MCM8) and yellow (MCM9), while the ACL are highlighted in purple (MCM8) and red (MCM9). The long helix of MCM8 was marked as $\alpha 5$. (C) Ribbon diagram showing the top and side views of gMCM8/9 hexamer with three-fold symmetry axis.

Data collection and processing	NTD	CTD
Magnification		130,000
Voltage (kV)		200
Electron dose (e ⁻ /Å ²)		50
Frame		32
Under-focus range (μm)		1.7-2.2
Pixel size (Å)		1.0
Symmetry imposed		C3
FSC threshold		0.143
Relion processing		
Initial particle images (no.)		965k
Final particle images (no.)		144,202
Global map resolution (Å)		6.6
NTD map resolution (Å)	4.2	
CTD map resolution (Å)		5.4
Cryosparc processing		
Initial particle images (no.)		2,134k
Global map resolution (Å)		4.31
Final particle images (no.)	290,080	72,948
NTD map resolution (Å)	3.66	
CTD mean map resolution (Å)		5.21
Model composition		
Initial model used (PDB code)	7DP3 and 7DPD	3JA8
Non-hydrogen atoms	13410	12687
Protein	1698	1671
Ligands	0	0
R.m.s. deviations		
Bond lengths (Å)	0.002	0.003
Bond angles (°)	0.593	0.843
Validation		
MolProbity score	1.73	2.25
Clashscore	8.36	29
Rotamers outliers (%)	0.00	0.00
Ramachandran plot		
Favored (%)	95.64	95.61
Allowed (%)	4.3	4.33
Outliers (%)	0.06	0.06

Table 1.

Cryo-EM 3D reconstruction and refinement of the gMCM8/9 complex.

The assembly of gMCM8 and gMCM9 in the NTD ring

As our structural analysis revealed that the gMCM8/9 hexamer is the trimerization of the MCM8/9 heterodimer, this leads to the formation of three dimer interfaces within each MCM8/9 heterodimer and three putative trimer interfaces between each neighboring MCM8/9 heterodimer (**Figure 2A**). The high-resolution structure of the gMCM8/9 NTD ring allows us to analyze these two types of interfaces in detail, which serve as the structural basis of MCM8/9 assembly.

The dimer interface between NTDs of gMCM8 and gMCM9 is largely composed of the β -strands from the OB domain of gMCM9 and the flanking ZF and OB domain of gMCM8. Both the polar and hydrophobic interactions between respective β -strands are responsible for the binding of the MCM8/9 dimer. For example, the side chains of R220_{gMCM9} and Q248_{gMCM9} form hydrogen bonds with the carbonyl groups in the main chains of I172_{gMCM8} and T226_{gMCM8}, respectively. In addition, the main chain of T119_{gMCM9} also forms hydrogen bond with the neighboring carbonyl group in the side chain of D228_{gMCM8}. Furthermore, the benzene ring of F253_{gMCM9} inserts into a hydrophobic pocket surrounded by residues L179, P217, F181, L201 and Y199 from gMCM8 and W250 from gMCM9, which further stabilize the interaction between gMCM8 and gMCM9 (**Figure 2B**, S6A and S6B).

Compared to the strong interaction between the dimer interface, the interaction within trimer interface is weaker. The trimer interface is dominated by electrostatic interaction and hydrogen-bonding interactions. The side chain of R152_{gMCM8} forms salt bridges with both the side chains of D200_{gMCM9} and D232_{gMCM9}. In addition, the side chain of K158_{gMCM8} forms a hydrogen bond with the side chain of E37_{gMCM9} (**Figure 2C**, S6C and S6D). Based on the preceding analysis, we assume that opening a “gate” on the trimer interface for DNA loading will be easier than on the dimer interface. Taken together, these interactions facilitate and stabilize the assembly of the NTD hexameric ring of gMCM8/9.

Structural comparison of MCM8/9 and MCM2-7

To explore the functional state of gMCM8/9, we performed structural comparisons between gMCM8/9 and various MCM2-7-containing complexes. We first compared the gMCM8/9 structure with MCM2-7 in the OCCM complex by aligning their respective C-tier rings. As shown in **Figure 3A**, the NTDs of gMCM8/9 fit well to those of MCM7, MCM3 and MCM5, but rotated slightly in the following MCM2, MCM6 and MCM4. This minor variation is due to the opening “gate” of the MCM5-MCM2 interface in the OCCM complex. When superimposing the hexameric gMCM8/9 complex onto one hexamer of the MCM2-7 DH, we found that the two C-tier rings aligned well while their N-tier rings had a $\sim 30^\circ$ clockwise rotation (**Figure 3B** and **3C**). Such rotation could be caused by the two hexameric complex's different assembly patterns. A more plausible explanation is that the assembly of the MCM2-7 DH induces the conformational change in its N-tier ring (10). We also compared the gMCM8/9 structure with that of MCM2-7 in the apo CMG complex and found that the NTD ring has a similar translation and rotation but slighter than comparison with MCM2-7 DH (data not shown). As the NTD ring of MCM2-7 that transits from the DH to the CMG just requires a small domain translation (29). Overall, our structure of gMCM8/9 was conformationally similar to the structure of MCM2-7 complex in its loading intermediate OCCM rather than that in the DH. This is also consistent with previous finding that the recombinant MCM8/9 complex alone exhibited limited helicase activity *in vitro* (24). Therefore, we presume the current structure of gMCM8/9 may represent a loading state of MCM8/9 helicase.

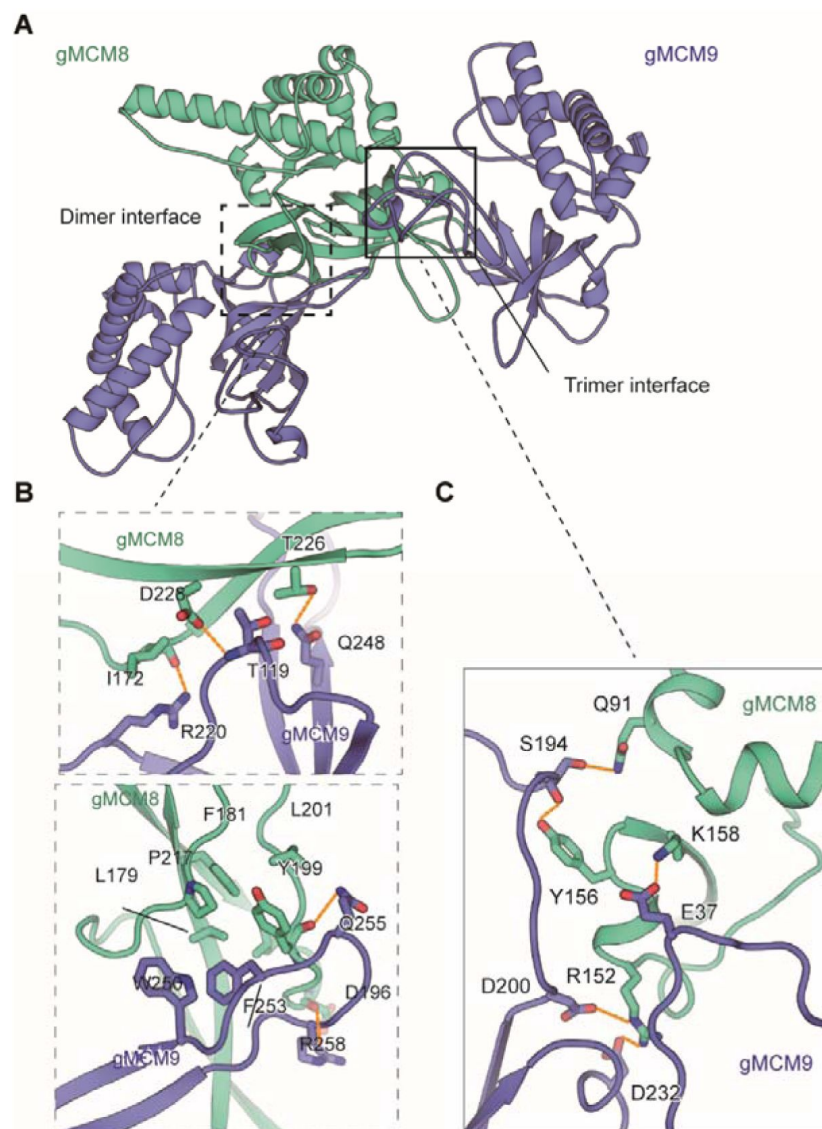


Figure 2.

The assembly analysis of the gMCM8/9 NTD ring.

(A) The cutoff structure of the gMCM8/9 NTD presents in cartoon. The dimer interface and trimer interface were indicated by boxes with dotted line and solid line, respectively. (B, C) The dimer interface (B) and trimer interface (C) were mediated by hydrophobic interaction and polar interactions. The interaction details between gMCM8 and gMCM9 in two interfaces are shown in stereo view.

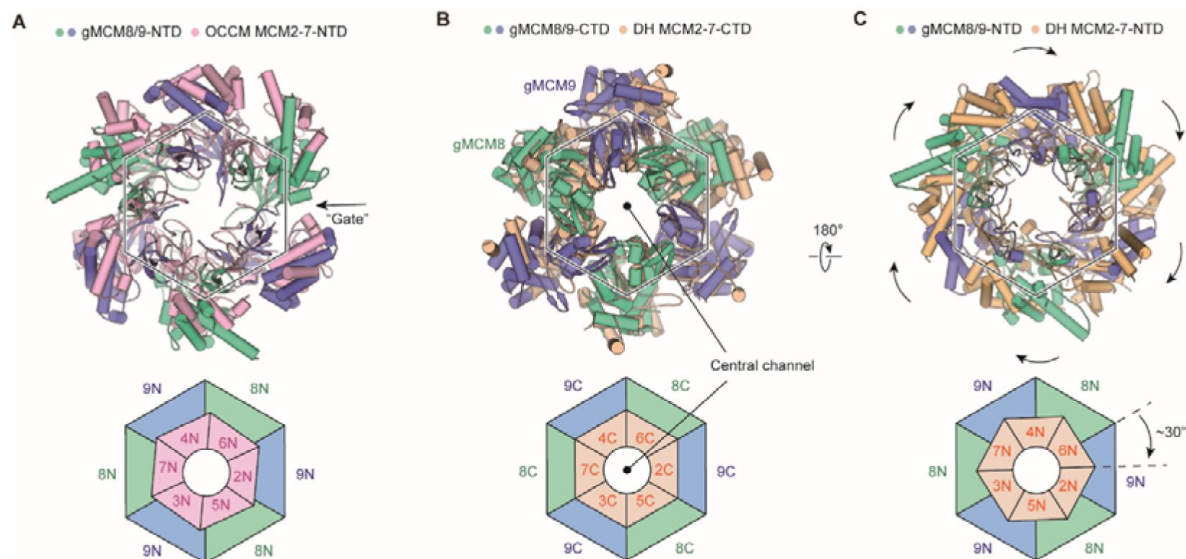


Figure 3.

Structural comparison of gMCM8/9 with MCM2-7-containing intermediates.

(A) The gMCM8/9 hexamer (greencyan and slate) was aligned to MCM2-7 hexamer (lightpink) from the OCCM complex. The N-tier rings comparison are presented here with a slightly rotation beginning at MCM5-MCM2 “Gate” and the following MCM6 and MCM4. The “Gate” was indicated by the black arrow. (B, C) Structural superposition of the gMCM8/9 hexamer to the MCM2-7 double hexamer (wheat) by aligning their respective C-tier ring. The bottom view (B) and top view (C) are shown respectively. Note the $\sim 30^\circ$ clockwise rotation of the gMCM8/9 N-tier ring comparing to that of MCM2-7.

HROB activates MCM8/9 probably by inducing conformational change of its NTD ring

To explore the activation mechanism of hMCM8/9, we examined its activity in the presence of OB-fold containing protein HROB (Figure 4A). Using a helicase assay, we found that the HROB-MBM motif was unable to stimulate the helicase activity of hMCM8/9. However, its CTD, which includes the OB fold domain, could significantly activate hMCM8/9 (Figure 4B). The HROB-CTD not only directly interacts with hMCM8/9 NTD, but also forms a complex with it, as confirmed by the GST pulldown assay (Figure 4C).

We collected the complex formed by the HROB-CTD and hMCM8/9 NTD proteins described above and reconstructed their structure using cryo-electron microscopy (Figure 4D, Table 2, Materials and Methods). Unfortunately, the density map of HROB-CTD was not observed in the reconstruction. And the 3.95 Å structure of hMCM8/9 NTD ring was reconstructed with *C*₁ symmetry, in contrast to the structure of gMCM8/9, which has *C*₃ symmetry (Figure S7). We also found that the structure of hMCM8/9 NTD ring underwent a large conformational change when compared to the previously described hMCM8/9 NTD (27). As a result, the two conformations were designated as Conformation I (hMCM8/9 NTD at 6.6 Å) and Conformation II (hMCM8/9 NTD at 3.95 Å) (Figure 4E and 4F). In line with our observation, when superimposed these two conformations of hMCM8/9 NTD, we found that one of the trimer interfaces in Conformation II was expanded by ~5 Å (Figure 4G). This expansion further pushed its neighboring MCM8/9 heterodimers to shift to either side. The left dimer translates downward by ~8 Å and tilts by ~6° while the right one shifts by ~7 Å and tilts by ~8° (Figure 4I and 4J). Interestingly, the ZF of MCM8 shifts by ~3 Å to the left and the ZF of MCM9 shifts by ~4.5 Å to the right, resulting in an overall expansion of ~6 Å (Figure 4G). These changes are consistent with our analysis for the trimer interface of gMCM8/9, which is easier to open for accommodating DNA.

Notably, the N-tier ring of MCM2-7 from the CMG-DNA complex translates laterally by ~12 Å and tilts by ~12° when compared to the structure of MCM2-7 DH (10). Without DNA, the C-tier ring of MCM2-7 from the apo CMG also adopts two different conformations including tilted and untilted, indicating a conformational change of MCM2-7 during activation (29). Based on the above observations and the stimulation effect of HROB on the helicase activity of MCM8/9, we propose that HROB activates MCM8/9 by inducing a conformational change of its NTD ring.

Spatial distribution and functional role of the OB fold hairpins

As the hairpin loops of OB fold domains are essential for DNA binding and/or unwinding in the reported hexameric MCM helicase complexes (14), we analyzed the two types of hairpin loops from OB domains in the NTD rings of gMCM8/9 and hMCM8/9. The first type of OB hairpins of gMCM8 and gMCM9 (refer to OB-hp) protrude into the central chamber in a three-fold symmetry manner and narrow down the channel (Figure 5A). In particular, the OB-hp of gMCM8/9 form two layers. The three OB-hp of gMCM8 form the upper layer and the three gMCM9 OB-hp comprise the lower layer (Figure 5C). Importantly, the tip of the gMCM8 OB-hp contains several positively charged residues (K337, K339, K342) while the gMCM9 OB-hp contains continuous polar residues (H254, Q255, D256) (Figure 5D), which are conserved in hMCM8/9 (R345, K347, K350 in hMCM8 and Q252, Q253, D254 in hMCM9). As seen in the recently reported cryo-EM structure of the *S. cerevisiae* CMG on a forked DNA, certain conserved residues, such as K364 and K367 from MCM7 OB hairpin, R449 and R451 from MCM4 OB hairpin loop and polar residues Q308 and N307 from MCM3 OB fold domain, are involved in the interaction with forked DNA and may contribute to its separation (4). We artificially docked a forked DNA into the central channel to generate a gMCM8/9-DNA model and found that the OB-hps of gMCM8 are capable to closely contact with it and insert their highly positively charged terminal loops into the major or minor grooves of the DNA strand, implying that they could be involved in substrate DNA processing and/or unwinding (Figure 5C).

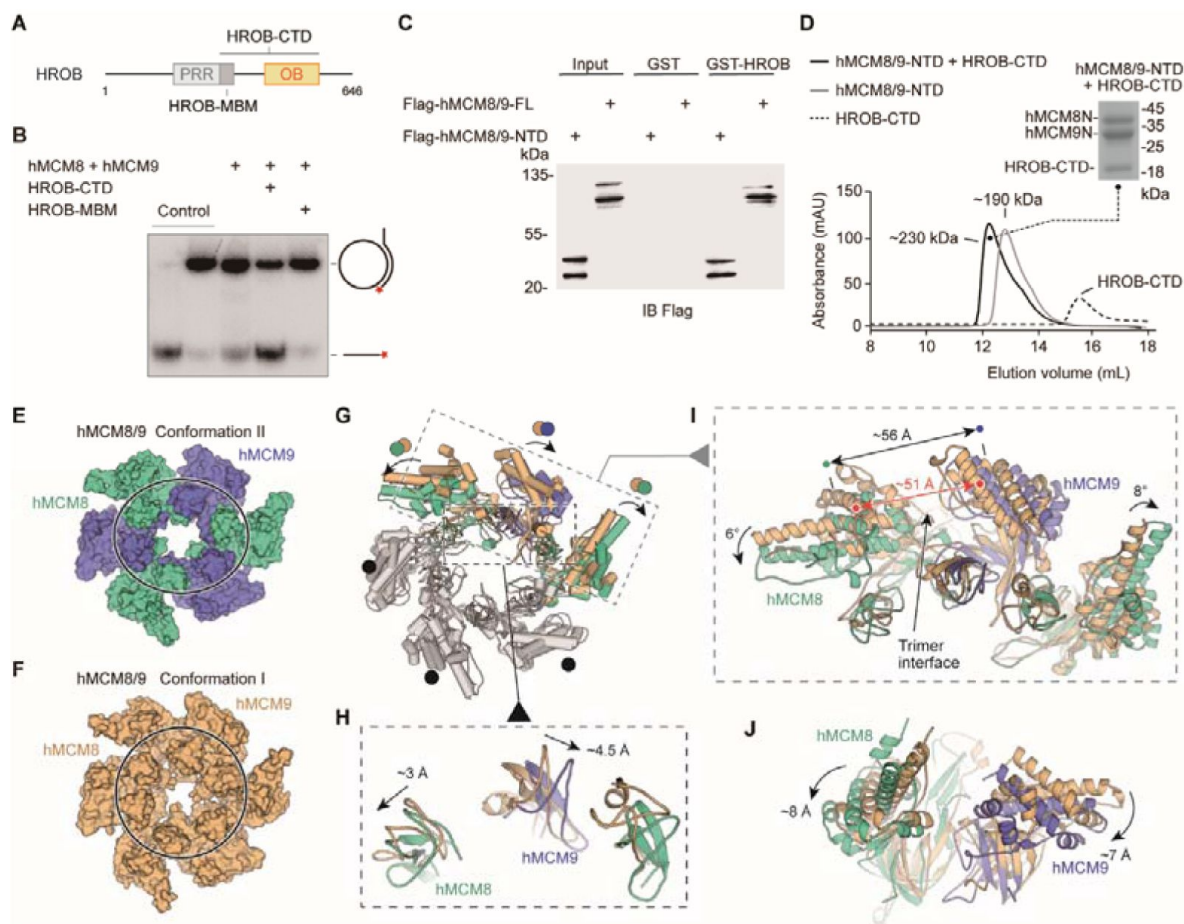


Figure 4.

HROB induces conformational change of hMCM8/9 complex.

(A) Domain organization of human HROB. PRR, proline-rich region, grey; MBM, MCM8/9 binding motif, dark grey; OB, OB fold domain, yellow. The HROB-CTD consisting of the HROB-MBM and OB domain is indicated. (B) The OB domain is required for MCM8/9 helicase in DNA unwinding. Representative autoradiograph of the DNA unwinding reaction conducted using a ^{32}P -labeled ssDNA oligo annealed to the M13mp18 as DNA substrate in the presence of HROB-MBM or HROB-CTD with purified hMCM8/9. Control, DNA substrate without proteins. (C) HROB-CTD interacts with MCM8/9-NTD. Detection by western blot of MCM8/9-FL or MCM8/9-NTD co-precipitated by bead-bound GST or GST-HROB. (D) The diagram of gel filtration shows the co-purified protein complex of HROB-CTD and MCM8/9-NTD. Their complex also indicated in SDS-PAGE. (E) Reconstructed cryo-EM map of MCM8/9 NTD ring in Conformation II. MCM8, greencyan; MCM9, slate. The shape of the channel is oval. (F) The structure of hMCM8/9-NTD from previous study. MCM8/9, orange. (G-J) Structural superposition of the hMCM8/9-NTD in Conformation I and Conformation II. The translations of the Zinc fingers were showed in (H). The ~ 5 Å expansion of the trimer interface of hMCM8/9-NTD in Conformation II was shown in (I). The shift distance and angles of MCM8/9 heterodimer were also shown in (I) and (J), respectively.

Data collection and processing	
Magnification	105,000
Voltage (kV)	300
Electron dose (e ⁻ /Å ²)	50
Frame	32
Under-focus range (μm)	1.7-2.2
Pixel size (Å)	0.827
Symmetry imposed	<i>C1</i>
FSC threshold	0.143
Cryosparc processing	
Initial particle images (no.)	1,855k
Final particle images (no.)	95,257
Global map resolution (Å)	3.95
Model composition	
Initial model used (PDB code)	7DP3 and 7DPD
Non-hydrogen atoms	13878
Protein and DNA residues	1749
Ligands	0
R.m.s. deviations	
Bond lengths (Å)	0.003
Bond angles (°)	0.682
Validation	
MolProbity score	1.99
Clashscore	19.18
Rotamers outliers (%)	0.06
Ramachandran plot	
Favored (%)	96.6
Allowed (%)	3.28
Outliers (%)	0.06

Table 2.

Cryo-EM 3D reconstruction and refinement of the hMCM8/9 NTD ring.

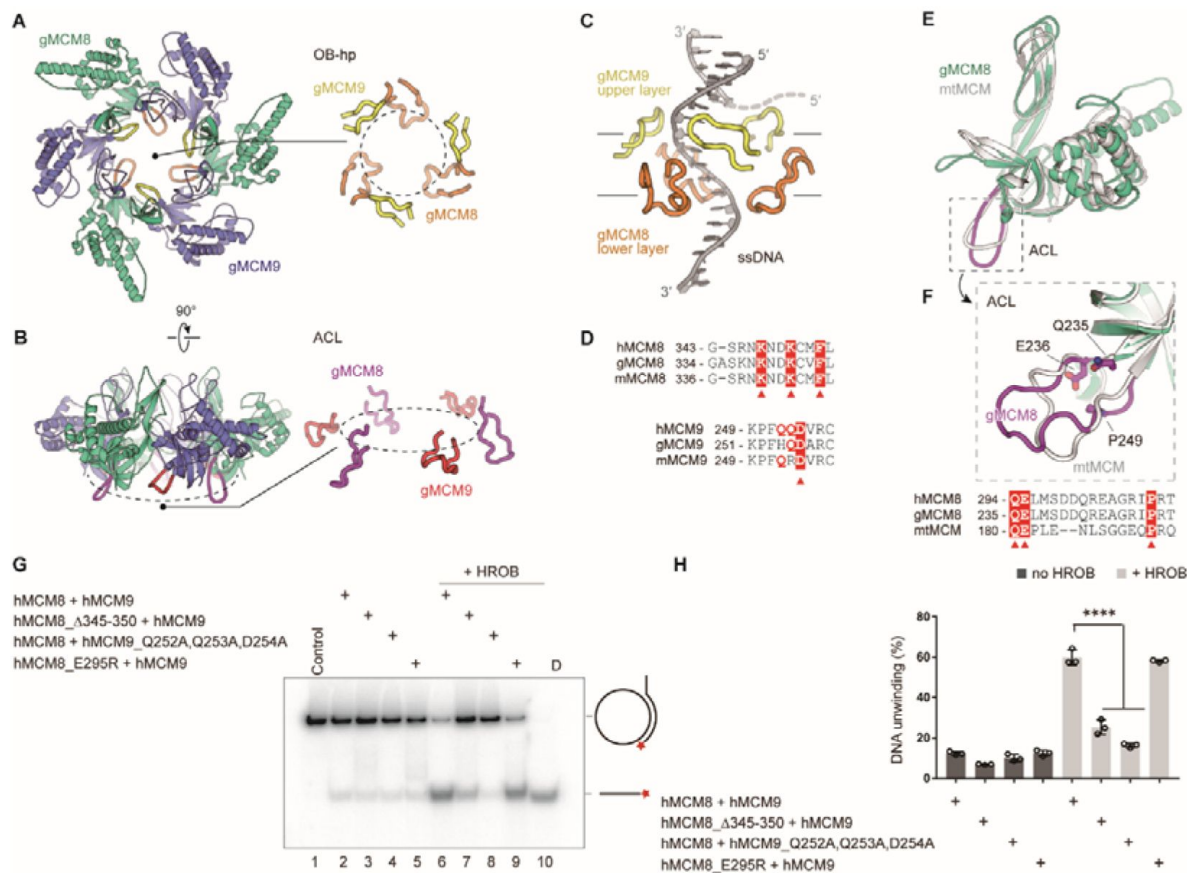


Figure 5.

The spatial distribution of OB fold hairpins and their functional roles in DNA unwinding.

(A, B) Top and side views of the structure of the gMCM8/9 NTD ring. OB-hps of gMCM8 and gMCM9 are highlighted in orange and yellow, respectively (A); ACLs are highlighted in purple and red as indicated (B). (C) Contacts between OB-hps and forked DNA illustrated through fitting a forked DNA fragment into the hexameric gMCM8/9 central channel. While encircling the DNA, the OB-hps of the gMCM9 form the upper layer and that of gMCM8 constitute the lower layer. (D) Sequence alignments of the OB-hps of MCM8 and MCM9 from different species. h, human; m, mouse; g, chicken (*Gallus gallus*). The highly conserved residues are labeled with red triangles. (E, F) Structure superposition of the gMCM8 NTD/ mtMCM NTD (gMCM8, greencyan; mtMCM, PDB: 1LTL, grey). The ACL is highlighted by purple (gMCM8) (E). Structure-based alignment of MCM from different species are shown below the structures and key residues are labeled with red triangles (F). (G) Representative autoradiograph of the DNA unwinding reaction conducted using a 32 P-labeled ssDNA oligo annealed to the M13mp18 as DNA substrate in the presence of HROB with purified MCM8/9 or mutants as indicated. (H) Graphical representation of the percentage of DNA unwinding in reactions conducted as in (G). The mean $\pm$ SD of three independent experiments is presented. Statistical analysis was conducted using one-way ANOVA (** p <0.01, *** p <0.001).

The OB domains of gMCM8 and gMCM9 also harbor another type of interesting hairpin loop known as the allosteric communication loops (ACLs) (Figure 5B). These loops are located on the opposite side of the OB-hp but extend to the CTD ring of gMCM8/9. Several studies have found that the ACLs of *M. thermotrophicus* MCM (mtMCM) and *Sulfolobus solfataricus* MCM (SsoMCM) helicases facilitate interactions between NTDs and AAA+ catalytic domains of hexameric helicases as well as modulate the positioning of the OB-hp and then regulate the helicase activities (15,16). As shown in Figure 5E and 5F, the ACLs show a high degree of conservation when performing superposition of gMCM8 NTD onto the structure of mtMCM (15). The sequence alignment of the ACLs also highlights some key residues including Q235, E236 and P249 in gMCM8 (Q294, E295, P308 in hMCM8) that are conserved in mtMCM and important for the mtMCM helicase activity (Figure 5E and 5F) (15). Thus, we propose that the ACLs are required for the helicase activity of MCM8/9.

To investigate the role of OB domain hairpins in regulating the helicase activities of MCM8/9, we deleted the tip residues (345-RNKNDK-350 in hMCM8) to shorten the OB-hp or mutated the polar residues (Q252, Q253, D254 in hMCM9) to change the charge property of OB-hp. Then we performed an *in vitro* helicase assay based on plasmid-based substrate (30,31), using the recombinant hMCM8/9 complex and corresponding mutants (Figure S8). In the helicase assay, we observed that both the wild type (WT) hMCM8/9 complex and its mutants only exhibited limited DNA unwinding activity. In the presence of HROB, the WT hMCM8/9 exhibited ~6-fold stimulation of the helicase activity, while the mutants deleting the tip residues of the hMCM8 OB-hp or mutating that of hMCM9 OB-hp both showed decreased helicase activities. However, the hMCM8 ACL mutant (hMCM8_E295R) corresponding to that in mtMCM had no significant change even though in the presence of HROB, implying that the MCM8's NTD doesn't interact with its CTD directly through ACL hairpins (Figure 5G and 5H). Collectively, the above structural analyses and biochemical experiments suggest that the OB-hps of MCM8/9 play an important role in DNA unwinding.

Structural dynamic study of the gMCM8/9 CTD ring

Since the C-tier of MCM2-7 undergoes conformational change coupling with ATP hydrolysis (32), we propose that the CTD AAA+ domain of MCM8/9 is also mobile for DNA unwinding, serving as a motor domain. Notably, the electron density of gMCM8/9 CTD ring is a little blurry and its structure was reconstructed at 5.21 Å (Figure S1 and S9). On the basis of our cryo-EM dataset, we performed structural flexibility analyses using multi-body refinement to investigate the dynamics of the gMCM8/9 complex. In this method, we used separate focused refinements and generated movies to describe the most important motions on the relative orientations of the rigid bodies, which allows us to gain insight into the molecular motions of gMCM8/9, particularly regarding the structural flexibility of its CTD ring (33–35). As expected, twelve components were generated, each with a discrete number of independently moving bodies, providing a characterization of the motions in gMCM8/9 CTD ring. As shown in the movie 1, the CTD AAA+ motor ring reciprocally rotates right and left relative to the NTD ring. It would be very interesting to see similar rotary movement along DNA, which may underlie the translocation/unwinding mode for MCM8/9 helicase.

The N-C linkers is required for MCM8/9's helicase activity

Unlike MCM8/9, such rotary motion between NTD and CTD rings has yet to be observed in the structures of MCM2-7. We analyzed the linkers connecting the NTD and CTD (N-C linkers) of MCM8/9 and found that these N-C linkers, which are 30-40 residues long, are longer than those in MCM2-7. In MCM2-7, the N-C linker of MCM 3 is the shortest, which may restrict the rotary motion of MCM2-7 (Figure S10). To test whether the N-C linkers are crucial for the relative motion between N-tier and C-tier rings of MCM8/9, we shortened the N-C linkers by deleting several amino acids in loop regions of hMCM8 (MCM8Δ369-377) or hMCM9 (MCM9Δ283-287) (Figure S11). In the helicase assay *in vitro*, we found that both single and combined deletions have significantly lower

unwinding ability than the WT MCM8/9 complex. The presence or absence of HROB had no effect on this result, indicating that the N-C linkers are required for the MCM8/9 helicase activity (**Figure 6A** and **6B**).

We next studied whether the N-C linkers are important for the *in vivo* functions of MCM8 or MCM9. To do so, we employed a chemoresistance assay because MCM8 and MCM9 had been reported to mediate cellular resistance to the DNA cross-linking reagent cisplatin, which induces DNA interstrand crosslinks that are mostly repaired by HR (17,19,36). As shown in the cell-viability assay, we found a notable reduction in survival of *MCM8*- or *MCM9*-deficient DT40 cells following cisplatin treatment compared to the WT DT40, indicating that the KO cells exhibit significant sensitivity to cisplatin. Expression of WT MCM8 or MCM9 cDNAs in their respective KO cells almost completely complemented chemoresistance. In contrast, expression of their N-C linker deletion mutants had minimal complementary effects (**Figure 6C**, 6D, 6E and 6F). These results suggest that the N-C linkers of MCM8/9 are required for its cellular resistance to DNA-damaging reagents.

Discussion

In this study, we reconstructed two structures of the MCM8/9 helicase complex at near-atomic resolution. We found that MCM8 and MCM9 form a single heterohexameric ring with a centric channel that allows the DNA to pass through. The OB hairpin loops within the centric channel may be involved in interacting with and separating substrate DNA. The N-C linkers of MCM8/9 play important roles in helicase activity. Notably, the regulator HROB induces conformational change of MCM8/9 NTD ring, which may underlie the helicase activation.

MCM8/9 and MCM2-7 are both MCM proteins with highly conserved domain organization and sequence similarity (3,37). Compared with the heterohexameric structure of MCM2-7, the MCM8/9 hexamer is arranged in an alternate mode with the three-fold symmetry axis. Interestingly, it was reported that the *Escherichia coli* helicase DnaB also exists in a quaternary state with *C*3 or *C*6 symmetry and can change the symmetry states reversibly depending on the pH value of the buffers or the presence of nucleotides (38). It has been proposed that the symmetry transition plays a role in the loading of regulators on DNA and is directly linked to DNA translocation (39,40). Similarly, we doubt that MCM8/9 changes its symmetry from *C*3 to *C*1 symmetry in the presence of HROB. In other words, helicases may orchestrate the inner channels by controlling the relative positions of subunits, assisting in DNA loading and unwinding.

Taking inspiration from the structural and unwinding mechanisms of MCM2-7, we also investigated the key elements of MCM8/9 involved in DNA interaction or separation. In the central channel of the MCM2-7 complex, several structural elements involved in DNA binding and strand separation were illustrated by a series of CMG structures complexed with different types of DNA (4). According to recent research, the OB hairpin loops of MCM3, MCM4, MCM6 and MCM7 form “dam-and-diversion tunnel” to block and divert the lagging strand from the leading strand (4). For the MCM8/9 complex, the OB-hps of MCM8 and MCM9 are essential for DNA unwinding. They form a two-layered structure with three-fold symmetry. In which, three OB-hps of MCM9 form the upper layer, while three OB-hps of MCM8 form the lower layer, suggesting a synergistic mechanism for MCM8/9 OB hairpins to interact with DNA. When modelled a fork DNA into the channel, the OB-hp loops make very close contact with the DNA strand that passes through the channel, capable of inserting their loops into the major or minor grooves of the DNA in a sequential manner. While in the Conformation II of hMCM8/9 NTD ring, the OB-hps of hMCM8 and hMCM9 lose their *C*3 symmetry without significant conformational change. This symmetry transition could be caused by HROB while inducing trimer interface expansion. In addition, the highly conserved residue F363 from OB hairpin loop of MCM7 serves as a strand separation pin to

unwind forked DNA via making a π - π interaction with DNA (41 [↗](#)). The highly conserved phenylalanine was also found on the OB-hps of MCM8 and MCM9 (F353 of hMCM8, F251 of hMCM9) (Figure 5D [↗](#)). They will most likely perform functions similar to those found in MCM2-7.

We also present a new conformation (Conformation II) of the MCM8/9 NTD ring induced by HROB, which may enhance the DNA binding ability of MCM8/9 complex. We found that when MCM8/9 is complexed with the CTD domain of HROB, it becomes more stable and exhibits significantly higher helicase activity. The HROB CTD domain is about 20 kDa and is composed of a flexible loop and an OB fold domain. The flexible loop containing ~20 amino acids is sufficient to interact with the MCM8/9 complex, but it alone is unable to stimulate the helicase activity of MCM8/9 (24 [↗](#)). Our helicase assay demonstrated that the OB fold domain of HROB is required for MCM8/9 activation. The OB fold domain is a nucleic acid-binding motif that likely help to load the DNA into the central channel of MCM8/9. Induced by HROB, the structure of MCM8/9 NTD ring convert its C3 symmetry to C1 and two of the MCM8/9 heterodimers translate and tilt following the expansion of the trimer interface. When the MCM2-7 double hexamer encircles the double-stranded origin DNA, the MCM5-MCM2 interface characterizes as a “gate” that undergoes open and closed conformation change to engage the DNA entry (9 [↗](#)). The expansion of the MCM8/9 trimer interface may involve a transient conformation state to facilitate the DNA entry into the inner channel of the MCM8/9 hexamer. Thus, HROB not only aids in the loading of DNA into MCM8/9, but also acts as an activator, collaborating with MCM8/9 to promote DNA unwinding.

Benefit from the rapid development of cryo-electron microscopy technology over the past decade, a dozen structures of MCM2-7-containing intermediates have been resolved including the OCCM, DH and CMG with/ without DNA (4 [↗](#)). Based on these structures, the hand-over-hand rotary mode and the “dam-and-diversion tunnel” model had been proposed for MCM2-7 helicase in DNA translocation and unwinding (4 [↗](#), 12 [↗](#)). For MCM8/9 helicase, there is currently no established model to describe its unwinding mechanism. Here, our structural and biochemical studies have provided some hints in this regard. First of all, the structural dynamics analysis reveals that the MCM8/9 CTD ring rotates relative to the NTD ring. Secondly, the length of the MCM8 and MCM9 N-C linkers are surprisingly much longer than those of MCM2-7 complex, which may permit the rotations of MCM8/9. The fact that single-site structure-guided mutations on the ACL loops has no effect on the helicase activity of MCM8/9 further supports the idea that the N-C linkers, rather than the ACL loop, mediated the interaction between the N-tier ring and the C-tier ring of MCM8/9. Last but not least, HROB can convert inactive MCM8/9 into an active helicase by recruiting and inducing conformational change in the MCM8/9 NTD ring.

Although a rotary model is in sight and our findings also hint at a distinct unwinding mechanism of MCM8/9, many questions remain unsolved. The most pressing issue is that the type of DNA substrate unwound by MCM8/9 is unknown. In vivo sequencing and in vitro DNA binding results are required to screen appropriate substrate. Furthermore, as the HROB only stimulate ~60% of the MCM8/9 helicase in vitro, we believe that other regulators remain to be discover for the full activation of MCM8/9 helicase. Based on these results, the high-resolution structures of MCM8/9 with DNA in the presence of ADP or ATP are necessary to fully clarify the unwinding mechanism of MCM8/9. Despite these limitations, the MCM8/9 complex structures provided in this study will be a rich source of information for further research into the mechanism of DNA helicase in homologous recombination, as well as for the analysis of multiple disease mutations in MCM8/9 associated with premature ovarian failure as well as cancers.

Material and methods

Plasmids

The human MCM8/9 genes were obtained by reverse transcription of mRNA from HeLa cells. The gene fragments coding MCM8/9 of *Gallus gallus* were also from corresponding reverse transcripts and the total RNA was extracted from the chicken DT40 cell line. Human MCM8_61-840 and MCM9_1-684 were inserted into the pFastbac-Dual vector (Invitrogen). *Gallus gallus* MCM8_50-830 and MCM9_1-691 were inserted into the pFastbac-1 vector, respectively. All the MCM8 were fused with 6× His tag at their C-terminus for subsequent affinity purification. The genes coding hMCM8/9 NTDs (MCM8_61-376, MCM9_1-276) and relative mutants were cloned into multiple cloning site I (MCS I) of pRSFDuet-1 vector (Novagen). All of the mutations were introduced by the standard PCR-based mutagenesis method described before (42). Human HROB (NM_001171251.3) was obtained by reverse transcription of mRNA from 293T cells. HROB-CTD_391-580 was inserted into pET28S-SUMO vector which expresses the N-terminal SUMO tag. All constructs were confirmed by DNA sequencing.

Protein expression and purification

To express the core hMCM8/9 and gMCM8/9 complex proteins, recombinant baculoviruses were prepared using the Bac-to-Bac expression system (Invitrogen). The proteins were expressed in *Trichoplusia ni* (BTI-Tn5B1–4, Hi5) insect cells for 60 hours at 27°C before harvesting. The cell pellet was collected by centrifugation at 800× g and then freshly frozen by liquid nitrogen and stored at -80 °C before use. For purification, the cell pellet was resuspended with lysis buffer (50 mM HEPES, pH 8.0, 350 mM KCl, 50 mM sodium glutamate, 10% glycerol and 30 mM imidazole) and lysed by cell homogenizer (Avestin Emulsiflex C3). Then the lysate was clarified by centrifugation at 16,000× rpm for 30 min. The supernatant was mixed with nickel-NTA resin (Novagen) and was continuously stirred for 1 hour at 4 °C for adequate binding. The beads were then collected into a column and extensively washed with lysis buffer to remove undesired proteins. Target proteins were then eluted down with elution buffer (50 mM HEPES, pH 7.8, 80 mM KCl, 20 mM sodium glutamate, 5% glycerol and 300 mM imidazole). The proteins were further purified by HiTrap Heparin HP 5ml column (GE Healthcare) and size exclusion column (Superose 6 Increase 10/300 GL, GE Healthcare) in buffer (20 mM HEPES, pH 7.8, 150 mM KCl, 1 mM DTT and 5% glycerol). The strategies of purification of MCM8/9 mutants were as the same as that used for the core hMCM8/9 complex.

For expression of HROB-CTD, 10 mL BL21 codon plus strain containing the recombinant pET28a-HROB-CTD plasmid was inoculated into 1 L LB medium. The bacteria solution was firstly cultured at 37°C until the OD600 ≈ 0.8 and then cooled down to 16°C. The expression of recombinant protein was induced by IPTG (isopropyl-β-d-thiogalactoside) at a concentration of 0.2 mmol/L and kept expressing for another 20 hours. Subsequently, the cells were harvested and disrupted by a high-pressure cell homogenizer in lysing buffer. After centrifugation, the supernatant was mixed with nickel-NTA beads and retained for 1 h at 4 °C with rolling. Before elution, the SUMO tag was cleaved by Ulp Protease and finally the HROB-CTD protein was eluted in the buffer containing 50 mM HEPES, pH 7.8, 80 mM KCl, 20 mM sodium glutamate, 5% glycerol. Following Ni-NTA affinity purification, the eluted protein was further purified by HiTrap Heparin HP 5ml column (GE Healthcare) and size exclusion chromatography (HiLoadTM16/600 Superdex TM pg, GE Healthcare) in the same buffer using for hMCM8/9. The GST tag HROB-CTD was also expressed in BL21 (DE3) *E. coli* cells and purified using GSH-Sepharose affinity chromatography (GE Healthcare) following by the size-exclusion chromatography. And the HROB-MBM peptides (HROB_391-413) were commercial synthesized by China Peptide Co., Ltd.

Cryo-EM sample preparation

The fresh gMCM8/9 proteins were used to prepare the cryo-EM sample. Briefly, 0.8 μL crosslinker BS(PEG)₉ (Thermo scientific) and 10 μM protein were incubated for 1 hour at 4°C and filtered by Superose 6 in buffer (20 mM HEPES, pH 7.8, 150 mM KCl) after centrifugation. The fractions were identified by SDS-PAGE and the peak fraction was diluted to about 0.3 mg/mL to prepare cryo-EM samples. 4 μL fresh sample was applied on glow-discharged Cu holey carbon grids (Quantifoil R 1.2/1.3) and incubated for 1 min at room temperature. Grids were then blotted for 3 s in 100% humidity at 4°C and plunged frozen in liquid ethane cooled by liquid nitrogen using Vitrobot (Thermo).

For the hMCM8/9-HROB complex, the hMCM8/9 NTDs and HROB proteins were mixed by the ratio 1:3. 4 μL fresh sample (0.48 mg/mL) was applied on glow-discharged Au holey carbon grids (Quantifoil R 0.6/1) and the following operation were the same as describe above for preparing the sample of gMCM8/9 complex.

Cryo-EM data acquisition

The grids of gMCM8/9 were loaded onto a transmission electron microscope operated at 200 kV (FEI Talos Arctica) in liquid nitrogen temperatures. Images were recorded on a Gatan K2 camera in counting mode, at a nominal magnification of 29,000 $\times$. The total dose rate was about 50 $\text{e}^-/\text{\AA}^2$ for each micrograph stack. The total dose was fractionalized to 32 frames equally, corresponding to a physical pixel size of 1.0 $\text{\AA}$ (0.5 $\text{\AA}$ super-resolution pixel size). The defocus range was set from -1.7 to -2.2 μm . 2,891 raw movie micrographs were collected using the serialEM software (43 [43](#)).

The grids of hMCM8/9 NTD complex were loaded onto a transmission electron microscope operated at 300kV (FEI Titan Krios) equipped with a K3 direct electron detector. The parameter settings of total dose and defocus range were as same as that of gMCM8/9. While the images' physical pixel size is 0.827 $\text{\AA}$ and the super-resolution pixel size is 0.4135 $\text{\AA}$. 4078 raw movie micrographs were collected using the Thermo Fisher Scientific EPU software.

Single particle data processing

The data set of gMCM8/9 was processed by Cryosparc-3.2 (44 [44](#)). All the movies were processed using patch motion correction and patch CTF estimation. A total of 2,134,191 particles were automatically picked by template picker and 1,574,821 particles were produced after the inspect particle picks treatment. Three parallel runs of 2D classification ($K = 100$) were performed using the data (binned 4) with a box size at 256 pixels. A small subset of the “good” particles with clear secondary-structure features was selected from 2D classification results and was used to generate 4 initial models. The 434,320 particles in the best initial model class were re-extracted with original pixel size and further refined to a 6.7 $\text{\AA}$ low-resolution reconstruction using non-uniform refinement (New) (45 [45](#)). Four resolution gradient templates generated from non-uniform refinement were used for the multi-reference heterogeneous refinement, resulting in a 4.31 $\text{\AA}$ global reconstruction (290,080 particles) renamed consensus map. The NTD map was improved to 3.73 $\text{\AA}$ with local refinement followed by applying a tight N-terminal mask with $C3$ symmetry. Local CTF refinement further improved the resolution to 3.66 $\text{\AA}$. For the CTD map, a reference was morphed from a predicted gMCM8/9 CTD model using AlphaFold (Figure S12) (46 [46](#)). Before processing CTD local refinement (applying $C3$ symmetry), the projection of NTD particles from the 3.66 $\text{\AA}$ NTD map were subtracted from the 4.31 $\text{\AA}$ consensus map, and generated a 6.57 $\text{\AA}$ CTD hexamer map. Finally, 3D classification and local refinement resulted in a 5.21 $\text{\AA}$ (mean resolution, validation from Phenix (47 [47](#))) CTD map using 72,948 particles. The 3.66 $\text{\AA}$ NTD map and 5.21 $\text{\AA}$ CTD map were aligned to the consensus map, respectively, and then merged together to generate a final map.

The EMDB accession code is EMD-32346 and the PDB entry ID is 7W7P. The image processing and 3D reconstruction steps were illustrated in Figure S13.

RELION-3.1.1 (48 [↗](#)) was also used to process the same data, resulting in a 4.2 Å NTD map and a 5.4 Å CTD map. All the movie frames were motion-corrected by MotionCorr2 (49 [↗](#)) with a patch alignment of 5×5. CTF parameters were estimated using Gctf (50 [↗](#)). 965,175 particles were auto-picked and extracted to 4×binned particles, which rescales the initial box size at 256 pixels to the final box size at 64 pixels. The particles were subjected to 2 rounds of 2D classifications (K = 100). 388,378 particles with clear secondary-structure features were selected for 3D initial model generation. Three parallel runs of multi-reference 3D classification (K = 9) were performed and the best class was merged. 144,202 particles were re-extracted at original pixel size and a final map was imported as reference and were submitted to the final 3D auto-refinement. Finally, we yielded a 6.6 Å global 3D density map estimated by the gold-standard Fourier shell correlation at a correlation cutoff value of 0.143.

As the density of the gMCM8/9 CTD was a little blurred, we then used the multi-body refinement program (51 [↗](#)) to process the dataset. 6.6 Å gMCM8/9 global map's NTD and CTD were subtracted as the first and the second subtracted body respectively using Chimera (52 [↗](#)). The corresponding masks were generated from the final map using mask creation in RELION. The multi-body refinement resulted in a 4.2 Å 3D density map of gMCM8/9 NTD and a 5.4 Å 3D density map of gMCM8/9 CTD. It generated 12 components, with each including a discrete number of independently moving bodies. The first five components explained 86.77% of the variance in the data. The image processing and 3D reconstruction steps were illustrated in Figure S14.

The data set of hMCM8/9 NTD complex was processed by Cryosparc-3.3. In brief, a total of 1,855,585 particles were automatically picked by template picker and 677,960 particles were produced after the inspect particle picks treatment. After the processing of 2D classification, Ab-Initio Reconstruction and Heterogeneous refinement (K=3), the 95,257 particles in the best 3D class were re-extracted with original pixel size and further refined to a 3.95 Å resolution reconstruction (C1 symmetry) using Non-uniform refinement (New). The image processing and 3D reconstruction steps were illustrated in Figure S15.

Model building and refinement

For the gMCM8/9 model building, the gMCM8 NTD and gMCM9 NTD monomer models were predicted through SWISS-MODEL by providing the hMCM8 NTD (PDB: 7DP3) and hMCM9 NTD (PDB: 7DPD) structural template, respectively. Firstly, the predicted structures were used as the initial model directly docking into the EM map. As the gMCM8/9 NTD has a C3 symmetry axis, another two copies were generated by horizontal rotating for subsequent rigid body-fitting into the 3D density map with program Chimera. Secondly, using the automatic refinement plugin function in COOT (53 [↗](#)) to refine the initial model. The final model was refined against the corresponding map using PHENIX (54 [↗](#)) in real space with secondary structure and geometry restraints. The gMCM8/9 CTD hexamer model was predicted as above described and the processing of docking and refinement was the same as the gMCM8/9 NTD model.

For the hMCM8/9 NTD complex model building, in brief, hMCM8 NTD (PDB: 7DP3) and hMCM9 NTD (PDB: 7DPD) models were used as the initial model directly docking into the EM map. The other processing procedure was similar to that of the gMCM8/9 model building. All the structures of the gMCM8/9 complex were validated through examination of the Molprobit scores and statistics of the Ramachandran plots. Molprobit scores were calculated as described in **Table 2** [↗](#).

Helicase assays

To prepare the substrate, the oligonucleotide (5'-(dT)₄₀GTTTCCAGTCACGACG-TTGTAACGACGGCCAGTGCC-3') containing a 40 nt region complementary to the M13mp18(+) stand and a 40 nt oligo-dT at the 5' end was labeled at the 3' terminus with [α -³²P] dCTP (Perkin Elmer) and annealed to the single-stranded DNA M13mp18(24 [↗](#)). 0.1 nM (in molecules) DNA substrates were mixed with 5 µg recombinant MCM8/9 complex as well as its mutants as indicated

within each 15 µl volume reaction in the helicase buffer (25 mM HEPES, pH 7.5, 1 mM magnesium acetate, 25 mM sodium acetate, pH 5.2, 4 mM ATP, 0.1 mg/ml BSA, 1 mM DTT). 2.5 µg HROB was used as an activator. To avoid re-annealing, the reaction was supplemented with a 100-fold unlabeled oligonucleotide. The reactions were then incubated at 37 °C for 60 min and stopped by adding 1 µl of stop buffer (0.4% SDS, 30 mM EDTA, and 6% glycerol) and 1 µl of proteinase K (20 mg/ml, Sigma) into the reaction for another 10 min incubation at 37 °C. The products were separated by 15% polyacrylamide gel electrophoresis in 1× TBE buffer and analyzed by the Amersham typhoon (Cytiva).

Cell-viability assay

Cell-viability assays were performed as described previously (17). Briefly, the *MCM8* or *MCM9* KO DT40 cell lines were transfected with plasmids containing wildtype *MCM8*, *MCM9* or the indicated mutants by electroporation. 1.0×10^3 cells were seeded into each well of a 96-well plate and treated with a range of concentrations of cisplatin. After 48 h of incubation, the cell viability was measured by the Cell counting kit 8 (CCK8) according to the standard procedures. The absorbance readings at 405nm were conducted using an Epoch Microplate Spectrophotometer (BioTek Instrumentals, Inc., Winooski, VT, USA). All experiments were performed at least 3 times.

Pull Down Assay

The HEK293T cells transfected with Flag-hMCM8/9-FL or Flag-hMCM8/9-NTD were cultured overnight and washed twice with cold phosphate-buffered saline (PBS). Cell pellets were resuspended with lysis buffer (20 mM Tris, pH7.5, 150 mM NaCl, 5mM EDTA, 0.5% NP-40, 10% glycerol, protease inhibitor cocktail (Roche, 04693132001)). After incubation for 45 min at 4°C with gentle agitation, the whole-cell lysates were collected by centrifugation ($12,000 \times g$ for 15 min, at 4 °C). GST beads coupled with 2 µg GST-HROB or GST alone were then incubated with an equal volume of above HEK293T cell lysates at 4°C for 4h. The beads were washed four times with lysis buffer. Proteins bound to the beads were separated by SDS–PAGE and subsequently immunoblotted with anti-Flag antibody (Cytiva).

Statistical analysis

The statistical analysis was performed using Prism (GraphPad Software Inc.) on the data from at least three independent experiments. Statistical significance was assessed by two-tailed unpaired Student's *t*-test. $P \leq 0.05$ was considered significant.

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Data availability

The 3D cryo-EM map of the gMCM8/9 complex and the hMCM8/9 NTD ring have been deposited to the EMDB database with accession number EMD-32346 and EMD-33989, respectively. The corresponding atomic models have been deposited at the RSCB PDB under the accession code 7W7P and 7Y0X, respectively.

Author contributions

Z.W. and J.Z. designed and performed the experiments. J.Z., Z.W. and Y. W. performed protein purification, cryo-EM sample preparation, data collection and structure model building. Z.W and Y.Z. performed the helicase assay. Z.L performed cell survival assay. Y.L., H.L., Z.W., D.X. and H.L. analyzed the data. Z.W. and H.L prepared the figures. Z.W., H.L. and Y. L. wrote the paper. All authors reviewed the results and approved the final version of the manuscript.

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Declaration of interests

The authors declare no competing interests.

References

1. Brosh R.M., Matson S.W (2020) **History of DNA Helicases** *Genes (Basel)* **11**
2. Heyer W.D., Ehmsen K.T., Liu J (2010) **Regulation of homologous recombination in eukaryotes** *Annu Rev Genet* **44**:113–139
3. Maiorano D., Lutzmann M., Méchali M (2006) **MCM proteins and DNA replication** *Current opinion in cell biology* **18**:130–136
4. Yuan Z., Li H (2020) **Molecular mechanisms of eukaryotic origin initiation, replication fork progression, and chromatin maintenance** *The Biochemical journal* **477**:3499–3525
5. Yuan Z. *et al.* (2017) **Structural basis of Mcm2-7 replicative helicase loading by ORC-Cdc6 and Cdt1** *Nature structural & molecular biology* **24**:316–324
6. Miller T.C.R., Locke J., Greiwe J.F., Diffley J.F.X., Costa A (2019) **Mechanism of head-to-head MCM double-hexamer formation revealed by cryo-EM** *Nature* **575**:704–710
7. Li N., Zhai Y., Zhang Y., Li W., Yang M., Lei J., Tye B.K., Gao N (2015) **Structure of the eukaryotic MCM complex at 3.8 Å** *Nature* **524**:186–191
8. Abid Ali F., Renault L., Gannon J., Gahlon H.L., Kotecha A., Zhou J.C., Rueda D., Costa A. (2016) **Cryo-EM structures of the eukaryotic replicative helicase bound to a translocation substrate** *Nat Commun* **7**
9. Samel S.A., Fernández-Cid A., Sun J., Riera A., Tognetti S., Herrera M.C., Li H., Speck C (2014) **A unique DNA entry gate serves for regulated loading of the eukaryotic replicative helicase MCM2-7 onto DNA** *Genes Dev* **28**:1653–1666
10. Noguchi Y., Yuan Z., Bai L., Schneider S., Zhao G., Stillman B., Speck C., Li H (2017) **Cryo-EM structure of Mcm2-7 double hexamer on DNA suggests a lagging-strand DNA extrusion model** *Proc Natl Acad Sci U S A* **114**:E9529–e9538
11. Georgescu R., Yuan Z., Bai L., de Luna Almeida Santos R., Sun J., Zhang D., Yurieva O., Li H., O'Donnell M.E. (2017) **Structure of eukaryotic CMG helicase at a replication fork and implications to replisome architecture and origin initiation** *Proc Natl Acad Sci U S A* **114**:E697–e706
12. Eickhoff P. *et al.* (2019) **Molecular Basis for ATP-Hydrolysis-Driven DNA Translocation by the CMG Helicase of the Eukaryotic Replisome** *Cell Rep* **28**:2673–2688
13. Douglas M.E., Ali F.A., Costa A., Diffley J.F.X (2018) **The mechanism of eukaryotic CMG helicase activation** *Nature* **555**:265–268
14. Slaymaker I.M., Chen X.S (2012) **MCM structure and mechanics: what we have learned from archaeal MCM** *Sub-cellular biochemistry* **62**:89–111
15. Sakakibara N., Kasiviswanathan R., Melamud E., Han M., Schwarz F.P., Kelman Z (2008) **Coupling of DNA binding and helicase activity is mediated by a conserved loop in the MCM protein** *Nucleic Acids Res* **36**:1309–1320

16. Barry E.R., Lovett J.E., Costa A., Lea S.M., Bell S.D (2009) **Intersubunit allosteric communication mediated by a conserved loop in the MCM helicase** *Proc Natl Acad Sci U S A* **106**:1051–1056
17. Nishimura K., Ishiai M., Horikawa K., Fukagawa T., Takata M., Takisawa H., Kanemaki M.T (2012) **Mcm8 and Mcm9 form a complex that functions in homologous recombination repair induced by DNA interstrand crosslinks** *Mol Cell* **47**:511–522
18. Lutzmann M. *et al.* (2012) **MCM8- and MCM9-deficient mice reveal gametogenesis defects and genome instability due to impaired homologous recombination** *Mol Cell* **47**:523–534
19. Morii I., Iwabuchi Y., Mori S., Suekuni M., Natsume T., Yoshida K., Sugimoto N., Kanemaki M.T., Fujita M (2019) **Inhibiting the MCM8-9 complex selectively sensitizes cancer cells to cisplatin and olaparib** *Cancer Sci* **110**:1044–1053
20. McKinze D.R., Gomathinayagam S., Griffin W.C., Klinzing K.N., Jeffries E.P., Rajkovic A., Trakselis M.A (2021) **Motifs of the C-terminal domain of MCM9 direct localization to sites of mitomycin-C damage for RAD51 recruitment** *J Biol Chem* **296**
21. Park J. *et al.* (2013) **The MCM8-MCM9 complex promotes RAD51 recruitment at DNA damage sites to facilitate homologous recombination** *Mol Cell Biol* **33**:1632–1644
22. Lee K.Y., Im J.S., Shibata E., Park J., Handa N., Kowalczykowski S.C., Dutta A (2015) **MCM8-9 complex promotes resection of double-strand break ends by MRE11-RAD50-NBS1 complex** *Nat Commun* **6**
23. Hustedt N. *et al.* (2019) **Control of homologous recombination by the HROB-MCM8-MCM9 pathway** *Genes Dev* **33**:1397–1415
24. Huang J.W. *et al.* (2020) **MCM8IP activates the MCM8-9 helicase to promote DNA synthesis and homologous recombination upon DNA damage** *Nat Commun* **11**
25. Brewster A.S., Wang G., Yu X., Greenleaf W.B., Carazo J.M., Tjajadi M., Klein M.G., Chen X.S (2008) **Crystal structure of a near-full-length archaeal MCM: functional insights for an AAA+ hexameric helicase** *Proc Natl Acad Sci U S A* **105**:20191–20196
26. Griffin W.C., Trakselis M.A (2019) **The MCM8/9 complex: A recent recruit to the roster of helicases involved in genome maintenance** *DNA Repair (Amst)* **76**:1–10
27. Li J., Yu D., Liu L., Liang H., Ouyang Q., Liu Y (2021) **Structural study of the N-terminal domain of human MCM8/9 complex** *Structure* **29**:1171–1181
28. Fletcher R.J., Bishop B.E., Leon R.P., Sclafani R.A., Ogata C.M., Chen X.S (2003) **The structure and function of MCM from archaeal M. Thermoautotrophicum** *Nature structural biology* **10**:160–167
29. Yuan Z., Bai L., Sun J., Georgescu R., Liu J., O'Donnell M.E., Li H (2016) **Structure of the eukaryotic replicative CMG helicase suggests a pumpjack motion for translocation** *Nature structural & molecular biology* **23**:217–224
30. Lee J.K., Hurwitz J. (2001) **Processive DNA helicase activity of the minichromosome maintenance proteins 4, 6, and 7 complex requires forked DNA structures** *Proc Natl Acad Sci U S A* **98**:54–59

31. Traver S., Coulombe P., Peiffer I., Hutchins J.R., Kitzmann M., Latreille D., Mechali M (2015) **MCM9 Is Required for Mammalian DNA Mismatch Repair** *Mol Cell* **59**:831–839
32. O'Donnell M.E., Li H (2018) **The ring-shaped hexameric helicases that function at DNA replication forks** *Nature structural & molecular biology* **25**:122–130
33. Nakane T., Scheres S.H.W (2021) **Multi-body Refinement of Cryo-EM Images in RELION** *Methods in molecular biology (Clifton, N.J.)* **2215**:145–160
34. Liu Y., Huynh D.T., Yeates T.O (2019) **A 3.8 Å resolution cryo-EM structure of a small protein bound to an imaging scaffold** *Nat Commun* **10**
35. Singh S.K., Gui M., Koh F., Yip M.C., Brown A (2020) **Structure and activation mechanism of the BBSome membrane protein trafficking complex** *eLife* **9**
36. Kanemaki M.T (2013) **The dimeric Mcm8-9 complex of *Xenopus laevis* likely has a conserved function for resistance to DNA damage** *Cell Cycle* **12**:1338–1339
37. Maiorano D., Cuvier O., Danis E., Mechali M (2005) **MCM8 is an MCM2-7-related protein that functions as a DNA helicase during replication elongation and not initiation** *Cell* **120**:315–328
38. Donate L.E., Llorca O., Barcena M., Brown S.E., Dixon N.E., Carazo J.M (2000) **pH-controlled quaternary states of hexameric DnaB helicase** *J Mol Biol* **303**:383–393
39. Yang S., Yu X., VanLoock M.S., Jezewska M.J., Bujalowski W., Egelman E.H (2002) **Flexibility of the rings: structural asymmetry in the DnaB hexameric helicase** *J Mol Biol* **321**:839–849
40. Núñez-Ramírez R., Robledo Y., Mesa P., Ayora S., Alonso J.C., Carazo J.M., Donate L.E (2006) **Quaternary polymorphism of replicative helicase G40P: structural mapping and domain rearrangement** *J Mol Biol* **357**:1063–1076
41. Baretic D., Jenkyn-Bedford M., Aria V., Cannone G., Skehel M., Yeeles J.T.P (2020) **Cryo-EM Structure of the Fork Protection Complex Bound to CMG at a Replication Fork** *Mol Cell* **78**:926–940
42. Weng Z., Situ C., Lin L., Wu Z., Zhu J., Zhang R (2019) **Structure of BAI1/ELMO2 complex reveals an action mechanism of adhesion GPCRs via ELMO family scaffolds** *Nat Commun* **10**
43. Mastronarde D.N (2005) **Automated electron microscope tomography using robust prediction of specimen movements** *J Struct Biol* **152**:36–51
44. Punjani A., Rubinstein J.L., Fleet D.J., Brubaker M.A (2017) **cryoSPARC: algorithms for rapid unsupervised cryo-EM structure determination** *Nat Methods* **14**:290–296
45. Punjani A., Zhang H., Fleet D.J (2020) **Non-uniform refinement: adaptive regularization improves single-particle cryo-EM reconstruction** *Nat Methods* **17**:1214–1221
46. Varadi M. *et al.* (2022) **AlphaFold Protein Structure Database: massively expanding the structural coverage of protein-sequence space with high-accuracy models** *Nucleic Acids Res* **50**:D439–d444

47. Liebschner D. *et al.* (2019) **Macromolecular structure determination using X-rays, neutrons and electrons: recent developments in Phenix**. *Acta crystallographica. Section D Structural biology* **75**:861–877
48. Scheres S.H (2012) **RELION: implementation of a Bayesian approach to cryo-EM structure determination** *J Struct Biol* **180**:519–530
49. Zheng S.Q., Palovcak E., Armache J.P., Verba K.A., Cheng Y., Agard D.A. (2017) **MotionCor2: anisotropic correction of beam-induced motion for improved cryo-electron microscopy** *Nat Methods* **14**:331–332
50. Zhang K (2016) **Gctf: Real-time CTF determination and correction** *J Struct Biol* **193**:1–12
51. Nakane T., Kimanius D., Lindahl E., Scheres S.H (2018) **Characterisation of molecular motions in cryo-EM single-particle data by multi-body refinement in RELION** *eLife* **7**
52. Pettersen E.F., Goddard T.D., Huang C.C., Couch G.S., Greenblatt D.M., Meng E.C., Ferrin T.E (2004) **UCSF Chimera--a visualization system for exploratory research and analysis** *J Comput Chem* **25**:1605–1612
53. Emsley P., Lohkamp B., Scott W.G., Cowtan K (2010) **Features and development of Coot** *Acta Crystallogr D Biol Crystallogr* **66**:486–501
54. Afonine P.V., Grosse-Kunstleve R.W., Echols N., Headd J.J., Moriarty N.W., Mustyakimov M., Terwilliger T.C., Urzhumtsev A., Zwart P.H., Adams P.D (2012) **Towards automated crystallographic structure refinement with phenix.refine** *Acta Crystallogr D Biol Crystallogr* **68**:352–367

Article and author information

Zhuangfeng Weng

Shenzhen Key Laboratory for Systems Medicine in Inflammatory Diseases, School of Medicine, Shenzhen Campus of Sun Yat-sen University, Shenzhen, Guangdong 518107, China
ORCID ID: [0000-0002-7079-0293](https://orcid.org/0000-0002-7079-0293)

Jiefu Zheng

Shenzhen Key Laboratory for Systems Medicine in Inflammatory Diseases, School of Medicine, Shenzhen Campus of Sun Yat-sen University, Shenzhen, Guangdong 518107, China

Yiyi Zhou

Shenzhen Key Laboratory for Systems Medicine in Inflammatory Diseases, School of Medicine, Shenzhen Campus of Sun Yat-sen University, Shenzhen, Guangdong 518107, China

Zuer Lu

State Key Laboratory of Protein and Plant Gene Research, School of Life Sciences, Peking University, Beijing, China

Yixi Wu

Shenzhen Key Laboratory for Systems Medicine in Inflammatory Diseases, School of Medicine, Shenzhen Campus of Sun Yat-sen University, Shenzhen, Guangdong 518107, China

Dongyi Xu

State Key Laboratory of Protein and Plant Gene Research, School of Life Sciences, Peking University, Beijing, China
ORCID iD: [0000-0001-5711-2618](https://orcid.org/0000-0001-5711-2618)

Huanhuan Li

Department of Colorectal Surgery, The Sixth Affiliated Hospital, Sun Yat-sen University, Guangdong Institute of Gastroenterology, Guangdong Provincial Key Laboratory of Colorectal and Pelvic Floor Diseases, Guangzhou, Guangdong 510655, China

Huanhuan Liang

Pharmaceutical Sciences (Shenzhen), Sun Yat-sen University, Shenzhen, Guangdong 518107, China

For correspondence: lianghh26@mail.sysu.edu.cn

ORCID iD: [0000-0002-1803-3121](https://orcid.org/0000-0002-1803-3121)

Yingfang Liu

Shenzhen Key Laboratory for Systems Medicine in Inflammatory Diseases, School of Medicine, Shenzhen Campus of Sun Yat-sen University, Shenzhen, Guangdong 518107, China, Department of Colorectal Surgery, The Sixth Affiliated Hospital, Sun Yat-sen University, Guangdong Institute of Gastroenterology, Guangdong Provincial Key Laboratory of Colorectal and Pelvic Floor Diseases, Guangzhou, Guangdong 510655, China

For correspondence: liuyingf5@mail.sysu.edu.cn

ORCID iD: [0000-0002-1835-0397](https://orcid.org/0000-0002-1835-0397)

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Editors

Reviewing Editor

Volker Dötsch

Goethe University, Frankfurt am Main, Germany

Senior Editor

Volker Dötsch

Goethe University, Frankfurt am Main, Germany

Reviewer #1 (Public Review):

MCM8 and MCM9 are paralogues of the eukaryotic MCM2-7 proteins. MCM2-7 form a heterohexameric complex to function as a replicative helicase while MCM8-9 form another hexameric helicase complex that may function in homologous recombination-mediated long-tract gene conversion and/or break-induced replication. MCM2-7 complex is loaded during the low Cdk period by ORC, CDC6, and Cdt1, when the origin DNA may intrude into the central channel via the MCM2-MCM5 entry "gate". In the S phase, MCM2-7 complex is activated as CMG helicase with the help of CDC45 and GINS complex. On the other hand, it still remains unclear how MCM8-9 complex is loaded onto DNA and then activated.

In this study, the authors first investigated the cryo-EM structure of chicken MCM8-9 (gMCM8-9) complex. Based on the data obtained, they suggest that the observed gMCM8-9 structure might represent the structure of a loading state with possible DNA entry "gate". The authors further investigated the cryo-EM structure of human MCM8-9 (hMCM8-9) complex in the presence of the activator protein, HROB, and compared the structure with that obtained without HROB1, which the authors published previously. As a result, they suggest that MCM8-9 complex may change the conformation upon HROB binding, leading to helicase activation. Furthermore, based on the structural analyses, they identified some important residues and motifs in MCM8-9 complex, mutations of which actually impaired the MCM8-9 activity in vitro and in vivo.

Overall, the data presented would support the authors' conclusions and would be of wide interest for those working in the fields of DNA replication and repair. One caveat is that most of the structural data are shown only as ribbon model without showing the density map data obtained by cryo-EM, which makes accurate evaluation of the data somewhat difficult.

Addition after review of the revised manuscript: The authors have made a reasonable attempt to address the points raised by the reviewers, by which the paper is significantly improved.

- <https://doi.org/10.7554/eLife.87468.2.sa1>

Reviewer #2 (Public Review):

MCM8 and MCM9 together form a hexameric DNA helicase that is involved in homologous recombination (HR) for repairing DNA double-strand breaks. The authors have previously reported on the winged-helix structure of the MCM8 (Zeng et al. BBRC, 2020) and the N-terminal structure of MCM8/9 hexameric complex (MCM8/9-NTD) (Li et al. Structure, 2021). This manuscript reports the structure of a near-complete MCM8/9 complex and the conformational change of MCM8/9-NTD in the presence of its binding protein, HROB, as well as the residues important for its helicase activity.

The presented data might potentially explain how MCM8/9 works as a helicase. However, additional studies are required to conclude this point because the presented MCM8/9 structure is not a DNA-bound form and HROB is not visible in the presented structural data. Taking into these accounts, this work will be of interest to biologists studying DNA transactions.

A strength of this paper is that the authors revealed the near-complete MCM8/9 structure with 3.66Å and 5.21Å for the NTD and CTD, respectively (Figure 1). Additionally, the authors discovered a conformational change in the MCM8/9-NTD when HROB was included (Figure 4) and a flexible nature of MCM8/9-CTD (Figure S6 and Movie 1).

The revised version of "Structural and mechanistic insights into the MCM8/9 helicase complex" by Weng et al. includes only very minor changes in the text and incorporates two additional supplementary figures (S8 and S11) illustrating the size of MCM8/9 mutants.

In the previous version, I raised two important concerns that required addressing. 1) The presented structures exclusively depicted the unbound forms of DNA. It is crucial to elucidate the structure of a DNA-bound form. 2) The MCM8/9 activator, HROB, was not visible in the structural data. Although HROB induced a conformational change in MCM8/9-NTD, it is essential to visualize the structure of an MCM8/9-HROB complex.

The authors neither addressed nor provided new data in response to these issues. Consequently, I maintain my initial stance and have no further comments on the revised

version.

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Author Response

The following is the authors' response to the original reviews.

Reviewer #1 (Public Review):

MCM8 and MCM9 are paralogues of the eukaryotic MCM2-7 proteins. MCM2-7 form a heterohexameric complex to function as a replicative helicase while MCM8-9 form another hexameric helicase complex that may function in homologous recombination-mediated longtract gene conversion and/or break-induced replication. MCM2-7 complex is loaded during the low Cdk period by ORC, CDC6, and Cdt1, when the origin DNA may intrude into the central channel via the MCM2-MCM5 entry "gate". In the S phase, MCM2-7 complex is activated as CMG helicase with the help of CDC45 and GINS complex. On the other hand, it still remains unclear how MCM8-9 complex is loaded onto DNA and then activated.

In this study, the authors first investigated the cryo-EM structure of chicken MCM8-9 (gMCM89) complex. Based on the data obtained, they suggest that the observed gMCM8-9 structure might represent the structure of a loading state with possible DNA entry "gate". The authors further investigated the cryo-EM structure of human MCM8-9 (hMCM8-9) complex in the presence of the activator protein, HROB, and compared the structure with that obtained without HROB1, which the authors published previously. As a result, they suggest that MCM8-9 complex may change the conformation upon HROB binding, leading to helicase activation. Furthermore, based on the structural analyses, they identified some important residues and motifs in MCM8-9 complex, mutations of which actually impaired the MCM8-9 activity in vitro and in vivo.

Overall, the data presented would support the authors' conclusions and would be of wide interest for those working in the fields of DNA replication and repair. One caveat is that most of the structural data are shown only as ribbon model without showing the density map data obtained by cryo-EM, which makes accurate evaluation of the data somewhat difficult.

We thank the reviewer for the positive comments on our work. For evaluating all the structural data, in our revised manuscript, we have presented the density maps of the cryo-EM structures of the gMCM8/9 complex in supplementary figure S5 and S6. In addition, the 3D cryo-EM map of the gMCM8/9 complex and the hMCM8/9 NTD ring have been deposited to the EMDB database with accession number EMD-32346 and EMD-33989, respectively. The corresponding atomic models have been deposited at the RSCB PDB under the accession code 7W7P and 7Y0X, respectively. All these data have been released in May 2023.

Reviewer #2 (Public Review):

MCM8 and MCM9 together form a hexameric DNA helicase that is involved in homologous recombination (HR) for repairing DNA double-strand breaks. The authors have previously reported on the winged-helix structure of the MCM8 (Zeng et al. BBRC, 2020) and the Nterminal structure of MCM8/9 hexameric complex (MCM8/9-NTD) (Li et al. Structure, 2021). This manuscript reports the structure of a near-complete MCM8/9 complex and the conformational change of MCM8/9-NTD in the presence of its binding protein, HROB, as well as the residues important for its helicase activity.

The presented data might potentially explain how MCM8/9 works as a helicase. However, additional studies are required to conclude this point because the presented MCM8/9 structure is not a DNA-bound form and HROB is not visible in the presented structural data. Taking into these accounts, this work will be of interest to biologists studying DNA transactions.

A strength of this paper is that the authors revealed the near-complete MCM8/9 structure with 3.66Å and 5.21Å for the NTD and CTD, respectively (Figure 1). Additionally, the authors discovered a conformational change in the MCM8/9-NTD when HROB was included (Figure 4) and a flexible nature of MCM8/9-CTD (Figure S6 and Movie 1).

The biochemical data that demonstrate the significance of the Ob-hp motif and the N-C linker for DNA helicase activity require careful interpretation (Figures 5 and 6). To support the conclusion, the authors should show that the mutant proteins form the hexamer without problems. Otherwise, it is conceivable that the mutant proteins are flawed in complex formation. If that is the case, the authors cannot conclude that these motifs are vital for the helicase function.

A weakness of this paper is that the authors have already reported the structure of MCM8/9NTD utilizing human proteins (Li et al. Structure, 2021). Although they succeeded in revealing the high-resolution structure of MCM8/9-NTD with the chicken proteins in this study, the two structures are extremely comparable (Figure S2), and the interaction surfaces seem to be the same (Figure 2).

Another weakness of this paper is that the presented data cannot fully elucidate the mechanistic insights into how MCM8/9 functions as a helicase for two reasons. 1) The presented structures solely depict DNA unbound forms. It is critical to reveal the structure of a DNA-bound form. 2) The MCM8/9 activator, HROB, is not visible in the structural data. Even though HROB caused a conformational change in MCM8/9-NTD, it is critical to visualize the structure of an MCM8/9HROB complex.

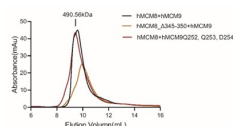
We appreciate the reviewer's comments on our work. Regarding the first weakness mentioned above, the previously reported cryo-EM structure of hMCM8/9 NTD ring was achieved with a resolution of 6.6 Å. At this level of resolution, we were only able to observe the overall shape of the structure and a partial representation of the protein's secondary structure. It is hard for us to discern any specific details regarding the interaction interface between MCM8 and MCM9. In this study, we solved the structure of gMCM8/9 NTD ring with a resolution of 3.67 Å. We believe that the higher resolution of gMCM8/9 NTD structure provides a significant advantage in analyzing the interaction surface between MCM8 and MCM9. This improved resolution has enabled us to gain valuable insights into the assembly mechanism of the MCM8/9 hexamer, representing a significant step forward in our understanding of the MCM8/9 helicase complex. In response to the second weakness raised by the reviewer, we fully agree with the reviewer that high-resolution structures of the MCM8/9 complex with DNA or HROB are necessary to elucidate the mechanism of this helicase complex. We are actively working towards obtaining these complex structures using cryo-EM and X-ray crystal diffraction.

Moreover, we would like to address the reviewer's concern regarding the mutant proteins used in the in vitro helicase assays. We have conducted additional experiments to confirm that these mutant proteins do not impair the formation of the MCM8/9 hexamer. Specifically, we performed size exclusion chromatography (SEC) analyses of the wild-type (WT) MCM8/9 complex, as well as MCM8 and MCM9 mutant proteins (Author response image 1). The results demonstrated that all the proteins behaved consistently and displayed similar SEC profiles during the purification process. Notably, the N-C linker deletion mutant (hMCM8_Δ369-377+MCM9_Δ283-287) combining the MCM8 and MCM9 N-C linker deletions also behaved

similarly with WT MCM8/9 (Author response image 2). These findings strongly suggest that the mutations in the OB-hps regions and the N-C linkers do not disrupt the hexamer formation of the MCM8/9 complex. Author response image 1 and Author response image 2 have been included into the supplementary figure S8 and S11, respectively.

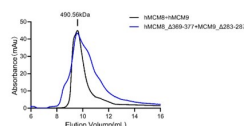
Author response image 1.

SEC profiles of WT and OB-hps mutants of MCM8/9 complex.



Author response image 2.

SEC profiles of WT and N-C linker mutant of MCM8/9 complex.



Reviewer #1 (Recommendations For The Authors):

I would like to provide some suggestions to improve the manuscript.

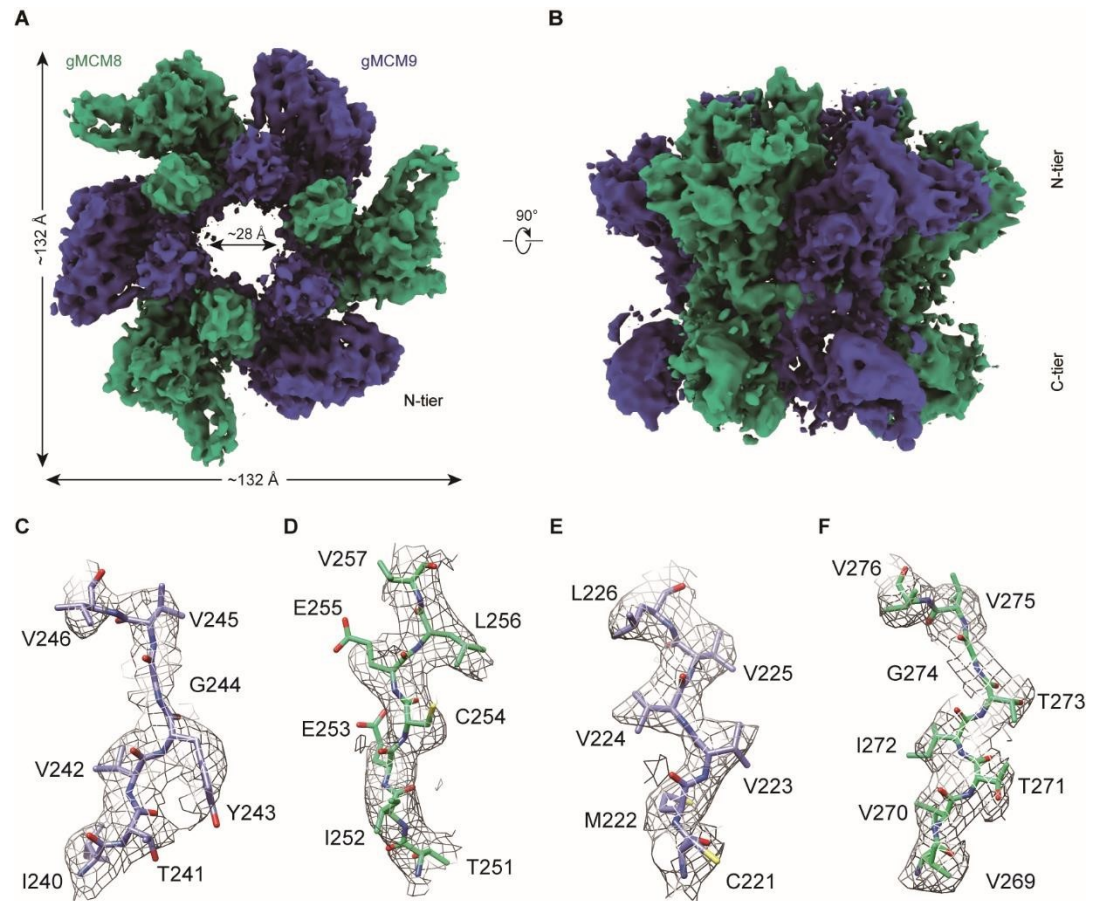
1. Throughout the manuscript, more density map data obtained by the cryo-EM should be shown for accurate evaluation of the data. For example, in Figure 1C, the authors state that inner channel of the gMCM8-9 hexamer is ~28 angstrom, apparently based on the ribbon model. This is not appropriate because the space upon ribbon model is not same as that upon the density map. For Figure 1B, they state that "The domain structures of gMCM8-9 fit well into their electron map". If so, please show the actual docking data. Also for Figure 2, the docking presentation between the side chains in the ribbon model and the density map should be shown.

We sincerely appreciate the reviewer for the constructive suggestions. In addition to releasing our structural data in the EMDB and PDB, we have also followed the reviewer's suggestions to included more density map data in the supplementary material. In fact, when calculating the diameter of the inner channel of the MCM8/9 hexamer, we also measured that upon the density map (Author response image 3. A and B), which is consistent with our report in our manuscript. To further evaluate the structure of MCM8/9, we have included additional docking structures based on the density map (Author response image 3. C-F). Moreover, for Figure 2, more docking presentation are provided and the key residues involved in the hydrophobic interactions were highlighted in a bold manner (Author response image 4). Author response image 3 and Author response image 4 have been included into the supplementary figure S5 and S6, respectively.

Author response image 3.

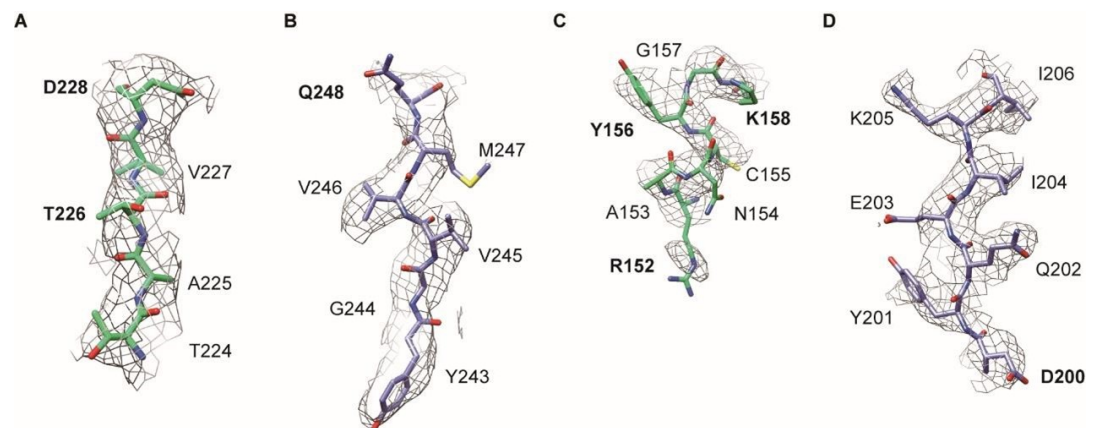
The cryo-EM structure of gMCM8/9. (A and B) Reconstructed cryo-EM map of gMCM8/9. The diameter of the inner channel of MCM8/9 was measured at ~28 Å. (C-F) Representative

regions of the cryo-EM structure of gMCM8/9 NTD are shown based on their density map. C, chain A (MCM9); D, chain B (MCM8); E, chain C (MCM9); F, chain D (MCM8).



Author response image 4.

Representative regions of the cryo-EM structure of gMCM8/9 NTD. (A and B), the region mediated hydrophobic interaction in figure 2B. A (MCM8), B (MCM9). (C and D), the region mediated hydrophobic interaction in figure 2C. C (MCM8), D (MCM9). The key residues were in bold.



1. Figures 4, 5, and 6: For helicase assay, more detailed experimental conditions (e.g. concentrations of DNA substrates and proteins used) should be presented. In addition, it should be described how Flag-hMCM8-9 complex (Figure 4C) was purified.

We sincerely appreciate the constructive suggestion provided by the reviewer. In the revised manuscript, we have included more experimental details in the helicase assays, including the concentrations of DNA substrates and proteins. The following paragraph describes the updated experimental procedure and also provided in the revise version of the manuscript.

Helicase assays: To prepare the substrate, the oligonucleotide (5'(dT)40GTTTTCCAGTCACGACG-TTGTAACGACGGCCAGTGCC-3') containing a 40 nt region complementary to the M13mp18(+) strand and a 40 nt oligo-dT at the 5' end was labeled at the 3' terminus with [α -³²P] dCTP (Perkin Elmer) and annealed to the single-stranded DNA M13mp18 (24). 0.1 nM (in molecules) DNA substrates were respectively mixed with 5 μ g recombinant MCM8/9 complex and its mutants as indicated within each 15 μ l volume reaction in the helicase buffer (25 mM HEPES, pH 7.5, 1 mM magnesium acetate, 25 mM sodium acetate, pH 5.2, 4 mM ATP, 0.1 mg/ml BSA, 1 mM DTT). 2.5 μ g HROB was used as an activator. To avoid re-annealing, the reaction was supplemented with a 100-fold unlabeled oligonucleotide. The reactions were then incubated at 37 °C for 60 min and stopped by adding 1 μ l of stop buffer (0.4% SDS, 30 mM EDTA, and 6% glycerol) and 1 μ l of proteinase K (20 mg/ml, Sigma) into the reaction for another 10 min incubation at 37 °C. The products were separated by 15% polyacrylamide gel electrophoresis in 1 $\times$ TBE buffer and analyzed by the Amersham typhoon (Cytiva).

In addition, to describe the expression of Flag-hMCM8/9 complex in Figure 4C, we have included the Pull-Down Assay in the “Material and Methods” section. The description is as follow: The HEK293T cells transfected with Flag-hMCM8/9-FL or Flag-hMCM8/9-NTD were cultured overnight and washed twice with cold phosphate-buffered saline (PBS). Cell pellets were resuspended with lysis buffer (20 mM Tris, pH7.5, 150 mM NaCl, 5mM EDTA, 0.5% NP-40, 10% glycerol, protease inhibitor cocktail (Roche, 04693132001)). After incubation for 45 min at 4°C with gentle agitation, the whole-cell lysates were collected by centrifugation (12,000 $\times$ g for 15 min, at 4 °C). GST beads coupled with 2 μ g GST-HROB or GST alone were then incubated with an equal volume of above HEK293T cell lysates at 4°C for 4h. The beads were washed four times with lysis buffer. Proteins bound to the beads were separated by SDS-PAGE and subsequently immunoblotted with anti-Flag antibody (Cytiva).

1. Figure 3C: This is just an assumed model. Please clearly state it in the manuscript.

We appreciate the reviewer’s comment. We guess the reviewer is referring to Figure 5C. As Figure 3C depicts the top view of the gMCM8/9 hexamer structurally aligned with the MCM2-7 double hexamer (wheat) by aligning their respective C-tier ring. On the other hand, Figure 5C represents an assumed model where we docked a forked DNA fragment into the central channel of the gMCM8/9 hexamer. To address this assumed model, we have made the following clarification in the revised manuscript: “We artificially docked a forked DNA into the central channel to generate a gMCM8/9-DNA model and found that the OB-hps of gMCM8 are capable to closely contact with it and insert their highly positively charged terminal loops into the major or minor grooves of the DNA strand, implying that they could be involved in substrate DNA processing and/or unwinding (Figure 5C)”.

1. Figure S1, C and D: The coloring of the gMCM8-9 CTD appears to show higher resolution than the NTD. May this be mispresentation?

We appreciate the reviewer's valuable feedback, and we have thoroughly re-evaluated Figure S1C and D. At the beginning, the local resolution distributions of the gMCM8/9 NTD and gMCM8/9 CTD were calculated using CryoSPARC. Upon re-examination, we found that the density maps of the gMCM8/9 CTD may be lower than 3.66 Å, because the density map of the gMCM8/9 CTD does not reveal more structural details than what is observed in the gMCM8/9 NTD. Thus, although the map shown in Figure S1D may appear to show a greater distribution of high-resolution regions, we would like to clarify that this discrepancy could be attributed to an optical illusion. We thank the reviewer for bringing this to our attention.

1. *Figure S9: Is the "mean resolution" 5.21 angstrom identical to the Gold standard FSC? If not, please estimate the resolution using FSC, like other maps in this paper.*

We thank the reviewer for the constructive suggestion. In response to this feedback, we would like to clarify the resolution estimation process for the gMCM8/9 CTD. Initially, we calculated the resolution of the gMCM8/9 CTD using the gold standard Fourier shell correlation (FSC) method, which yielded a resolution of 3.66 Å. However, upon further analysis, we identified an issue with the GSFSC Resolution curves, which led to an overestimation of the resolution based on the density map of the gMCM8/9 CTD. To ensure a more reliable and accurate estimation, we employed the Phenix software package to calculate the mean resolution during the refinement process of the gMCM8/9 CTD structure. The calculated mean resolution was determined to be 5.21 Å, which aligns more reasonably with the characteristics of the density map. To address any potential misunderstandings and provide clarity, we have explicitly labeled and described the evaluation process for this mean resolution in the "Single particle data processing" section of the Materials and Methods.

Minor points:

1. *Throughout the manuscript, there are several typographical and grammatical errors, which should be corrected. For example, in "Introduction", "GNIS complex" should be "GINS complex".*

We thank the reviewer for pointing out the typographical and grammatical errors. We have corrected the grammar errors and polished our manuscript with the help of native speakers.

Reviewer #2 (Recommendations For The Authors):

1. *"During HR repair, MCM8/9 was rapidly recruited to the DNA damage sites and colocalized with the recombinase Rad51 (21). It also interacted with the nuclease complex MRN (MRE11RAD50-NBS1) and was required for DNA resection at DSBs to facilitate the HR repair (Introduction)."*

There is a debate about whether MCM8/9-HROB colocalizes with RAD51 and whether it works upstream or downstream of RAD51 (Park et al. MCB, 2013; Lee et al. Nat Commun., 2015; Lutzmann et al. Mol Cell, 2012; Nishimura et al. Mol Cell, 2012; Natsume et al. G&D, 2017; Hustedt et al. G&D, 2019; Huang et al. Nat Commun., 2020).

We completely agree with the reviewer that previous studies have reported contradictory results regarding to the function of MCM8/9 in homologous recombination. Based on the structure information of MCM8/9, now we do not have direct evidence to resolve the ongoing debate. Nonetheless, based on our findings, we speculate that the MCM8/9 complex is likely involved in multiple steps within the process of homologous recombination. The structural insights provided by our study serve as a foundation for further investigations and may

contribute to a better understanding of the complex and multifaceted roles of MCM8/9 in homologous recombination repair.

1. I noted that the BioRxiv version 1 (<https://www.biorxiv.org/content/10.1101/2022.01.26.477944v1?versioned=true>) contains a near-complete MCM8/9 with human protein based on the crystal analysis. Because its structure is comparable to chicken MCM8/9 revealed by cryo-EM, I highly suggest including this data in the manuscript.

We would like to thank the reviewer for this suggestion. The resolution of the hMCM8/9 crystal structure presented in our previous BioRxiv version is 6.6 Å, which is a little low. Moreover, it cannot provide more information than the present cryo-EM structures of MCM8/9. We are dedicated to optimizing the crystal quality and implementing strategies to enhance the resolution of the structure. We hope to present an improved crystal structure of hMCM8/9 in our forthcoming article.