RESEARCH ARTICLE

Cryo-EM structure of an early precursor of large ribosomal subunit reveals a half-assembled intermediate

Dejian Zhou, Xing Zhu, Sanduo Zheng, Dan Tan, Meng-Qiu Dong, Keqiong Ye

1 Graduate School of Peking Union Medical College and Chinese Academy of Medical Sciences, Beijing 100730, China
2 Key Laboratory of RNA Biology, Institute of Biophysics, CAS Center for Excellence in Biomacromolecules, Chinese Academy of Sciences, Beijing 100101, China
3 National Institute of Biological Sciences, Beijing 102206, China
4 University of Chinese Academy of Sciences, Beijing 100049, China

Correspondence: yekeqiong@ibp.ac.cn (K. Ye)

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ABSTRACT

Assembly of eukaryotic ribosome is a complicated and dynamic process that involves a series of intermediates. It is unknown how the highly intertwined structure of 60S large ribosomal subunits is established. Here, we report the structure of an early nucleolar pre-60S ribosome determined by cryo-electron microscopy at 3.7 Å resolution, revealing a half-assembled subunit. Domains I, II and VI of 25S/5.8S rRNA pack tightly into a native-like substructure, but domains III, IV and V are not assembled. The structure contains 12 assembly factors and 19 ribosomal proteins, many of which are required for early processing of large subunit rRNA. The Brx1-Ebp2 complex would interfere with the assembly of domains IV and V. Rpf1, Mak16, Nsa1 and Rrp1 form a cluster that consolidates the joining of domains I and II. Our structure reveals a key intermediate on the path to establishing the global architecture of 60S subunits.

KEYWORDS ribosome assembly, cryo-EM, pre-60S ribosome, nucleolar

INTRODUCTION

Correct formation of the ribosome is essential for protein synthesis in cells. The ribosome is composed of a small 40S subunit and a large 60S subunit (SSU and LSU) in eukaryotes. Ribosome assembly is a highly complicated process that engages more than 200 protein assembly factors (AFs) and many snoRNAs (Woolford and Baserga, 2013). These factors temporarily associate with ribosomal subunits and drive their maturation. In the yeast S. cerevisiae, assembly of both subunits begins in the nucleolus with the transcription of a long precursor RNA (pre-rRNA) that encodes 18S, 5.8S and 25S rRNAs and four external and internal transcribed spaces (ETS and ITS). The 5′ region of the pre-rRNA is co-transcriptionally assembled into the 90S pre-ribosome or SSU processome, which is the early precursor of SSU (Dragon et al., 2002; Grandi et al., 2002; Osheim et al., 2004; Chaker-Margot et al., 2015; Zhang et al., 2016). Following cleavage of the pre-rRNA at the A0, A1 and A2 sites, the 90S pre-ribosome is released from the pre-rRNA and transformed into a pre-40S ribosome. Cryo-EM structures of 90S pre-ribosome show that the nascent 40S subunit is assembled into several isolated units that have yet to acquire the global architecture of 40S subunit (Komprobst et al., 2016; Barandun et al., 2017; Chaker-Margot et al., 2017; Cheng et al., 2017; Sun et al., 2017).

The 60S subunit is composed of 25S, 5.8S and 5S rRNAs, and 46 ribosomal proteins (RPLs). The 25S/5.8S rRNAs are divided into six domains that intertwine to form a monolithic structure (Ben-Shem et al., 2011; Klinge et al., 2011). The earliest pre-60S ribosome is co-transcriptionally...
formed on the 3' LSU region of pre-rRNA in a stepwise manner (Chen et al., 2017). The pre-60S progressively matures as it transits from the nucleolus to the nucleoplasm and cytoplasm. The maturation of pre-60S is coupled to the dynamic association of approximately 90 assembly factors, hierarchical incorporation of RPLs and sequential processing of ITS1 and ITS2. The development of pre-60S at the nucleoplasm and cytoplasm has been characterized to great extents, particularly by recent cryo-EM studies (Pena et al., 2017). An early nucleoplasmic state is characterized by a foot structure assembled on the partially processed ITS2 and an immature central protuberance (CP) (Bradatsch et al., 2012; Leidig et al., 2014; Wu et al., 2016). Further evolution of pre-60S leads to removal of the foot, maturation of the CP and association of the Rix1-Ipi1-Ipi2 subcomplex and the AAA-ATPase Rea1 (Barrio-Garcia et al., 2016). Rea1 hydrolyzes ATP and drives the release of Ytm1 and Rsa4 (Ulbrich et al., 2009; Bassler et al., 2010). The pre-60S acquires competence for nuclear export at the late nucleoplasmic stage and undergoes final maturation in the cytoplasm (Matsuo et al., 2014; Greber et al., 2015; Weis et al., 2015; Ma et al., 2017; Malyutin et al., 2017).

In all previously determined cryo-EM structures of pre-60S, the structural core has already assumed the shape of mature subunit. Therefore, the key question of how the global architecture of LSU is established is still unknown. In this study, we determine the cryo-EM structure of pre-60S at an early nucleolar state. The structure is only half-assembled at domains I, II and VI and represents a key intermediate in the formation of global architecture of LSU.

**RESULTS**

**Structure determination**

We affinity purified pre-60S using the bait protein Rpf1 fused to a C-terminal tandem affinity purification (TAP) tag. Mass spectrometry analysis showed that the Rpf1-TAP particle displayed a similar protein composition as the previously characterized nucleolar Ssf1-TAP and Nsa1-TAP particles (Fatica et al., 2002; Kressler et al., 2008; Chen et al., 2017) (Fig. S1, Supplementary Dataset 1). Cryo-EM images were collected in a Titan Krios 300 kV electron microscope equipped with a direct electron detector (Fig. S2). After three rounds of 3D classification, 98,155 particles were selected to reconstruct a density map at an overall resolution of 3.65 Å (Figs. 1A and S3). Large side chains of protein and RNA bases are resolved in the structure core, whereas some peripheral regions including many AFs are of lower resolution (Figs. S3 and S4). The structure was built based on the cryo-EM structure of Nog1-TAP pre-60S (Wu et al., 2016), crystal structures (Brx1, Ebp2, Nsa1), homology models (Has1 and Rpf1) and *de novo* modeling (Mak16 and Rpf1) (Fig. S4 and Table S1). The Rpf1-TAP sample was also analyzed by chemical crosslinking and mass spectrometry (CXMS), yielding 28 intermolecular and 110 intramolecular crosslinks (Supplementary Dataset 2, Fig. S5). These intermolecular crosslinks assisted the assignment of AFs (Table S1) and also suggested the position of unmodeled AFs, such as Dbp10, Drs1, Nop2 and Mak11 (Fig. S5). The final model was refined in real space and displayed good geometry (Table S2).

**Overall structure**

The cryo-EM density map and structural model of Rpf1-TAP pre-60S are shown in Fig. 1. The most striking feature of the structure is that only half of LSU is present. Domains I, II and VI assemble into a native-like substructure, but domains III, IV and V are totally absent (Fig. 2). A few peripheral elements in domain II that form the CP (H38, ES12) and the P stalk (H43, H44) and interact with domains III, VI and V (H33-H35) are also disordered (Fig. 2E). Therefore, the structure represents an early assembly intermediate before the global architecture of 60S is established. A total of 12 AFs and 19 RPLs were modeled in the map (Figs. 1B, 2C and 2D). Five AFs, namely, Nop7, Rip7, Nop15, Cic1 and Tif6, are also present in the early nucleoplasmic pre-60S (Bradatsch et al., 2012; Leidig et al., 2014; Wu et al., 2016) and adopt similar conformation in the Rpf1-TAP pre-60S. Seven AFs Has1, Brx1, Ebp2, Rpf1, Mak16, Nsa1 and Rpf1 were newly located (Fig. S3).

**Assembly of ITS2**

The N-terminal region of ITS2 is associated with Cic1, Nop15, Rip7 and Nop7, forming the foot structure (Figs. 1 and 3A). The foot structure is similar to that of the nucleoplasmic pre-60S structure (Bradatsch et al., 2012; Leidig et al., 2014; Wu et al., 2016), except that Nop53 is not present (Fig. 3B). Nop53 recruits the exosome for processing 7S pre-rRNA (Thoms et al., 2015; Falk et al., 2017). Nop53 is of low abundance in the Rpf1-TAP particle (Fig. S1) and likely associates at a later stage.

**Has1**

Has1 is a DEAD-box RNA helicase involved in both small and large ribosomal subunit assembly (Emery et al., 2004; Dembowski et al., 2013). RNA helicases are thought to unwind RNA structures or disassemble RNA-protein complexes, hence driving the evolvement of pre-ribosomes. In the cryo-EM structure, Has1 is situated next to Cic1 and interacts with H16 of 25S rRNA through its two RecA domains (Fig. 3A). DEAD-box helicases unwind short RNA duplexes by local strand separation (Yang et al., 2007). In a proposed mechanistic model (Mallam et al., 2012), RecA2 first associates with the RNA duplex, and binding of ATP to RecA1 then induces closure of two RecA domains, causing RecA1 to displace one RNA strand. In the pre-60S structure, H16 is bound to Has1 at a separate site from where single- and double-stranded RNAs are bound to DEAD-box.
helicas (Fig. 3C and 3D) (Del Campo and Lambowitz, 2009; Mallam et al., 2012). In addition, the ATP-binding pocket of Has1 is empty, and its two RecA domains are not arranged in the same closed conformation as in ternary complex structures of helicase, ssRNA and ATP analog. These structural features suggest that H16 serves as a docking site rather than an unwinding substrate for Has1.

Structure of the Brx1 and Ebp2 complex

Brx1 is a Brix domain protein that interacts with Ebp2 (Wehner and Baserga, 2002; Shimoji et al., 2012). We determined the crystal structure of Brx1 (residues 26–255) in complex with Ebp2 (residues 195–293) at 2.3 Å resolution with de novo phasing (Table S3). Brx1 adopts an α/β fold with pseudo 2-fold symmetry (Fig. 4A), similar to other Brix domain proteins (Ng et al., 2005; Asano et al., 2015; Kharde et al., 2015; Madru et al., 2015; Sa-Moura et al., 2017). The structural core of Brx1 is composed of two curved β-sheets that form a half-open β-barrel sandwiching two α-helices (α1 and α3). The surface of the N-terminal sheet holds helix α2, whereas the surface of the C-terminal sheet makes an intermolecular interaction with helix α2’ of Ebp2. In addition, the Ebp2 sequences that flank helix α2’ wrap around the Brx1 structure, forming an extensive interface. The α2’ helix of Ebp1 packs against the N-terminal sheet of Brx1, and the β1’ strand of Ebp2 expands the C-terminal sheet of Brx1.

In the cryo-EM map, the Brx1-Ebp2 complex perfectly fits into a density bound to domain I (Figs. 1 and S4B). Brx1 interacts with a flat surface formed by H13 and H21. The position of Brx1 would clash with H68 and H88 in domains IV and V in the fully assembled 60S (Fig. 3B). Thus, the Brx1-Ebp2 complex must be released from the current position for ribosome assembly to proceed.

There are six Brix domain proteins, namely, Imp4, Rpf2, Rpf1, Brx1, Ssf1 and Ssf2, in the yeast S. cerevisiae. Imp4 is involved in small subunit assembly, whereas the others function in large subunit assembly. The structures of Rpf2, Imp4, Rpf1 (see below) and Brx1 have now been determined in the context of pre-ribosomes (Leidig et al., 2014; Komprobst et al., 2016; Wu et al., 2016; Barandun et al., 2017; Chaker-Margot et al., 2017; Cheng et al., 2017; Sun et al., 2017), allowing for comparison of their conserved and variable features in protein and RNA recognition (Fig. 4B–E). In all four proteins, an α-helix of their partner proteins binds with the C-terminal sheet of the Brix domain. Outside the core interface, the intermolecular interaction is highly variable. Brx1 and Rpf2 form an extensive interface with their binding partners Ebp2 and Rsr1, but Rpf1 and Imp4 primarily bind only a single α-helix of Mak16 and Mpp10. The RNA binding interface is also extremely diverse for the four Brix proteins. No consensus sequence or structure is apparent for the bound RNAs. Nevertheless, one protein region that consistently binds RNA is located at the junction between strand β1 and helix α1 and the turn connecting strands β3 and β4. The diverse interactions with protein and RNA allow each Brix protein to play specific roles in ribosome assembly.
D2 factors

Rrp1, Mak16, Rpf1 and Nsa1 form a cluster bound to a junction region of domains I and II at the solvent side (Fig. 5A–F). The four proteins are co-assembled into a pre-rRNA fragment ending at domain II (termed D2 factors) (Chen et al., 2017), consistent with their binding sites. These AFs are not present in the nucleoplasmic pre-60S and Nsa1 is released by the AAA ATPase Rix7 (Kressler et al., 2008).

Mak16 occupies a central position of the cluster and is mostly buried. Mak16 is composed of a knot-like N-terminal domain (NTD), a two-layered middle domain (MD) and an extended C-terminal helix (Fig. 5B). The NTD fits snugly into a cavity formed by H45 and H46 and ribosomal proteins L4 and L32. The MD is covered by the first helix of expansion segment 7 (ES7a) RNA. The C-terminal α-helix of Mak16 binds to the Brix domain protein Rpf1. This interaction was also demonstrated by yeast 2-hybrid, in vitro pull-down and gel filtration assays (Fig. S6) (McCann et al., 2015; Bassler et al., 2017). In addition to binding Mak16, Rpf1 also binds L32, Nsa1 and the base of helix ES7a. Nsa1 contains a WD domain (Lo et al., 2017) and is docked at H19, ES7a and Rpf1. Rrp1 folds into a super-helical structure and contacts L4, L18, the N-terminus of L7, Mak16 and RNA helix ES7a. By binding both domains I and II, these D2 factors appear to

Figure 2. Structure of rRNAs and RPLs in Rpf1-TAP pre-60S and mature 60S subunit. (A and B) Structure of rRNAs in Rpf1-TAP pre-60S (A) and mature 60S (B). The subunit and solvent side views are shown. Domains I–VI of 25S are colored slate, cyan, green, orange, pink and purple, respectively. ITS2, 5.8S and 5S are colored dark blue, red and brown, respectively. Central protuberance (CP), peptidyl transferase center (PTC), polypeptide exit tunnel and P-stalk are labeled. (C and D) RPLs in Rpf1-TAP pre-60S (C) and mature 60S (D). The solvent side view is shown. RPLs present in pre-60S are color coded and those missing in pre-60S are shown in black in (D). The early, middle and unclassified RPLs are labelled in red, blue and black, respectively. (E) Secondary structure model of 5.8S, ITS2 and 25S RNA. The modeled regions are colored as in (A) and (B) and the unmodeled regions are colored gray.
strengthen the joining of the two domains at early assembly stages when many interactions in the mature subunit are missing.

The ribosomal structure at the D2 factor binding region closely resembles the mature subunit (Fig. 5F). However, in the mature subunit, the N-terminal tail of L6 inserts into the cavity initially occupied by Mak16, and ES7a becomes disordered in the absence of AF contacts. ES7a is a eukaryote-specific element in 25S rRNA and appears to function in ribosome assembly, which differs dramatically between bacteria and eukaryotes, rather than in protein translation.

**DISCUSSION**

We have determined the cryo-EM structure of pre-60S at an early nucleolar state. The structure demonstrates that the global architecture of 60S is first established by domains I, II and VI, which constitute the major part of the solvent shell of 60S. Part of the solvent shell (domain III) and the entire inner layer at the subunit interface (domains IV and V) have not been integrated. The functional sites of 60S, the peptidyl transferase center and the polypeptide exit tunnel, are not assembled at this stage. The absence of domains III, IV and V does not mean that these domains are completely unfolded. They could fold and associate with AFs and RPLs (Chen et al., 2017) but are too flexible to be visualized in the cryo-EM structure. The Rpf1-TAP particle contains ~40 AFs of appreciable abundance (Fig. S1). Those AFs not present in our structure may be flexible or associate with other states of pre-60S. The map also contains unassigned structures and unmodeled weak densities.

In the Rpf1-TAP pre-60S, domains I, II and VI together with 19 RPLs are tightly packed as in the mature subunit. All AFs barely interfere on the formed ribosomal structure. By contrast, in the 90S pre-ribosome, all four domains of the 40S subunit are isolated and are organized by extensive interactions with a large number of AFs, the 5' ETS and U3 snoRNA (Komprobst et al., 2016; Barandun et al., 2017; Chaker-Margot et al., 2017; Cheng et al., 2017; Sun et al., 2017).
Many AFs and RPLs have been classified according to their role in pre-rRNA processing (Sahasranaman et al., 2011; Talkish et al., 2012; Woolford and Baserga, 2013; Gamalinda et al., 2014; de la Cruz et al., 2015). A3-factors (or early factors) are required for processing of 27SA3 to 27SB pre-rRNA, B-factors (or middle factors) are required for C2 cleavage of ITS2 and late factors are required for processing of 7S pre-rRNA. Based on functional classification, binding dependence and structural location of RPLs, a hierarchical model has been proposed for 60S assembly with assembly beginning at the solvent side (Gamalinda et al., 2014). The Rpf1-TAP pre-60S structure is half-assembled at the solvent side and contains 8 of 13 AFs and all of 12 RPLs that are classified as early A3-factors (Figs. 2C and S1). This suggests that the structure present in this study corresponds to the predicted early assembly intermediate and is required for ITS1 processing.

By analyzing the association of AFs and RPLs with plasmid-derived pre-rRNA fragments of increasing length, the earliest precursor of 60S was found to form in a stepwise manner (Chen et al., 2017). The D2 factors Rpf1, Mak16, Nsa1 and Rrp1, which are initially recruited by a pre-rRNA fragment ending at domain II, are found to bind at the junction of domains I and II. This suggests that domains I and II
could already associate despite the absence of other domains. This suggests a sequential assembly model for the Rpf1-TAP pre-60S. Domains I and II first combine into a structural core that is late joined by domain VI.

Note added

While this work was prepared for publication, the cryo-EM structures of nucleolar pre-60S in multiple states were reported (Kater et al., 2017). Our structure is mostly similar with state B derived from Nsa1 and Ytm1 affinity purification.

MATERIALS AND METHODS

Purification of Rpf1-TAP particles

The Rpf1-TAP strain (BY4741, RPF1-TAP::HisMX3) was purchased from Open Biosystems. Six liters of cells were grown at 30 °C in YPD medium (1% yeast extract, 2% peptone, 2% glucose) to OD_{600} of 1–2. The cells were harvested and re-suspended in lysis buffer (20 mmol/L HEPES-KOH, pH 8.0, 110 mmol/L KOAc, 40 mmol/L NaCl, 0.1% NP-40) supplemented with one tablet of EDTA-free protease inhibitor cocktail (Roche). The cells were lysed by a high pressure cell disruptor (JNBIo) and clarified by centrifugation at 6,000 × g for 10 min. The supernatant was filtered through a 0.45 µm membrane and incubated with 50 mg of IgG-coated magnetic Dynabeads (Invitrogen) for 30 min. The beads were washed with lysis buffer twice and incubated with TEV protease in cleavage buffer (10 mmol/L Tris-HCl, pH 8.0, 100 mmol/L NaCl and 2 mmol/L DTT) at 4 °C for 4–8 h. The released complex was concentrated to ∼50 µL.

Cryo-EM data collection and image processing

To prepare vitrified specimen, 3 µL of sample (OD_{260} = 3–5) was applied to glow-charged holey carbon grids (GiG R422) at 4 °C and 100% humidity using an FEI Vitrobot Mark IV. The grids were blotted for 1–3 s and rapidly plunged into liquid ethane. The cryogenic sample was first screened on a Talos F200C 200kV electron microscope equipped with a Ceta camera (FEI). A dataset was collected with a pixel size of 2.27 Å. Images were processed with RELION (Scheres, 2012a, b). Particles were picked, cleaned and subjected to 3D classification using the mature yeast 40S ribosome...
as the initial model (Ben-Shem et al., 2011). A density map with clear structural feature was obtained at 11.7 Å resolution. This map was used as the initial model for subsequent image processing.

High-resolution images were recorded on a Titan Krios (FEI) operated at 300 kV and equipped with a Falcon III detector. A total of 1983 images were collected with SerialEM (Mastronarde, 2005) in low-dose mode at a pixel size of 1.38 Å. Images were recorded in 23 movie frames with an exposure time of 1.5 s and a total dose of 30 e Å² and corrected for motion with MOTIONCORR (Li et al., 2013). Contrast transfer function (CTF) values were measured with CTFFIND4 (Rohou and Grigorieff, 2015). A total of 1,786 good-quality images with a defocus range of 1–4 µm were chosen for particle picking in RELION. Particles were extracted with a box size of 360 pixels and downsized 4-fold for 3D classification. Three rounds of 3D classification were performed with a mask of 400 Å diameter (Fig. S2). The classes with high resolution and similar features were combined for further classification or refinement. The final 3D refinement was conducted on 98,155 particles with the gold-standard Fourier shell correlation (FSC) method (Scheres and Chen, 2012). A density map at 3.65 Å resolution was obtained after post-processing, correction of modulation transfer function of the detector and B-factor sharpening. FSC curves were calculated from data half-sets using soft masks and corrected for the effect of the mask using high-resolution noise substitution. The reported resolution was based on the gold-standard FSC = 0.143 criterion. Local resolution was estimated with ResMap (Kucukelbir et al., 2014). To calculate FSC between model and map, the model was converted into a density map in Chimera, which was resampled to the grid of the map used for refinement. FSC between two maps was calculated with relion_image_handler. For cross-validation, the model was re-refined against a map reconstruction from half the data (Amunts et al., 2014).

Model building

The cryo-EM structure of Nog2-TAP pre-60S (PDB code 3JCT) was fit into the density as the starting model (Wu et al., 2016). The RNA and protein components without density were removed. The unmodeled density was assigned with assistance of the BALBES-MOLREP pipeline (Brown et al., 2015) and CXMS data. The crystal structures of the Brx1 and Ebp2 complex determined here and the Nsa1 structure (Lo et al., 2017) were fitted as rigid body. Homology models of Rpf1 and Has1 were created by SWISS-MODEL (Biasini et al., 2014). Rrp1 and Mak16 were manually built. A few unassigned densities were modeled as poly-alanine chains, and additional weak densities were not modeled. Nucleotides 438–493 of helix ES7a were modeled based on the structural prediction of RNAComposer (Popenda et al., 2012). The structure was further adjusted in COOT (Emsley and Cowtan, 2004) and refined in real space with secondary structure and geometry restraints using PHENIX (Adams et al., 2010). Structural figures were prepared with PyMOL 1.7 (Schrödinger, LLC) and Chimera (Petterson et al., 2004).

Expression and purification of the Brx1 and Ebp2 complex

The genes of Brx1 (total 291 residues) and Ebp2 (total 427 residues) were amplified from yeast genomic DNA and cloned with the transfer-PCR approach (Erijman et al., 2011). Ebp2 and its fragments were cloned into a modified pET28a vector (Novagen) and fused to an N-terminal His6-Smt3 tag. Brx1 and its fragments were cloned into a modified pETDuet-1 vector and fused to an N-terminal His6-GST tag followed by a PreScission cleavage site. The constructed plasmids were verified by DNA sequencing. Native or selenomethionine (SeMet)-labeled Brx1 and Ebp2 were co-expressed in the E. coli Rosetta 2(DE3) strain and co-purified with HisTrap, heparin and gel filtration chromatography according to the previously described procedure (Zheng and Ye, 2014). The interaction between the two proteins was confirmed by GST pull-down assay.

Crystallization and structural determination of the Brx1 and Ebp2 complex

Several fragments of Ebp2 and Brx1 were tested for crystallization. High quality crystals were grown from a complex containing residues 186–295 of Ebp2 and residues 26–259 of Ebp2. Both the native and SeMet-labeled complexes (15 mg/mL in 10 mmol/L Tris-HCl, pH 8.0 and 500 mmol/L NaCl) were crystallized in 0.1 mol/L Bis-Tris, pH 5.5 and 1.8 mol/L ammonium sulfate using the vapor diffusion hanging drop method. The crystals were cryo-protected in 20% glycerol made in the reservoir solution and flash frozen in liquid nitrogen.

Diffraction data were collected at the beamLine BL17U of Shanghai Synchrotron Radiation Facility and processed with HKL2000 (Otwinowski and Minor, 1997). Two types of crystals that belonged to C2 or P2₁ space group were obtained at the same condition. The structure was first solved by the single-wavelength anomalous diffraction (SAD) method based on a C2, Se-derivative dataset at 2.8 Å resolution using SHARP (Vonrhein et al., 2007) (Table S3). The resultant electron density map was of moderate quality. The experimental phases were iteratively combined with the phases calculated from a partially built model to improve the map. A better dataset at 2.3 Å resolution was later collected on the native P2₁ crystal and used for final refinement. The model was built in COOT (Emsley and Cowtan, 2004) and refined with PHENIX (Adams et al., 2010). The final model contains two copies of the complex composed of Ebp2 residues 195–293, Brx1 residues 26–65, 70–195 and 210–255, 308 water molecules and one sulfate ion.

Mass spectrometric analysis

Mass spectrometric analysis was conducted as previously described (Zhang et al., 2016). The total spectral counts per 100 residues (SCPHR) were calculated for each identified protein and further normalized against Brx1, Ebp2, Erb1, Ytm1, Nop7, Cic1 and Has1, yielding the relative spectral abundance factor (RSAF) (Supplementary Dataset 1).

CXMS

CXMS was performed as described (Sun et al., 2017). The Rpf1-TAP sample containing ~10 µg of total proteins was crosslinked with BS3 or DSS, precipitated with acetone and digested by trypsin. The sample was analyzed by LC-MS/MS on an EASY-nLC 1000 system interfaced to a Q-Exactive HF mass spectrometer (Thermo Fisher Scientific). Cross-linked peptides were identified with pLink (Yang et al., 2012).
Protein interaction analysis of Rpf1 and Mak16

Yeast two-hybrid and GST pull-down assays were conducted as described (Zheng et al., 2014). Rpf1 was cloned into a modified pET28a vector (Novagen) and fused to an N-terminal His$_6$-Smt3 tag. Mak16 and its fragments were cloned into a modified pETDuet-1 vector and fused to an N-terminal His$_6$-GST tag followed by a PreScission cleavage site. For GST pull-down assay, Rpf1 was expressed and purified by HisTrap chromatography, Ulp1 cleavage of the His$_6$-Smt3 tag, heparin and gel filtration chromatography. Mak16 was purified by HisTrap chromatography without cleavage of the tag. To purify the complex, Rpf1 and Mak16 were co-expressed and co-purified according to the previously described procedure (Zheng and Ye, 2014).

Accession numbers

The cryo-EM density map, coordinates and structural factors have been deposited to EMDB and PDB under accession numbers EMDB-6878, 5Z3G and 5Z1G.

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