Subunit architecture of general transcription factor TFIIH

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Subunit architecture of general transcription factor TFIIH


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Structures of complete 10-subunit yeast TFIIH and of a nested set of subcomplexes, containing 5, 6, and 7 subunits, have been determined by electron microscopy (EM) and 3D reconstruction. Consistency among all the structures establishes the location of the “minimal core” subunits (Ssl1, Tfb1, Tfb2, Tfb4, and Tfb5), and additional densities can be specifically attributed to Rad3, Ssl2, and the TFIIK trimer. These results can be further interpreted by placement of previous X-ray structures into the additional densities to give a preliminary picture of the RNA polymerase II preinitiation complex. In this picture, the key catalytic components of TFIIH, the Ssl2 ATPase/helicase and the Kin28 protein kinase are in proximity to their targets, downstream promoter DNA and the RNA polymerase C-terminal domain.


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struction (RCT) method (37). The resolution of the reconstructions was estimated to be 25–35 Å by the Fourier shell correlation method (38) (Fig. S2).

For each preparation of holoTFIIH and subcomplexes, reference-free alignment, classification, and averaging produced predominant class averages as well as additional minor class averages that differed substantially from one another, likely due to compositional and conformational variability, deviations from preferred orientation, and contamination (Fig. S1). When the predominant class averages and corresponding 3D reconstructions of all four complexes were compared, a striking consistency and clear ordering was observed (Fig. 2). Each complex exhibited the features of that preceding it in the series, with extra density in a well-defined region, attributable to the addition of a subunit or subunits to the structure. The locations of Rad3, Ssd2, and TFIIK and their molecular outlines were revealed in this manner.

The unfixed core TFIIH preparation produced a predominant class average that exhibited features of the minimal core preparation with extra density attributable to Rad3 (Fig. 2B). The unfixed core + Ssd2 preparation produced four predominant class averages that exhibited features of the core preparation, three with extra density attributable to the Ssd2 subunit in different conformations, suggesting flexibility (Fig. 2C). One of the class averages was identical to the class average obtained from the core preparation, due to the loss of Ssd2, suggesting its tendency to dissociate in the absence of cross-linker (Fig. S1C). Similarly, two predominant class averages were identified for the fixed core + Ssd2 complexes that varied only in the Ssd2 region, suggesting two distinct conformations of Ssd2 (Fig. 3B and C “core + Ssd2 and core + Ssd2+”; class averages in Fig. S1D). Just as gradient fixation prevented dissociation of Ssd2 from the core, mild fixation in the holoTFIIH preparation also stabilized the association of TFIIK sufficiently to produce class averages that exhibited the features of the core + Ssd2 preparation with extra density attributable to the TFIIK kinase trimer, enabling 3D reconstruction of the complete 10-subunit TFIIH complex (Fig. 2D). Despite fixation, the holoTFIIH preparation also produced some class averages that were identical to the class averages obtained from the core + Ssd2 preparation, due to the loss of TFIIK from many of the particles (Fig. S1E). When the datasets were split into larger numbers of classes, small differences among class averages were observed due to flexibility of the connections between Rad3 and the minimal core, between Ssd2 and the core (Fig. 2C and Figs. S1 and S3), and between Rad3 and TFIIK (Fig. S3).

Calmodulin-Au Labeling and Flexibility of Subunits. To confirm the location of TFIIK, holoTFIIH bearing a calmodulin-binding peptide at the C terminus of Tfb3 was labeled with a calmodulin-gold cluster conjugate. The preparation was sedimented through a glycerol gradient containing cross-linker, and class averages of EM images obtained as described above displayed a dense spot between the regions attributed to Rad3 and TFIIK (Fig. 2E). Specificity was established by absence of the dense spot in class averages obtained from control preparations that included EDTA to chelate the calcium and prevent binding of the calmodulin-gold clusters to the calmodulin-binding peptide at the C terminus of Tfb3. These results confirm the presence of Tfb3 and also the

![Image](https://www.pnas.org/cgi/doi/10.1073/pnas.1105266109)

**Fig. 1.** SDS/PAGE of purified TFIIH complexes. Gels were stained with Coomassie blue and bands identified as TFIIH subunits (subunit names and predicted molecular weights indicated; see Table S3 for mass spectrometry analysis).

**Fig. 2.** TFIIH complexes. Class averages (Upper) and RCT reconstructions (Lower) of TFIIH complexes. (A) Minimal core, lacking Rad3. (B) Core, with additional density revealing the location of Rad3. (C) Unfixed core + Ssd2 data with additional density revealing the location of Ssd2 in several conformations indicating the flexibility of the Ssd2 subunit. (D) Fixed holoTFIIH with additional density revealing the location of the TFIIK kinase trimer. (E) Fixed holoTFIIH with Tfb3-calmodulin-Au (holoTFIIH+gold), indicating location of the TFIIK kinase trimer. In gold is shown the inverse volume corresponding to the negative density attributed to the gold particle attached to calmodulin bound to the CBP at the C terminus of Tfb3. Density that appears to be a connection between the trimer and the top of Rad3 is presumably due to Tfb3. RCT reconstructions were low-pass filtered to 35 Å and thresholded to match predicted volumes based on molecular weights (Table S4). The number of particles used to produce each class average and 3D reconstruction is indicated. The class average for fixed holoTFIIH has a less distinct appearance possibly due to inferior staining caused by residual glycerol or variability in cross-linking, because this dataset was obtained from a preparation isolated on a glycerol gradient containing glutaraldehyde. The class average for fixed holoTFIIH+gold has an improved appearance due to extensive washing of the grid to remove glycerol prior to staining. Scale bar, 10 nm (applies to all class averages and reconstructions in Figs. 2 and 3). (F) Fitting of RCT reconstructions. Panel 1 shows the minimal core volume fitted into the core volume, revealing the extra density for Rad3. Panel 2 shows the core volume fitted into one of the fixed core + Ssd2 volumes, revealing the extra density for Ssd2. Panel 3 shows the core volume fitted into the holoTFIIH volume. Panel 4 shows one of the fixed core + Ssd2 volumes fitted into the holoTFIIH volume, showing the extra density for the TFIIK kinase trimer. Fixed core + Ssd2 volumes are shown in F rather than unfixed volumes as in C for consistency with the fixed holoTFIIH in F.
assignment of the adjacent density to TFIIK. There appears to be a connection between TFIIK and the top of Rad3 that is presumably due to Tfb3 (Fig. 2 D and E). Besides placing the C terminus of the Tfb3 subunit of TFIIK at the interface with Rad3, the class averages revealed alternative conformations, in which TFIIK is rotated as much as 90° around its connection with Rad3, bringing it into proximity with Ssl2 (Fig. S3). Variability in the conformation of Ssl2 is apparent as well (Figs. 2 and 3 and Fig. S3). This is consistent with a proposed mechanism in which the C terminus of XBP rotates to interact with dsDNA (11) (Fig. S3).

Comparison and Docking with Previously Determined Structures. In addition to the structural analysis of holoTFIIH and subcomplexes, and calmodulin-Au labeling, a third line of support for the locations of subunits comes from comparison and docking of previous structures determined by EM and X-ray crystallography to the volumes from EM and 3D reconstruction shown here. The core TFIIH structure determined from 2D crystals (28) is similar in overall size and shape but appears to have captured a different conformation than the core volume reported here (Fig. 3A), and proposed locations of subunits by crude volume segmentation of the 2D crystal structure are also consistent with our subunit localizations. Observed differences in shape are likely to reflect either dramatic conformational changes or artifacts of the 2D crystallographic and/or single-particle specimen preparation methods.

The class averages and reconstructed volumes for the core + Ssl2 complex reported here are remarkably similar to those of the previously published human TFIIH structure (27) (Fig. 3C). This resemblance between yeast and human structures demonstrates evolutionary conservation and strongly suggests the absence of the kinase module in the human structure. Although it was reported to complete TFIIH, the human reconstruction has a volume that is only sufficient to accommodate the yeast core + Ssl2 without the kinase trimmer (Fig. 3C). It should also be noted that the localization of the subunits within our structures is partly inconsistent with the human TFIIH mapping by antibody labeling (27). While the human Ssl2 homolog, XBP, was localized with antibodies in the same general region identified as Ssl2 in this study, the human Kin28 homolog, cd7 (component of the kinase trimmer) and human Ssl1 homolog, p44 were both localized with antibodies in the region that has been clearly identified as Rad3 in this study, and the Rad3 human homolog, XPD, was localized with antibodies in the region that has been clearly identified as part of the minimal core in this study. The localizations that we report here are likely to be accurate because of the large and obvious density differences between the complexes, while the antibody labeling was limited by the small size of the antibodies and correspondingly small and ambiguous density differences. It is also possible that the epitopes for antibody binding, located at the subunit termini, are distant from the bulk of the subunit mass, leading to the misassignment of density to particular subunits.

X-ray crystal structures for Rad3 homologs from Sulfolobus tokodaii (29), Sulfolobus acidocaldarius (30), and Thermoplasma acidiphilum (31) resemble closely the three-lobe density that we have mapped here by comparison of the EM structures of the core and minimal core complexes (Fig. 2, Fig. 3, and Fig. 4). The crystal structure of an Ssl2 homolog in Archaeoglobus fulgidus (11) fits well into the extra density that we have mapped here by comparison of the EM structures of the core + Ssl2 and core complexes (Fig. 2, Fig. 3, and Fig. 4). Crystal structures are also available for the Kin28 and the Ccl1 human homologues Cdk7 (32) and cyclinH (33), respectively, which can be accommodated in the extra density that we have mapped here by comparison of the EM structures of the holoTFIIH and core + Ssl2 complexes and that we have assigned to the TFIIK kinase trimmer, with space remaining for Tfb3 (Fig. 2, Fig. 3, and Fig. 4).

Fig. 3. Comparison of TFIIH reconstructions with previously determined EM and X-ray structures. (A) On the left (labeled “2D core”) is shown the reconstructed volume of core TFIIH from EM of 2D crystals in stain with speculative designations based on segmentation of volumes for: Rad3 colored green; Ssl1 colored light gray; and Tfb1-4 colored orange (28). On the right (labeled “RCT core”) is shown the reconstructed volume of core TFIIH from this work with four different viewing angles (see also Fig. 2). The X-ray crystal structure of XPD (PDB ID code 2V5F) is fitted with Chimera (65), correlation coefficient = 0.98, to the Rad3 density revealed by the comparison between the core and core-Rad3 (minimal core) reconstructions (Fig. 2). XPD only fits well in one orientation, with the C-terminal region of XPD contacting the predicted location of Ssl1 (Fig. 4B). (B) Fitting of 20 Å resolution models computed with Chimera (65) from X-ray crystal structures. XPD (2V5F), the homolog of Rad3, is shown in green fitted to each of the TFIIH reconstructions (from left to right): core, core + Ssl2, core + Ssl2*, holoTFIIH, and holoTFIIH+gold. The two fixed core + Ssl2 classes suggest the presence of two distinct conformations of Ssl2 (referred to as core + Ssl2 and core + Ssl2*). XBP, the homolog of Ssl2, containing N- and C-terminal domains (2FWR) is shown in blue fitted to the Ssl2 density in the core + Ssl2 reconstruction. The individual N-terminal domain (2F24) and C-terminal domain (2F21) are shown in blue fitted together into the Ssl2 density in the core + Ssl2 reconstruction (see also Fig. S3). Cyclin H (1KW1), the homolog of Ccl1, is shown in magenta; and CDK7 (1UA2), the homolog of Kin28, is shown in red. Both are shown fitted together into the kinase trimmer density in the holoTFIIH and holoTFIIH+gold reconstructions. (C) Comparison of reconstructed volumes of human TFIIH from EM (27) (gray) to yeast TFIIH from this work. Yeast TFIIH structures from left to right: core + Ssl2 (blue), core + Ssl2* (blue), holoTFIIH (violet), and holoTFIIH+gold (violet). The core + Ssl2 structures show similarity to the human reconstruction but provide additional density, particularly for Ssl2. The holoTFIIH structures reveal the kinase trimmer that is absent in the human TFIIH reconstruction.

In the case of Rad3, the X-ray crystal structure may be oriented in the 3D volume on the basis of additional information. The N-terminal region of the human homolog of Rad3, XPD, was shown to interact with the C-terminal region of the human homolog of Tfb3, MAT1, by four-hybrid analysis (18). As already
mentioned, labeling with a calmodulin-gold conjugate placed the C terminus of Tfb3 at the interface with Rad3 (Fig. 2E). Other studies have shown that the C-terminal region of XPD interacts with the N-terminal region of the human homolog of Ssl1, p44 (40). Disease-causing mutations in the N- and C-terminal regions of XPD have been shown to destabilize its interactions with MAT1 and p44 and interfere with activity in transcription and DNA repair (18, 39, 43). The Rad3 homolog crystal structure could be fitted in the reconstructed volume from EM with its N-terminal region near the C terminus of Tfb3 and its C-terminal region adjacent to the likely location of Ssl1, constraining it to a single orientation in the volume. When fitted in this orientation, both the dimensions and the overall trilobate shape of the crystal structure are remarkably complementary with the appearance of this density in the EM structure (Fig. 3A). This result illustrates the bridging of Tfb3 to Ssl1 by Rad3, and highlights the role of Rad3 in anchoring the kinase trimer to the core TFIIH (Fig. 3B and Fig. 4B).

**Discussion**

TFIIH is a multisubunit enzyme that plays multiple roles in transcription, DNA repair, and the cell cycle. Its subunit organization underlies these diverse roles. The main findings of this work concern the arrangement of the TFIIH subunits (Fig. 2, Fig. 3, and Fig. 4). From a nested set of subcomplex structures, we have discerned the densities due to individual subunits. This extends our understanding of the organization of TFIIH beyond previous approximate locations of subunits determined by antibody labeling with the use of affinity tags or by volume segmentation. The delineation of subunit densities, in turn, provides a basis for docking previously described X-ray structures of individual subunits. The result is a first approximation to the atomic structure of complete TFIIH.

Fine structural features cannot be determined from the 3D reconstructions alone, due to various limitations: Embedding in stain results in loss of detail and flattening, and the random conical tilt reconstruction method suffers from a missing cone of information. Nevertheless, the fitting of X-ray structures, as well as consistency with previous electron microscopy (27, 28), testify to the validity of the 3D reconstructions (Fig. 3).

Our studies of TFIIH form part of a larger effort to determine the structure of the entire pol II transcription initiation complex (PIC). Toward this end, we combined our 3D reconstruction with structures of pol II and other components previously obtained to give a preliminary picture of the PIC (Fig. 4). In previous work, the X-ray crystal structure of a pol II—TFIIIB complex and that of a TBP—TFIIB—TATA box DNA complex were combined by superposition of the TFIIH components (44). The TATA box DNA was extended with straight B-form DNA without steric clash with pol II. The trajectory of the DNA downstream of the TATA box was consistent with that previously reported for PICs (45, 46). We introduced our TFIIH reconstruction in the picture on the basis of known interactions of TFIIH subunits with other components of the PIC (Fig. 4A). Ssl2 has been shown by protein–DNA cross-linking to interact with DNA approximately 30 base pairs downstream from the transcription start site (9, 10, 13, 46). We found two orientations of TFIIH that placed Ssl2 at this location and brought the Kin28 protein kinase in proximity to its substrate, the C-terminal domain of the Rpb1 subunit of pol II. Of these two orientations, only one was consistent with the known interaction of Tfb1 (47, 48) and Ssl2 (49, 50) with TFIIIE, and the interactions of TFIIE and TFIIF with pol II and DNA in PICs from chemical cross-linking (46, 51, 52, 64). In the resulting picture (Fig. 4C), the two ATPase/helicases, Ssl2 and Rad3, bracket the promoter and interact on opposite sides of pol II. Rad3 is in proximity to DNA, near the location of the transcription bubble in a transcribing complex, as has been suggested on the basis of biochemical work (9). Both Rad3 and TFIIK are placed near the proposed location of the Mediator head module (53) (see Fig. S4B) consistent with suggested interactions that may be important for regulation of PIC assembly and transcription initiation (54). In this manner, the findings from previous studies were combined with the present results to model the relationship between TFIIH and the other components of the transcription machinery. This model of the PIC (Fig. 4C) represents the association of TFIIH with the other components of the PIC prior to strand separation and formation of the “transcription bubble.” It is assumed that TFIIE, TFIIF, and, in some cases, TFIIA and TFIIIS will also be present. The precise arrangement of components doubtless changes over the course of strand separation and the initiation of transcription.

**Materials and Methods**

**Yeast Strains.** *Saccharomyces cerevisiae* protease-deficient strain CB010 was modified by the introduction of tandem affinity purification (TAP) tags at the C termini of Tfb3, Tfb4, and Ssl2 as described (55, 56). Briefly, homologous recombination cassettes were produced by amplifying the coding sequence from pBS1479 for a linker, a calmodulin-binding peptide, a tobacco etch virus (TEV) cleavage site, dual Protein A elements, a stop codon and a Trp selection marker, with primers (from IDT) containing sequences complementary to the C-terminal regions of Tfb3, Tfb4, and Ssl2, as well as sequences for amplifica-
tion from the plasmid (Table S2). An approximately 10-fold increase in the yield of stable holoTFIIH complex was achieved through the deletion of the YOR352W gene product, found to copurify with TFIIH complexes containing Ssl2 (Table S3 and SI Methods). YOR352W knockout strains were generated by a variation of the homologous recombination method using disrupcassettes produced by amplifying the coding sequence from pFLG (57) for Kan/Geneticin selection marker cassette flanked byloxP sites (for recycling) with primers (from IDT) containing sequences complementary to the upstream and downstream regions of YOR352W as well as sequences for amplification from the plasmid (Table S2). Transformations were performed by the LIoAc method and colonies were tested by PCR with internal and external primers to confirm the insertion locus and orientation, followed by Western blotting. The subunit composition of isolated TFIIH complexes was verified via mass spectrometry (see Table S3).

TFIIH Purification. TAP-tagged HoloTFIIH and TFIIH complexes were isolated fromSaccharomyces cerevisiae lysates with IgG affinity chromatography followed by additional steps to separate subcomplexes and improve homogeneity. HoloTFIIH and core – Ssl2 complexes were purified via TAP-tag on the Tfb3 subunit using IgG chromatography followed by ion exchange chromatography, size exclusion chromatography, and glycerol gradient centrifugation. Core and minimal core complexes were purified via TAP-tag on the Tfb4 subunit using IgG chromatography followed by hydrophobic interaction chromatography, ion exchange chromatography, and size exclusion chromatography (see SI Methods for details).

Calmodulin-Au Labeling. Wheat germ (Triticum aestivum) calmodulin (A.G. Scientific, Inc.) was resuspended in labeling buffer (25 mM Hepes pH 6.5, 0.1 mM calcium chloride) and reduced with 1 mM TCEP for 1 h at 37 °C. Reduced calmodulin (5 mg/ml) was reacted 4 h at room temperature (20 to 25 °C) with an equal mass of approximately 1.5 nm diameter monolayer-protected gold particles (approximately 1.5:1.0 molar ratio), prepared as described (58) except with 3-mercaptobenzoic acid instead of 4-mercapto-
benzoic acid. The gold-subunit complex was prepared by treatment with 10 mM glutathione for 1 h at 37 °C. The product was flash frozen in liquid nitrogen and stored at ~80 °C. Calmodulin-Au was added with a three fold molar excess to holoTFIIH complex and incubated for 1 h at 4 °C before gradient centrifugation or size exclusion chromatography.

Specimen Preparation and Electron Microscopy. TFIIH complexes (1–15 mg/ml) were stored until use at ~80 °C in 20 mM Hepes, pH 7.6, 250 mM potassium acetate, 1 mM DTT or TCEP, 1 mM EDTA or 0.5 mM CaCl₂, and 5% (v/v) glycerol, or 20–30% glycerol if from glycerol gradient. Where indicated, complexes were sedimented through 10–40% glycerol gradients containing 0.1% glutaraldehyde (34) (Fig. S1 and SI Methods). Concentrations were adjusted to 10–50 μg/ml in degassed, filtered solution containing 20 mM Hepes, pH 7.6, 250 mM potassium acetate, 1 mM DTT or TCEP, and 1 mM EDTA or 0.5 mM CaCl₂, and 2–4 μl were applied to continuous carbon-coated specimen grids (prepared by floating a fresh, thin carbon film onto 300 mesh Cu/ Rh grids (Ted Pellla or EMS), drying, and glow discharge in the presence of argon plasma immediately before use). Grids were washed with water, immersed in 1% uranyl acetate solution, lifted through a piece of thin carbon floating on the solution to apply a second carbon layer, blotted, and dried.

Untilted and ~55° tilt-pair images of the TFIIH complexes were collected manually on SO163 film (Kodak) or automatically on CCD (4 K × 4 K Gatan Ultrasmcan™ 4000) under low-dose conditions (each exposure approximately 100 s). After exposure, the film was developed on a Tecnai F20 microscope (FEI) operating at either 120 kV (Scirps) or 200 kV (Stanford). Negatives were digitized with a Leascan 45 scanner (Leaf Systems) and binned to a final pixel size of 3.0 Å on the object scale; CCD images were recorded with a pixel size of 2.29 Å.

Image Processing. The SPIPER and Web software packages (Version 17) (35) were employed. The defocus of each micrograph was calculated and power spectra were evaluated visually for evidence of drift or astigmatism. Particles from each class of micrograph and CCD frames with estimated defocus between approximately 200 nm and 600 nm were selected for further processing. Tilted micrographs were assessed on an optical diffractometer to ensure that the image was entirely underfocused. For the holoTFIIH and core– Ssl2 datasets, particles in tilt-pair micrographs were selected manually and interactively using TitlPicker (59). For the core dataset, particles in tilt-pair micrographs were picked automatically using a combination of ApoDogPick-er.py (59) and ApTiltAutoPicker.py (59) (details in SI Methods). For the mini-
complex dataset, tilt-pairs were picked automatically using the “Dog Picking” and “Auto Align Tilt Pairs” routines within the Appion processing pipeline (59). All individual particle images were windowed, ramped, and normalized. Images were then band-pass filtered, retaining information between 21 Å and 330 Å, and a soft-edged circular mask was applied to remove information at the corners of the images. Particle were reference-free aligned and classified using a modified routine originally developed for analysis of conformational flexibility of fatty acid synthase (60). Improvements were made to the procedure to allow for mirroring of particles. An initial round of this alignment and classification routine was used to remove particles that were consistently assigned to poorly aligned classes of particles. After removal of these particles, the minimal core, core – Ssl2, holoT-
FIH, and holoTFIIH-CaM-Au datasets were composed of 7,104, 7,583, 9,112, and 9,400 single-particle images, respectively. These images were subjected to a second round of alignment and classification (61), producing 20 class averages. The final in-plane rotational align-
ment parameters were used to calculate projection angles for the tilted particles to produce random conical tilt (RCT) reconstructions (62). These reconstructions were improved through six iterations of shift refinement to center the tilted particle images, and then band-pass filtered, retaining information between 35 Å and 300 Å.

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