Cytoplasmic TAF2–TAF8–TAF10 complex provides evidence for nuclear holo–TFIID assembly from preformed submodules

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General transcription factor TFIID is a cornerstone of RNA polymerase II transcription initiation in eukaryotic cells. How human TFIID—a megadalton-sized multiprotein complex composed of the TATA-binding protein (TBP) and 13 TBP-associated factors (TAFs)—assembles into a functional transcription factor is poorly understood. Here we describe a heterotrimeric TFIID subcomplex consisting of the TAF2, TAF8 and TAF10 proteins, which assembles in the cytoplasm. Using native mass spectrometry, we define the interactions between the TAFs and uncover a central role for TAF8 in nucleating the complex. X-ray crystallography reveals a non-canonical arrangement of the TAF8–TAF10 histone fold domains. TAF2 binds to multiple motifs within the TAF8 C-terminal region, and these interactions dictate TAF2 incorporation into a core–TFIID complex that exists in the nucleus. Our results provide evidence for a stepwise assembly pathway of nuclear holo–TFIID, regulated by nuclear import of preformed cytoplasmic submodules.
Eukaryotic class II gene transcription is controlled by a plethora of proteins, which are preassembled in large multiprotein complexes, including RNA polymerase II, Mediator and the general transcription factors (GTFs) 1. The sequential nucleation of GTFs and Mediator on core promoter DNA initiates regulated class II gene transcription 2. The GTF TFIIID plays a central role in this process by linking cellular signalling events with regulatory DNA elements and the transcription machinery 3. Although a basal transcription system supporting initiation of transcription from TATA-box-containing promoters can be reconstituted with TATA-binding protein (TBP), TFIIA, TFIIIB, TFIIH, TFIIH and TFIIH in vitro, TFIIID is additionally required to respond to activators and for efficient supporting initiation of transcription from TATA-less promoters 4, 5. In mammalian cells, most of the expressed protein-coding gene promoters are occupied by TFIIID and loss of TFIIID components leads to embryonic lethality 6–9. TFIIID subunits are implicated in crosstalk with epigenetic modifications on nucleosomes and regulatory DNA elements in promoter regions 10, 11. Structural analysis of TFIIID by cryo-electron microscopy revealed the overall architecture of TFIIID and provided important insights into subunit assembly and promoter recognition at low to medium resolution 12–16.

Canonical human TFIIID consists of TBP and 13 TBP-associated factors (TAFs) 17. Other non-canonical TFIIID and TAF-containing complexes have been identified recently with key roles during spermatogenesis and stem cell development 18–20. A central scaffold of canonical TFIIID comprises two copies each of TAF4, 5, 6, 9, and 12, which were shown to form a symmetric core 12, 21. This core–TFIID complex was first identified in Drosophila melanogaster nuclei 21. TAF3, 4, 6, 8, 9, 10, 11, 12 and 13 contain histone fold domains (HFDs), which stabilize discrete heterodimers (TAF3–10, TAF4–12, TAF6–9, TAF8–10 and TAF11–13) (refs 22–25). Among these HFD pairs, the TAF8–10 heterodimer plays a key role in the TFIID assembly pathway, is critical for the integrity of holo–TFIID and also fulfills essential functions in early embryonic development 6, 8, 26, 27. Binding of TAF8–10 to core–TFIID triggers a transition from symmetry to asymmetry, which was proposed to prime the recruitment of TAF1, 2, 3, 7, 11, 13 and TBP to complete holo–TFIID 12.

Evidence from genetic and biochemical studies showed that knock-out of the TAF10 gene leads to impairment of mature TFIIID assembly in F9 EC cells and to dissociation of TFIIID in hepatocytes 6, 26, 27. Biochemical data suggested that TAF8 and TAF10 interact strongly and specifically with each other via their HFDs 28. Identification of human TAF8 uncovered high sequence similarities with the Drosophila protein PRODOS and the mouse TBN protein 28, 29. Mouse embryos carrying a mutation in TBN develop normally to the blastocyst stage but fail to develop further due to the lack of inner cell mass cells 8. Interestingly, the same phenotype was also found in TAF10-knockout mice strongly suggesting that TAF8 and TAF10 are both involved in controlling embryonic development at similar stages 6. The importance of this cooperative activity of TAF8 and TAF10 is supported by nuclear import assays, which showed that the transport of TAF10 from the cytoplasm to the nucleus depends on the nuclear localization signal (NLS) found at the carboxyl-terminal (C-terminal) end of TAF8 (ref. 30).

Human TAF2 (originally called either CIF150 or TAFII150) has been previously described as an essential cofactor for TFIIID-dependent transcription from promoters with initiator (Inr)-containing promoter elements 31–33. Later it was suggested that a trimeric TBP–TFI–TAF2 complex is minimally required for efficient utilization of the Inr and downstream promoter elements 11. TFIIID complexes containing or lacking TAF2 have been described 31, 34 further suggesting that different types of TFIIID complexes may exist in human cell nuclei. Recently, mutations in the TAF2-coding gene were shown to be associated with various neurological disorders 35, 36. Human TAF2 is predicted to adopt an aminopeptidase-like fold with an additional C-terminal unstructured region. Localization studies using immunopurified TFIIID showed that TAF2 is an integral part of the central lobe of the holo-complex 13.

While general functions of individual TFIIID subunits and the holo-complex are increasingly better understood, very little is known to date about how the cell assembles this essential multiprotein complex. The existence of physiological core–TFIID in the nucleus, containing a subset of TAFs, provides evidence that the holo-complex may be assembled in a regulated manner from stable, preformed partial TFIIID subassemblies. The dependence of some of the TAFs on each other for nuclear import and the critical role of the TAF8–10 pair in functional remodelling of core–TFIID imply that discrete submodules preassemble also in the cytoplasm of cells. However, direct evidence for the presence of subassemblies in the cytoplasm is lacking to date.

By immunoprecipitating TAF-containing complexes from different cellular compartments, we identify a novel endogenous TFIIID subcomplex formed by TAF2, 8 and 10 in the cytoplasm of human cells. We dissect cytoplasmic TFIIA2–8–10 biochemically and structurally. We elucidate the interactions that stabilize the complex and reveal a central role of TAF8 in its nuclear translocation. By X-ray crystallography, we demonstrate a non-canonical histone-fold domain pair arrangement between TAF8 and TAF10. Furthermore, we demonstrate that the TAF2–TAF8 interaction is not only crucial for formation of the cytoplasmic TFIIA2–8–10 complex, but also dictates incorporation of TAF2 into a physiological core–TFIID complex that exists in the nucleus.

Results

An endogenous cytoplasmic TAF2–8–10 complex. With the objective to better understand human TFIIID assembly and in particular the incorporation of TAFs into holo–TFIID, we carried out immunoprecipitations from HeLa cell cytoplasmic and nuclear extracts. To test the role of TAF2 in the assembly process, we raised a polyclonal antibody using highly purified recombinant human TAF2 protein for the immunization procedure. We ascertained specificity of the purified antibody against recombinant TAF2 and endogenous TFIIID by western blotting (Fig. 1a and Supplementary Fig. 1a). Using this antibody, we carried out co-immunoprecipitation experiments of endogenous TAF2 from the cytoplasm, where TAF2 is synthesized de novo, and from nuclear extracts, where TAF2 likely functions in the context of TFIIID. To identify proteins that co-precipitated with TAF2 we subjected the immunoprecipitated samples to proteomics analysis by using the multidimensional protein identification technology (MudPIT). MudPIT analysis of proteins co-precipitated with TAF2 from the nuclear fraction revealed the full set of TFIIID components (Fig. 1b and Supplementary Table 1). We observed differences in abundance of the individual TAFs, which may argue for the presence of distinct TAF2-associated TAF or TFIIID-like complexes in the nucleus. Strikingly, MudPIT analysis of TAF2-associated proteins from the cytoplasmic fraction identified only TAF8 and TAF10, whereas none of the other TAFs could be detected (Fig. 1b,c). We confirmed the presence of TAF2, TAF8 and TAF10 in the cytoplasm of HeLa cells by immunofluorescence experiments (Fig. 1d,e and Supplementary Fig. 1b).
These data suggest that a unique endogenous TAF2–TAF8–TAF10-containing TFIID building block exists in the cytoplasm.

TAF8 nucleates the TAF2–8–10 complex. To further analyse this TAF2–TAF8–TAF10 complex, we used highly purified recombinant human TAF2, TAF8 and TAF10 to reconstitute TAF2–8–10 in vitro. We produced recombinant TAF2 and the TAF8–10 pair separately in insect cells and tested complex formation by size-exclusion chromatography (SEC) experiments. SEC of a stoichiometric mixture of TAF2 and TAF8–10 showed a clear peak shift in retention volume towards earlier fractions as compared with the individual components (Fig. 2a). Analysis of the chromatographic fractions by SDS-polyacrylamide gel electrophoresis (SDS–PAGE) shows that all three polypeptides co-elute in the same fractions (Fig. 2a). We observed unusually high molecular weight estimates for the components TAF2 and TAF8–10, and also for the complete TAF2–8–10 complex, which exceed the calculated molecular weights of the proteins. These high estimates can be due to either oligomerization or elongated shapes of the specimens analysed. We therefore determined the oligomeric states of purified TAF2, TAF8–10 and the TAF2–8–10 complex by analytical ultracentrifugation sedimentation velocity and native mass spectrometry (MS) experiments. Sedimentation coefficients of 4.3 S, 2.3 S and 4.9 S were obtained for TAF2, TAF8–10 and TAF2–8–10, respectively (Fig. 2b). Continuous size-distribution analyses returned best-fit molecular weights of 140, 52 and 200 kDa. These values are in good agreement with monomeric TAF2, heterodimeric TAF8–10 and heterotrimeric TAF2–8–10 complexes, with subunit stoichiometries of 1:1 and 1:1:1 in case of the complexes.

Analysis of TAF2–8–10 by native MS revealed a predominant complex with an average molecular mass of 195,797 Da corresponding to a TAF2–8–10 heterotrimer containing one copy of each protein (Supplementary Fig. 2a and Supplementary Table 2). We subjected the TAF2–8–10 complex to collision-induced dissociation (CID) experiments in the mass spectrometer to probe for subunit interactions37. The resulting spectra reveal dissociation of the trimeric complex into TAF2–8 and TAF10 submodules (Fig. 2c and Supplementary Table 2). Notably, TAF2 and TAF10 do not interact under the conditions studied, since we did not observe a TAF2–10 species (Fig. 2c). We conclude from these data that TAF2, 8 and 10 assemble as a heterotrimeric complex with a 1:1:1 stoichiometry and that the complex is nucleated by TAF8 and stabilized by distinct TAF2–8 and TAF8–10 interactions.

TAF8 and TAF10 adopt a non-canonical histone fold dimer. We next dissected the interactions identified by CID. First, we determined the X-ray crystal structure of the TAF8–10 complex. Previous GST pull-down experiments suggested that the interaction between TAF8 and TAF10 is mediated by their HFDs, which are present in the amino-terminal (N-terminal) half of

Figure 1 | A TAF2–8–10 complex exists in the cytoplasm. (a) Purified, polyclonal anti-TAF2 antibodies specifically recognize recombinant and endogenous TAF2. Recombinant (rec.) purified TAF2 (10 ng, lane 1) and immunopurified TFIID (300 and 150 ng; lanes 2, 3) were loaded on an 8% SDS–PAGE, blotted and analysed by western blot assay. Protein size markers are indicated. (b) Abundances of individual proteins co-immunoprecipitated from nuclear or cytoplasmic HeLa cell extracts (grey or black bars, respectively) using purified polyclonal anti-TAF2 antibodies were compared in units of normalized spectral abundance factors (NSAFs). Each column is the average of two independent experiments and error bars represent range of the data. (c) Domain organization of TAF2, TAF8 and TAF10 in a schematic view. Grey rectangles indicate predicted, unstructured regions. The NLS of TAF8 is shown as a black bar. Numbers indicate first and last amino acids in each protein. (d) Immunofluorescence microscopy of HeLa cells. Nuclei are visualized by 4,6-diamidino-2-phenylindole (DAPI) staining (blue), TAF2 is displayed in green and TAF8 in red. The bottom panel shows images of control cells, which were treated with secondary antibodies only. Scale bar, 10 μm. (e) Immunofluorescence microscopy of HeLa cells as in d, but displaying TAF2 (green) and TAF10 (red).
TAF8 and the C-terminal half of TAF10 (refs 28,30). We co-expressed and purified full-length TAF8–10 complex in insect cells from a polypeptide construct38, subjected the complex to limited proteolysis and defined the core complex to TAF8 residues 28–120 and TAF10 residues 113–212 (hereafter referred to as TAF8ΔC and TAF10ΔN, respectively; Supplementary Fig. 3a).

We prepared this TAF8ΔC–TAF10ΔN core complex, but only obtained crystals diffracting X-rays to 5–6 Å resolution. We therefore tested various N- and C-terminal deletion constructs of the two proteins in crystallization experiments. A complex of TAF8–10 comprising TAF8 residues 25–120 and TAF10 residues 112–212 yielded crystals, which diffracted incident X-rays to 1.9 Å resolution (Supplementary Fig. 3b). We determined the structure of this complex by the Sulfur-SAD method and refined the model to a crystallographic R value of 20.5% and a free R value of 23.7% with excellent stereochemistry (Table 1). The final model includes TAF8 residues 28–120 and TAF10 residues 98–218 (hereafter referred to as TAF8 and TAF10, respectively; Supplementary Fig. 3a). SDS–PAGE analyses of the eluted samples are shown (below). Molecular masses of protein standards are indicated on the left of gel sections. Protein denominations are shown on the right. First lane shows the SEC input (IN). (b) Absorbance c(s) profiles from sedimentation velocity analytical ultracentrifugation experiments are plotted for TAF2 (green), TAF8–10 (blue) and TAF2-8-10 (purple). (c) Mass spectrum of TAF8–10 complex electrosprayed from an aqueous ammonium acetate solution under high collision energy for subunit dissociation. The MS spectrum reveals peaks with corresponding masses for a TAF2–10 dimer (blue dots), TAF2 subunit (green dots) and a predominant TAF2–8–10 complex (purple dots) centred at 4,000, 6,000 and 7,500 m/z, respectively. At 12,000 m/z is a TAF2–8 dimer (yellow dots) resulting from the dissociation of the TAF10 subunit (light blue dots) from the intact TAF2–8–10 complex. Proteins and protein complexes are shown schematically as coloured circles.

Figure 2 | Recombinant TAF2–8–10 complex. (a) TAF2, the TAF8–10 pair and a mixture of TAF2–8–10 were analysed by SEC. Elution profiles of TAF2 (green), TAF8–10 (blue) and TAF2–8–10 (purple) are plotted in relative absorption units at 280 nm versus elution volume (top). Fractions are numbered (top of graph). SDS–PAGE analyses of the eluted samples are shown (below). Molecular masses of protein standards are indicated on the left of gel sections. Protein denominations are shown on the right. First lane shows the SEC input (IN). (b) Absorbance c(s) profiles from sedimentation velocity analytical ultracentrifugation experiments are plotted for TAF2 (green), TAF8–10 (blue) and TAF2–8–10 (purple). (c) Mass spectrum of TAF8–10 complex electrosprayed from an aqueous ammonium acetate solution under high collision energy for subunit dissociation. The MS spectrum reveals peaks with corresponding masses for a TAF2–10 dimer (blue dots), TAF2 subunit (green dots) and a predominant TAF2–8–10 complex (purple dots) centred at 4,000, 6,000 and 7,500 m/z, respectively. At 12,000 m/z is a TAF2–8 dimer (yellow dots) resulting from the dissociation of the TAF10 subunit (light blue dots) from the intact TAF2–8–10 complex. Proteins and protein complexes are shown schematically as coloured circles.

Interestingly, the proteins TAF8 and TAF10 have similar L1 loop geometries, which are not found in other structures of related HFD-containing TAFs (Fig. 3d,e)22,25. In both proteins, a phenylalanine of loop L1 (F50 in TAF8 and F144 in TAF10) is embedded in a composite, hydrophobic cavity mainly formed by residues from helices α2/α3 of one protomer and helices α2/α3 of the other (Fig. 3d,e). The amino acids forming this hydrophobic cavity are remarkably similar in TAF8 and TAF10, suggesting an evolutionary interrelation between the two proteins (Supplementary Fig. 3d). To test the functionality of the TAF10 HFD and chimeric mutants thereof, we performed complementation assays in TAF10 null mouse F9 cells30. Interestingly, the human TAF10 HFD (residues 116–218) is fully functional in the complementation assay, whereas chimeric constructs, in which either the N-terminal region of TAF10 (residues 116–150) or the C-terminal region (residues 151–218) was substituted by sequences of the yeast TAF10 homologue, were not functional (Supplementary Fig. 3f, g).
TAF8–10
Native
TAF8–10
S-SAD
Importin α1/TAF8-NLS
Native
| Data collection | TAF8–10 | TAF8–10 | Importin α1/TAF8-NLS |
|-----------------|---------|---------|---------------------|
| Space group     | P3:21   | P3:21   | P2(2,2,2)           |
| Cell dimensions | 90, 90, 120 | 90, 90, 120 | 90, 90, 90 |
| Resolution (Å)  | 44.44–1.91 | 49.54–1.75 | 49.54–1.75 |
| Rmerge/Rf     | 16,793 (1.630) | 12,943 | 55.43 (5,436) |
| No. of atoms | 1,474 | 4,817 | 3,877 |
| Protein        | 1,404 | 3,366 | 483 |
| Ligand/ion     | 7 | 59.9 | 44.8 |
| Water          | 63 | 48 | 463 |
| B-factors      | 3,554 | 3,366 | 48 |
| Bond lengths (Å) | 0.003 | 0.008 | 1.158 |
| Bond angles (°) | 0.681 | 1.158 | 0.681 |

HFDs of TAF8 and TAF10 are dispensable for TAF2 binding.
We next analysed the physical interactions between TAF2 and the TAF8–10 heterodimer. We first tested the effects of deleting the intrinsically unstructured regions of TAF8 and TAF10 on TAF2 binding in pull-down assays with purified proteins. As a control, full-length TAF8–10 was co-precipitated with TAF2. We confirmed the interaction between TAF2 and the C-terminal tail of TAF8 by SEC. We utilized full-length TAF2 and a fusion protein of MBP with residues 105–310 of TAF8 and evidenced complex formation (Supplementary Fig. 4). These results are consistent with the CID data in native MS, which showed that only TAF8, and not TAF10, is directly interacting with TAF2.

TAF2 recognizes short motives in the TAF8 C-terminal region.
We characterized the TAF2–TAF8 interaction further by means of a peptide array. We monitored the binding of His-tagged TAF2 to peptide arrays covering residues 105–310 of TAF8 (Fig. 4a). Densitometric analysis of the arrays indicated that TAF2-binding clusters around four distinct regions; a short N-terminal region I covering TAF8 residues 105–125, a less well-defined region II including residues 147–202 and regions III and IV spanning residues 207–238 and 282–310, respectively (Fig. 4a).

We next analysed the individual contributions of these four TAF8 regions to TAF2 binding by surface plasmon resonance (SPR) experiments. We generated N- and C-terminal deletion constructs of TAF8 and fused them to MBP (Fig. 4b). We monitored the association and dissociation phases of the MBP–TAF8 truncations on TAF2-charged sensor chips and compared binding kinetics at identical analyte concentrations. An MBP–TAF8-fusion construct spanning the entire C-terminal region (TAF8 residues 105–310) showed a maximal association level of about 85 response units (RU) with fast on and off rates (Fig. 4b). A shorter MBP-fusion protein lacking region I (TAF8 residues 141–310) showed similar kinetics but a reduced maximal association level of about 85 response units (RU) with fast on and off rates (Fig. 4b).

On the basis of our peptide array and SPR results, we introduced TAF8 point mutants into the TAF8–10 polyprotein expression construct by substituting three triple amino-acid clusters spanning residues 185–187 (DVE), 222–224 (PYL) and 293–295 (PYL) with alanines. We produced and purified

Primary sequence comparison with two other TAF10-interacting proteins, TAF3 and human SPT7L, shows that similar residues can be also found in their HFDs, arguing for a conserved binding mode of these proteins known to interact with TAF10 (Supplementary Fig. 3d). Our structure underscores that HFDs in TAFs can adopt a variety of conformations, which may differ significantly from the canonical histone pairs found in the nucleosome.
Figure 3 | TAF8–TAF10 interactions. (a) Crystal structure of human TAF8–10 complex is depicted in a cartoon representation. Two orientations related by a vertical rotation of 90° are shown. TAF8 is coloured in blue and TAF10 in green. The disordered L2 loop of TAF10 is represented by a dotted line. Secondary structure elements and loops are labelled. The TAF8–TAF10 complex adopts a non-canonical HFD pair. (b–c) Close-up views of the interactions between TAF8 and TAF10. Key interacting residues are highlighted. All structure drawings were generated with PyMOL (http://www.pymol.org/). (f) Pull-down experiments of TAF2 fused to MBP analysing the interactions with TAF8–10, TAF8–TAF10ΔN and TAF8ΔC–TAF10 HFD pairs (see main text for details). Unfused MBP is included as a control. Input samples (top) and samples precipitated on amylose resin (bottom) were resolved on 4–12% gradient gels. Protein identities are shown on the right.

wild-type TAF8–10 and the mutated TAF8–10 complex and analysed TAF2 binding via SEC (Fig. 4c). In contrast to wild-type TAF8–10, formation of a trimeric TAF2–8–10 complex was not observed with the three triple amino-acid cluster TAF8–10 mutant, corroborating the results that we obtained with our peptide array and SPR experiments (Fig. 4c).

TAF8 promotes TAF2 incorporation in TFIID. We showed recently that a TFIID subcomplex comprising TAF4, 5, 6, 8, 9, 10 and 12 (hereafter referred as 7TAF) can be formed in vitro by binding TAF8–10 to a physiological nuclear core–TFIID complex, which constitutes an important intermediate in holo–TFIID assembly,12,21. We next asked whether the association of TAF2 to this 7TAF complex depends on the C-terminal region of TAF8, which we identified as responsible for TAF2 binding in the TAF2–8–10 complex. We produced and purified recombinant 7TAF and a 7TAFΔ complex, in which TAF8 is substituted by TAF8ΔC (Fig. 5a). We monitored binding of a mCherry-tagged TAF2 fusion protein to 7TAF and 7TAFΔ complexes using SEC. We introduced the mCherry tag on TAF2 to unambiguously separate the protein from TAF4 by SDS–PAGE. In all SEC experiments, we used stoichiometric amounts of TAF2 in relation to TAF8–10 or the truncated TAF8ΔC–TAF10 complex. Interestingly, TAF2 could be fully incorporated into the 7TAF complex, whereas TAF2 did not interact noticeably with the 7TAFΔ complex, in which the C-terminal TAF2-interaction region of TAF8 had been deleted (Fig. 5b).

Next we mapped the position of TAF2 on 7TAF. To this end, we determined a three-dimensional model of negatively stained 7TAF complexes bound to TAF2 (hereafter referred as 8TAF complex) by single-particle electron microscopy and compared the resulting structure to the reconstruction of the 7TAF complex we had determined previously12. We observed major density differences clearly positioned on only one side of the particle, indicating TAF2 location (Fig. 5c and Supplementary Fig. 5a). Interestingly, our 8TAF complex reconstruction resembles a precursor to the characteristic clamp shape of holo–TFIID, in contrast to the less elongated shape of 7TAF and core–TFIID (Supplementary Fig. 5b).

Next we sought to characterize possible alterations in the protein–protein interaction networks along the assembly pathway to holo–TFIID. In particular, we looked at the transition from 7TAF to 8TAF complexes on TAF2 binding by crosslinking and MS (CLMS) experiments. We crosslinked 7TAF and 8TAF complexes with the bifunctional reagent bis(sulfosuccinimidyl)-suberate, BS3 that targets mostly lysines41 (Supplementary Fig. 6a). Crosslinked complexes were separated from non-crosslinked species by SDS–PAGE, in-gel digested and crosslinked peptides were assigned to ion masses observed by
MS. We identified 37 protein–protein crosslinks for the 7TAF complex and 37 protein–protein crosslinks for the 8TAF complex with an overlap of 21 crosslinked peptides between the two complexes (Fig. 5d, Supplementary Fig. 6b–e and Supplementary Table 3). Our data suggest that TAF9 plays a central role in 7TAF complex architecture by interconnecting TAF4, 5, 6, 8 and 12 (Fig. 5d). In our CLMS data, prominent crosslinks between TAF8 and TAF10 were not present, consistent with the paucity of lysines within crosslinking distance, and the partly buried location of the TAF8–10 dimer within the 7TAF complex.

In the 8TAF complex, we observed crosslinks of the C-terminal region of TAF8 with residues on TAF2, which are predicted to map to the surface (Fig. 5d). In addition to its proximity to TAF8, TAF2 is also positioned closely to TAF5, 6 and 9.

Figure 4 | TAF8–TAF2 interactions. (a) His-tagged TAF2 binding to overlapping peptides of the TAF8 C-terminal region (residues 105–310) spotted onto nitrocellulose membranes (spots A2-G4, left) was analysed by utilizing a peptide array. Bound TAF2 was visualized by luminol reaction and signal intensities were plotted for each spot after background subtraction (right). Spots A1, G5 and G6 served as positive controls. TAF2 protein was omitted for the control membrane. The four major binding regions (I–IV) are indicated above the histogram. (b) SPR experiments with immobilized full-length TAF2 as ligand and MBP (control) as well as MBP fusions of TAF8 fragments 105–310, 141–310, 200–310 and 105–260 as analytes. TAF8 deletion constructs are schematically shown as bar diagrams (left). TAF2-interacting regions on TAF8 as identified in a are highlighted. SPR sensorgrams at identical analyte concentrations of 500 nM are plotted as RU versus time (right). (c) SEC analyses assessing the influence of TAF8 point mutations on TAF2 binding. Elution profiles for the indicated proteins and protein complexes are plotted on the left and SDS–PAGE analyses of each run are shown on the right. Molecular masses of protein standards are denoted on the left of the gels and protein names on the right.
and promotes crosslinking between TAF4 and 5 (Fig. 5d). Our data indicate that TAF2 is indeed anchored to the 7TAF complex via the TAF2-interacting region on TAF8 and that binding of TAF2 induces significant conformational changes that result in novel TAF–TAF interactions not present in the 7TAF complex.

TAF2–8–10 binds Importin α1 via the TAF8 NLS. Biochemical and cell biology experiments demonstrated that the C-terminal NLS within TAF8 is necessary for shuttling TAF8 and TAF10 from the cytoplasm to the nucleus in an Importin α/β-dependent fashion30. We asked whether the identified endogenous TAF2–8–10 complex would be capable of recruiting Importin α1 in vitro to form a nuclear import complex. To this end, we mixed highly purified TAF2–8–10 with a twofold molar excess of an Importin α1 variant lacking the Importin β-binding domain (Importin α1ΔIBB). We observed efficient complex formation in SEC elution profiles for indicated proteins and protein complexes are shown (top). Corresponding SDS–PAGE gel sections of peak fractions of each run are shown (bottom). Protein size markers are shown on the left; protein identities on the right. (a) TFIID components studied are shown as bar diagrams. Predicted low-complexity regions of the proteins are coloured in grey. A black bar denotes the TAF8 NLS. C-terminally truncated TAF8 (TAF8ΔC), which was used to reconstitute the 7TAFΔ complex, is depicted on the right. Numbers denote first and last amino acids for each protein. (b) Impact of the TAF8 truncation on TAF2 binding to 7TAF complexes. SEC elution profiles for indicated proteins and protein complexes are shown (top). Corresponding SDS–PAGE gel sections of peak fractions of each peak are shown (bottom). Protein size markers are shown on the left; protein identities on the right.

Figure 5 | TAF8 promotes TAF2 incorporation in TFIID. (a) TFIID components studied are shown as bar diagrams. Predicted low-complexity regions of the proteins are coloured in grey. A black bar denotes the TAF8 NLS. C-terminally truncated TAF8 (TAF8ΔC), which was used to reconstitute the 7TAFΔ complex, is depicted on the right. Numbers denote first and last amino acids for each protein. (b) Impact of the TAF8 truncation on TAF2 binding to 7TAF complexes. SEC elution profiles for indicated proteins and protein complexes are shown (top). Corresponding SDS–PAGE gel sections of peak fractions of each run are shown (bottom). Protein size markers are shown on the left; protein identities on the right. (c) Three-dimensional single-particle EM reconstruction of negatively stained 8TAF complex (grey mesh) superimposed on 7TAF complex (yellow, from ref. 12) is shown in two views related by a 90° rotation as indicated (arrow). Difference density attributed to bound TAF2 is highlighted in blue. (d) A black bar denotes the TAF8 NLS. C-terminally truncated TAF8 (TAF8ΔC), which was used to reconstitute the 7TAFΔ complex, is depicted on the right. Numbers denote first and last amino acids for each protein. (Image 67x514 to 320x602)

and (Image 87x380 to 308x487)

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The cryo-electron microscopy density envelope of core–TFIID complex is shown (adapted from ref. 12). On binding of Importin α1–TAF8 complex crystal structure. Magnified view of interacting residues of the major binding site of Importin α1 (grey) with residues of the NLS of TAF8 (blue). Importin α1 is shown in ribbon representation and the TAF8-NLS as a stick model. TAF8 residues R303 and R304 are not involved in contacting Importin α1 and are omitted for clarity. (c) Immunofluorescence microscopy of HeLa cells depleted of TAF8 (TAF8 siRNA) by RNAi or control cells (Control siRNA). Nuclei are visualized by 4',6-diamidino-2-phenylindole (DAPI) staining (blue). TAF2 is displayed in green and TAF8 in red. Arrows point to a non-transfected cell. Scale bar, 10 μm. (d) Cartoon model of cytoplasmic TAF2–8–10 complex and nuclear holo–TFIID assembly. The NLS of TAF8 is filled in black. The TAF2-interaction domain within TAF8 is highlighted by shading. TAF2, blue; TAF8, red; Importin α1, grey; Nuclease, orange. The TAF2–8–10 complex resides in the cytoplasm, whereas the physiological symmetric core–TFIID complex is found in the nucleus. The cryo-electron microscopy density envelope of core–TFIID complex is shown (adapted from ref. 12). On binding of Importin α1 (grey) to the TAF8 NLS, TAF2–8–10 translocates into the nucleus through a nuclear pore (arrow). In the nucleus, Importin α1 is released and TAF2, 8 and 10 associate with core–TFIID, to form intermediates including the asymmetric 8TAF complex along the holo–TFIID assembly pathway.

Discussion

Elucidation of the structure and function of multiprotein complexes in gene regulation is an intense focus of current research efforts. Whereas three-dimensional models of fully assembled multiprotein complexes derived from X-ray crystallography or single-particle cryo-electron microscopy provide a wealth of information on the architecture of such complexes, little is known about how the cell controls and regulates the ordered assembly of multiprotein gene regulatory complexes such as TFIID.

Several earlier studies described the existence of a variety of TFIID complexes with distinct subunit composition in different cell types conveying a concept of modular TFIID complexes such as TFIID. We next asked if the nuclear localization of TAF2 is dependent on the presence of TAF8 in vivo. Therefore, we knocked down endogenous TAF8 in HeLa cells by RNA interference (RNAi) treatment for 48 h and compared the nuclear/cytoplasmic distribution of TAF2 in TAF8 knockout cells with control cells by immunofluorescence (Fig. 6c). Short interfering RNA (siRNA) treatment leads to depletion of TAF2 in the nucleus and to an enrichment of TAF2 in the cytoplasm, suggesting that the import of TAF2 is controlled by TAF8 (Fig. 6c).

Taken together, our data suggest the presence of a nuclear import particle in which the TAF2–8–10 complex is bound by the major binding site of Importin α1 via the NLS of TAF8, poised to shuttle into the nucleus (Fig. 6d). On release of Importin α1, the TAF2–8–10 module then combines with core–TFIID (Supplementary Fig. 2b,c). Association of these preformed TFIID submodules leads to conformational rearrangements in the resulting intermediate TAF complex which enables formation of the functional nuclear holo–TFIID (Fig. 6d).
characterized the interactions stabilizing this cytoplasmic complex in an integrated approach combining native MS, X-ray crystallography, SPR, peptide arrays and biochemical and biophysical methods. Our experiments indicate that TAF2 interacts with the C-terminal unstructured region of TAF8 in vitro, substantiating protein–protein interaction mapping experiments of *Saccharomyces cerevisiae* TFIIĐ.\(^{49}\)

Previously, it was shown that TAF8 shuttles from the cytoplasm to the nucleus in an Importin α/β-dependent pathway and piggybacks TAF10 into the nucleus.\(^{30}\) Owing to the lack of a NLS, TAF10 cannot translocate to the nucleus on its own and depends on the NLS of its interaction partner, TAF8 (ref. 30). Deletion mutants suggested that Importin binding resides in the extreme C-terminus of TAF8 (ref. 30). We show that a tetrameric complex consisting of Importin α1, TAF2, TAF8 and TAF10 can be assembled from purified components in vitro suggesting a co-import mechanism for the three proteins. Our TAF2 and TAF10 cellular localization experiments support this mechanism, indicating that knockdown of TAF8 by RNAi not only alters the cellular localization of TAF10, but likewise the localization of TAF2.

To define the precise modes of interaction between Importin α1, TAF8 and TAF10, we solved the X-ray crystal structures of Importin α1 bound to the NLS of TAF8 on one hand, and of the HFD pair formed by TAF8 and TAF10 on the other. The crystal structure of the TAF8–10 complex reveals atypical histone folds of the two proteins and shows a combination of symmetric and asymmetric structural elements. Both TAFs share characteristic conformations of their L1 loops, which give rise to pseudosymmetric structures at the extremities of their HFDs. Otherwise, the presence of additional αN and αC helices render the TAF8–10 complex asymmetric. The pseudosymmetric L1 loops are characteristic for the TAF8–10 complex, since similar arrangements cannot be found in the crystal structures of the *Drosophila* TAF6–TAF9 and the human TAF4–TAF12 complex.\(^{22,25}\) suggesting that the overall shape and precise geometry of the complex is important for integration into core–TFIID.

The crystal structure of Importin α1 with the NLS of TAF8 reveals that TAF8 has a classical short monopartite NLS, which is recognized by the major binding site of Importin α1. Interestingly, phosphorylation of conserved serine residues C-terminal to canonical NLSs of different nuclear proteins either enhance, or abolish, the binding affinity of different importins, thus regulating nuclear import.\(^{50–52}\) Similarly, phosphorylation may also fine-tune nucleocytoplasmic shuttling of the TAF2–8–10 complex. Two serine residues predicted to be phosphorylated are located C-terminal to the NLS of TAF8. Therefore, the affinity of TAF8 to Importin α1 and consequently the nuclear import could be modified by phosphorylation. It will be interesting to see if such a regulatory mechanism by post-translational modification exists for the import of the TAF2–8–10 complex in vivo. Our crystal structure of Importin α1 bound to the TAF8 NLS further suggests that the cytosolic TAF2–8–10 complex together with Importin α1 constitutes an import particle responsible for delivering this building block into the nucleus. Likewise, our experiments indicate that this TAF2–8–10 building block is responsible for the incorporation of TAF2 in nuclear TFIIĐ.

A stable TFIIĐ core complex comprising two copies each of TAF4, 5, 6, 9 and 12 was identified in *Drosophila* and human cell nuclei.\(^{12,21}\) Previously, we postulated that the binding of TAF8–10 causes a rearrangement of the symmetric TFIIĐ core complex to an asymmetric particle, which is then capable of accommodating the remaining TAFs and TBP, each in single copy.\(^{12}\) We propose that association of the TAF2–8–10 complex with the preassembled nuclear core–TFIID involves an intricate network of interactions between the TAF8 C-terminal tail and TAF2 on one hand, and the globular HFD pair of the TAF8–10 complex and core–TFIID on the other. Our current data suggest that TAF8–10 may function alike a chaperone to regulate nuclear import and integration of TAF2 into core–TFIID. Note, however, that in the cytoplasmic extracts, apart from TAF8 and TAF10, we did not detect any of the other TFIIĐ components stably associated with endogenous TAF2. Therefore, we hypothesize that TAF1, 7, 11, 13 and TBP incorporate into the TFIIĐ structure probably at a defined, later step, after TAF2–8–10 has been acetylated. Moreover, notwithstanding the fact that we did not find either TAF1 or TBP associated with TAF2 in the cytoplasm in our co-IP coupled MS analyses, it still can be envisioned that the TAF2–8–10 complex is capable of nucleating the formation of the holo–TFIID complex, including the TAF1–TBP module, and thus promote transcription from Inr-containing core promoters.

Interestingly, TAF2-containing and TAF2-lacking, as well as TAF10-containing and TAF10-lacking, TFIIĐ complexes have also been extracted from human cells.\(^{22,23,45}\) Thus, in good agreement with the modular TFIIĐ assembly concept, our observations suggest that the here characterized TAF2–8–10 building block would act like a scaffold to incorporate in all canonical TFIIĐ complexes but, even in the nucleus, may exist as an independent regulatory entity. Future experiments will be required to elucidate the function(s) of holo–TFIID complexes versus complexes lacking TAF2–8–10. Along the same lines, it will also be interesting to test whether a TAF2–8–10 complex alone or in combination with core–TFIID can modulate transcription efficiency of Inr-dependent genes. Promoter architecture may at this junction control transcription regulation by gauging the assembly rate of holo–TFIID from building blocks.\(^{48}\) From a pharmaceutical point of view, it is to date entirely unclear whether or not neurological disorders caused by mutations in the TAF2 gene develop due to altered regulation of transcriptional activity or due to other currently unknown mechanisms. Future experiments will be required to elucidate if these TAF2 mutations may actually affect TAF8 binding and TFIIĐ assembly.

Our results support the view that stable partial TFIIĐ complexes—potentially have important functions of their own—might exist in the cell. These complexes may represent functional cytoplasmic or nuclear modules, which assemble into holo–TFIID in a stepwise fashion. Also, our results point to an important role of cytoplasmic–nuclear transport in holo–TFIID formation. We anticipate that such processes will likewise play important roles in regulating the assembly and activities of many other multiprotein complexes that direct gene transcription.

**Methods**

**DNA constructs.** Cloning of TAF2, TAF8 and TAF10 expression constructs in MultiBac plasmids pPBac,\(^{28}\) pFl and pIDC\(^{5}\) is detailed in the Supplementary Methods. Expression plasmids for subcomplexes TAF5–6–9 (pPbac–3TAF) and TAF4–12 (pIDF–412) and Importin α1\(^{22,25}\) were described previously.\(^{2,25}\) Truncated Importin α1 (residues 71–497) was generated by amplifying the coding region of Importin α1\(^{22,25}\) by PCR. All constructs were verified by DNA sequencing.

**Sequence alignments.** Alignments were generated using the ClustalW2 server\(^{55}\) and plotted with ESPript\(^{56}\). Protein sequences for human TAF5 (UniProt accession number Q5VVW9) and human TFIJD (Q9H464) were retrieved from the UniProt server (www.uniprot.org).

**Protein production and purification.** MBP–TAF8–fusion proteins were produced in *E. coli* Rosetta (DE3) cells (Novagen) and purified by metal affinity chromatography using TALON resin (Clontech) followed by size-exclusion chromatography on a Superdex 200 16/60 column (GE Healthcare; detailed in Supplementary Methods). Importin α1 constructs (residues 60–529 or residues 71–497) were produced and purified as described,\(^{2,25}\) except that *E. coli* Rosetta (DE3) cells (Novagen) were used. Production and purification of core–TFIID and 7TAF complexes was performed as described.\(^{12}\)
Proteins TAF2, MBP–TAF2, mCherry–TAF2 and TAF8–10 complex were produced using the MultiBac system. Expressed protein was captured via TALON resin (Clontech) from the cell lysate in batch. Proteins were further purified by ion exchange chromatography using a 5-ml SP-Sepharose HiTrap column (GE Healthcare) followed by gel filtration using Superdex200 10/300 or Superose6 10/300 columns (GE Healthcare, Supplementary Methods). Proteins were flash frozen in liquid nitrogen and stored at −80 °C in aliquots.

Binding experiments. SEC experiments were carried out using AKTA purifier or AKTA Micro systems (GE Healthcare) using Superdex200 10/300, Superdex200 PC3.2, Superose6 PC3.2 or Superose6 PC3.2 Increase columns. Binding experiments shown in Fig. 2a were performed in running buffer comprising 25 mM HEPES-NaOH, pH 7.5, 150 mM NaCl, 1 mM DTT (dithiothreitol). Runs in Figs 4c and 5b and Supplementary Fig. 2b, c were performed in buffer comprising 25 mM HEPES-NaOH, pH 7.5, 500 mM NaCl, 1 mM DTT. Crystallization and structure determination. Screening for crystallization conditions was performed at the High Throughput Crystallization (HTX) laboratory (EMBL-EBI, Supplementary Methods). Crystals of TAF8–10 complex (TAF8 residues 25–120 and TAF10 residues 112–212) were refined manually by mixing equal volumes of protein solution containing 15–25 mg ml−1 TAF8–10 in 25 mM Tris-HCl, 150 mM NaCl at pH 7.5 and crystallization solution containing 1.4 M NaK PO₄ at pH 7.6. Crystals grew in space group P3121 with cell dimensions a = 184.1 Å, b = 184.1 Å and c = 144.4 Å. Crystals were cryoprotected by adding 20% (v/v) glycerol and flash frozen in liquid nitrogen. Diffraction data were collected at 100 K on beamline PROXIMA 1 using a Pilatus 6 M detector (SOLEIL synchrotron, Gif-sur-Yvette, France) and were integrated and scaled using X-ray Detector Software (XDS)57. The structure of TAF8–10 was solved by the Sulfur-Augmented Molecular Replacement (SARMoR) approach. The TAF8–10 structure was built and refined using programs Coot61 and Phenix62, respectively, including TLS parameter, occupancy and individual B-factor refinements.

Surface plasmon resonance. Biosensor experiments were performed at 25 °C on a BIACORE 3000 (Biacore AB, Uppsala). TAF2 ligand was immobilized onto double β-alanine anchors and assembled using the SPOT technology (AG Molekulare Bibliotheken, Charité—Universitätsmedizin Berlin, Germany). Overlapping 20-mer peptides of TAF2 were synthesized via Fmoc (9-fluorenylethoxycarbonyl) chemistry on a 4370 Peptide Synthesizer (Millipore) to create three of amino acids between neighbouring spots. Low-density hexa-Histidine peptides were used as controls. Peptide membranes were blocked in blocking buffer (50 mM Tris-HCl, pH 7.6, 500 mM NaCl, 20% (v/v) sucrose, 3% (w/v) bovine serum albumin) for 1 h at 4 °C, washed with TBS (50 mM Tris-HCl, pH 7.6, 500 mM NaCl) and incubated for 1.5 h with His-tagged TAF2 (10 μg/ml) in blocking buffer or with blocking buffer alone. Membranes were incubated with mouse anti-His monoclonal primary antibody (Sigma-Aldrich, catalogue number H1029, dilution: 1:13 000) and peroxidase-conjugated anti-mouse secondary anti-body (Sigma-Aldrich, catalogue number A9596, dilution: 1:10 000) in blocking buffer. Membranes were washed three times with TBS between each incubation step. Luminol solution (Pierce) was added and luminescence detected on a KODAK 4000MM photomter. Images were analysed using the Dot Blot Analyzer tool in ImageJ.

Analytical ultracentrifugation. Sedimentation velocity experiments were performed in a Beckman XL-I analytical ultracentrifuge (Beckman Coulter). The purified proteins and protein complexes TAF2–TAF2–TAF8–TAF2–TAF2–10 and TAF8–10 were loaded into sample windows with 12-mm optical path length and spun in an An-60Ti rotor (Beckman Coulter). Absorbance at 280 nm was measured for 16 h at 42 000 r.p.m. and 10 °C. The data were analysed in terms of continuous size-distribution (c(s)) with the Sedit program58, considering 200 particles with sedimentation coefficients, s, between 0.1 and 20 S. A partial specific volume of 0.73 and svedberg ratio of 1.4 (TAF2, TAF2–8–10) and 1.6 (TAF8–10) were used. A regularization procedure with confidence level of 0.68 was applied. Sample densities and viscosities were determined with Sednery61 to 1.023 g/ml and 1.40 mPas (TAF2, TAF2–8–10) and 1.021 g/ml and 1.31 mPas (TAF8–10).

Native MS. Purified TAF8–2–10 complex (30 μl) was buffer exchanged into 500 mM ammonium acetate buffer (pH 7.5) using Amicon spin concentrators (Millipore). All MS experiments were performed on a Quadrupole Time-of-flight (Q-ToF) II mass spectrometer (Waters, Manchester, UK) in the positive ion mode.62 For data acquisition, 2 μl of the sample was injected into the mass spectrometer with gold-coated capillary needles made in-house using a needle pusher (Harvard Apparatus, Holliston, MA, USA). MS spectra were acquired using a capillary voltage of 1.7 kV and cone and collision voltages of 100 V. Time-of-flight and analyser pressures were at 5.6 × 10⁻⁶ and 4.2 × 10⁻⁶ mbar, respectively. Data sets were acquired and processed with Masslynx V4.1 software (Waters, UK) with minimal smoothing and no background subtraction. The recorded mass spectra were calibrated externally using 100 μg/ml calcium iodide in water.

SUMMARY

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alkylated and trypsin digested following standard procedures. Crosslinked peptides were fractionated using SCX-StageTips following published protocols for linear peptides described using C18 StageTips41.

**Mass spectrometry.** Peptides were analysed on an LTQ Orbitrap Velos mass spectrometer coupled with an UltiMate 3000 Rapid Separation LC system (Thermo Fisher Scientific). The column was packed into a spray emitter (75-μm inner diameter, 8-μm opening, 250-mm length; New Objectives) with C18 material (RepelSil-Pur C18 A-03 μm Dr Maisch, Ammerbuch-Entringen, Germany) using an air pressure pump (Proxeon Biosystems). Mobile phase A consisted of water and 0.1% formic acid. Mobile phase B consisted of 80% acetonitrile and 0.1% formic acid. Peptides were loaded onto the column with 2% B at 500 nl min⁻¹ flow rate and then increased to 4% B at 15 min and finally increased to 100% B in 139 min; then increase from 40 to 95% B in 11 min. The eluted peptides were directly sprayed into the mass spectrometer. Peptides were analysed using a high/low strategy: both MS spectra and MS2 spectra were acquired in the Orbitrap. MS spectra were recorded at 100,000 resolution. The eight highest intensity peaks with a charge state of three or higher were selected in each cycle for ion trap fragmentation. Fragments were produced using CID with 35% normalized collision energy and detected by the Orbitrap at 7,500 resolution. Dynamic exclusion was set to 90 s and repeat count was 1.

**Data processing.** The mass spectrometric raw files were processed into peak lists using MaxQuant (version 1.3.0.5) (ref. 41) at default parameters except for ‘top MS/MS peaks per 10 Da being set to 100. Search was conducted against TAF complex sequences using Xi software (version 1.3.355). Search parameters were MS accuracy, 6 ppm; MS/MS accuracy, 20 ppm; enzyme, trypsin; crosslinker, BS3 (including BS3 modification); max. missed cleavages, 4; fixed modification, carboxymethylation on cysteine; variable modifications, oxidation on methionine; crosslinkable amino acids, N-terminus, lysine, serine, tyrosine and threonine; fragments, b and y ions with loss of H₂O, NH₃ and CH₃OH. The data have been validated by 5% FDR with manual validation. UniProt protein accession numbers of the protein sequences used to search the database were as follows: TAF2 (Q6P1X5-1; with a sequence variation R785G; European Nucleotide Archive AAC68502.1), TAF3 (P49848-1), TAF5 (P49848-1), TAF4 (Q7ZC8-1), TAF9 (Q18624-1), TAF12 (Q16514-1). Crosslinks observed in artificially introduced sequences (for example, TEV cleavage and crosslinkable amino acids, N terminus, lysine, serine, tyrosine and threonine; fragments, b and y ions with loss of H₂O, NH₃ and CH₃OH) were directly sprayed into the mass spectrometer. Peptides were analysed using a high/low strategy: both MS spectra and MS2 spectra were acquired in the Orbitrap. MS spectra were recorded at 100,000 resolution. The eight highest intensity peaks with a charge state of three or higher were selected in each cycle for ion trap fragmentation. Fragments were produced using CID with 35% normalized collision energy and detected by the Orbitrap at 7,500 resolution. Dynamic exclusion was set to 90 s and repeat count was 1.

**Protein extract preparations and immunoprecipitation and MudPIT analyses.** HEK293 cell nuclear extract (NE) preparations and immunoprecipitations were done as described48 with minor modifications. Supernatant containing the cytoplasm was precipitated by adding stepwise 0.3 g ml⁻¹ ammonium sulfate under agitation (4°C, 30 min). Precipitated proteins were collected by centrifugation (30,000 g; 4°C, 20 min), resuspended and dialysed overnight.

Immunoprecipitated proteins were eluted from the protein G columns with 0.1 M glycine (pH 2.5) and quickly neutralized with 2 M Tris-HCl (pH 8.8) buffer.

**Protein identification.** Proteins were identified by database searching using SEQUEST with ThermoProteome Discoverer 1.4 (Thermo Fisher Scientific)49. Tandem mass spectra were searched against a human protein sequence database (from the Homo sapiens 2013-04-03 Swissprot release). In all searches, cysteine residues were considered to be fully carboxymethylated (≈ 57 Da statically added) and methionine to be oxidized (≈ 16 Da dynamically added). Relative protein abundance for each protein in a given sample was estimated by normalized spectral abundance factor22.

Normalized spectral abundance factor values were calculated from the spectral counts of each identified protein. Larger protein spectral peptide/spectra and therefore, spectral counts were divided by protein length to provide a spectral abundance factor (SAF). SAF values were then normalized against the sum of all SAF values in the corresponding run allowing the comparison of protein levels across different runs. The MS proteomics data were deposited to the ProteomeXchange Consortium66 via the PRIDE partner repository with the data set identifier PXD001427.

**Immunofluorescence.** Indirect immunofluorescence tests were performed as described50 with the following modifications: cells were fixed with 4% paraformaldehyde for 15 min at room temperature (RT) and then permeablisized with 0.1% Triton-X100 for 20 min at RT, incubated with 1% FBS for 1 h at RT with either an immunolabeled 3' endriminal peptide containing 11 amino acids (Homo sapiens 2013-04-03) and TAF2 (mouse monoclonal antibody (mAb) 1FR-1B6 (ref. 6); diluted 1:1,000) or anti-TAF2 + anti-TAF10 (mouse monoclonal antibody (mAb) 1FR-1B6 (ref. 6); diluted 1:1,000) antibody mix followed by incubation (RT, 1 h) with secondary antibody mix including Alexa488-conjugated goat anti-rabbit mAb (Life Technologies, catalogue number A-11034, diluted 1:800) and Alexa568-conjugated goat anti-mouse mAb (Life Technologies, catalogue number A-11004, diluted 1:3,000; detects either anti- TAF8 or anti-TAF10). As negative control, cells were incubated with secondary antibodies only to quantify background signal. Cells were imaged using a Veckashield mounting medium with DAPI (Vector laboratories Inc.). Images were analysed on a Leica widefield microscope equipped with a CoolSnap HQ camera (×63 or ×100 magnification). Fluorescence intensities measurements in the cell cytosplasm were performed using Fiji software; intensity values were normalized to background signals.

**siRNA transfection.** siRNAs targeting TAF8 (ON-TARGETplus SMARTpool siRNA J-015912-20, J015912-19, J015912-18, J015912-17; Dharmaco; Thermo-Scientific) and non-targeting control (D-001810-10-20) were transfected into HEK293 cells using Lipofectamine 2000 transfection reagent (Invitrogen). Cells were fixed for immunofluorescence experiments 48 h after transfection.

**Electron microscopy.** Specimen preparation. 8TAF sample was stabilized by mild glutaraldehyde crosslinking (Grafix®). Two-hundred μl purified 8TAF complexes were deposited on a 4-ml centrifugation tube containing a 10 to 30% glycerol and a 0.15% glutaraldehyde gradient followed by centrifugation (34,000 r.p.m., 18 h, 4°C) with a SW6 rotor (Beckman Coulter). Fragments containing stabilized sample were deposited onto a buffer exchange column (Zeba spin desalting columns, Pierce) to remove excess glycerol. Specimen was deposited onto a thin layer of carbon coated on an electron microscopy grid and negatively stained for 45 s with 2% of uranyl acetate. Images were acquired using a Tecnai F20 (FEI) equipped with a field emission gun operating at 200 kV. Images were recorded under low-dose condition (total dose of 40–50 e⁻/Å²) on a 2,048 × 2,048 CCD camera (Ultrascan 1000, Gatan Inc., Pleasanton) at a magnification of 50,000 resulting in a pixel spacing on the specimen of 0.21 nm.

**Random conical tilt reconstructions.** The initial reference volumes were obtained by random conical tilt using XMIPP39 and IMAGIC67 software packages. Two consecutive images of the same area were taken at 45° and 0° tilt angles under low-dose electron microscopy conditions. A total of 1,346 tilt pairs were selected manually using XMIPP. Untilted images were aligned using iteratively refined two-dimensional class averages as references and multivariate statistical analysis and Hierarchical Ascendant Classification for clustering into 50 class averages with IMAGIC. Fifty volumes calculated from two-dimensional classes were aligned, clustered and averaged using XMIPP MLtomato to compensate for the missing cone, resulting in five random conical tilt (RCT) reconstructions.

**Structure refinement.** The best volume was used as reference for refinement cycles using a data set of 35,145 untitled molecular images windowed with the Boxer application of the EMAN2 software package50 and coarsened by two resulting in a pixel spacing of 4.2 Å. Image sorting was found necessary to select the most homogenous particles since part of the structure was flexible and prevented convergence. Sorting was performed by using first XMIPP then subsequently the RELION software package57. Final 3D reconstruction was performed in RELION with 2,361 sorted particle results in a structure with 37 Å resolution as estimated by the 0.5 Fourier Shell Correlation criteria. Images were prepared using Chimera software (http://www.cgl.ucsf.edu/chimera).

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Author contributions

S.T., C.V., L.T. and I.B. designed and interpreted the experiments. S.T., C.V. and M.H. produced and purified all proteins and performed characterizations. S.T. determined the crystal structures. E.S. prepared the antibodies and carried out all the anti-TAF2 IPs. V.C. and M.F. performed MudPIT experiments and M.F. and L.T. analysed them. I.-O.E. and C.V.R. performed and analysed native mass spectrometry experiments. S.C. and L.T. prepared and interpreted cell-based experiments. G.P., C.S. and P.S. prepared EM grids, collected and analysed data and calculated reconstructions. J.Z. and J.R. performed and analysed CLMS experiments. S.T., L.T. and I.B. wrote the manuscript with input from all authors.

Additional information

Accession codes: Atomic coordinates and structure factors are deposited in the Protein Data Bank (PDB 4WV4; 4WV6). Mass spectrometry data has been deposited in the ProteomeXchange database (PXD001454; PXD001427).

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