Homozygous TAF8 mutation in a patient with intellectual disability results in undetectable TAF8 protein, but preserved RNA polymerase II transcription

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Abstract

The human general transcription factor TFIID is composed of the TATA-binding protein (TBP) and 13 TBP-associated factors (TAFs). In eukaryotic cells, TFIID is thought to nucleate RNA polymerase II (Pol II) preinitiation complex formation on all protein coding gene promoters and thus, be crucial for Pol II transcription. In a child with intellectual disability, mild microcephaly, corpus callosum agenesis and poor growth, we identified a homozygous splice-site mutation in TAF8 (NM_138572.2: c.781–1G>A). Our data indicate that the patient’s mutation generates a frame shift and an unstable TAF8 mutant protein with an unrelated C-terminus. The mutant TAF8 protein could not be detected in extracts from the patient’s fibroblasts, indicating a loss of TAF8 function and that the mutation is most likely causative. Moreover, our immunoprecipitation and proteomic analyses show that in patient cells only partial TAF complexes exist and that the formation of the canonical TFIID is impaired. In contrast, loss of TAF8 in mouse embryonic stem cells and blastocysts leads to cell death and to a global decrease in Pol II transcription. Astonishingly however, in human TAF8 patient cells, we could not detect any cellular phenotype,
significant changes in genome-wide Pol II occupancy and pre-mRNA transcription. Thus, the disorganization of the essential holo-TFIID complex did not affect global Pol II transcription in the patient’s fibroblasts. Our observations further suggest that partial TAF depletion, and/or an altered TFIID containing a mutated TAF8, could support human development and thus, the absence of holo-TFIID is less deleterious for transcription than originally predicted.

Introduction

In order for a gene to be transcribed, a transcriptional preinitiation complex (PIC) needs to assemble at its promoter. The first complex to bind the promoter is the general transcription factor, TFIID, consisting of the TATA-binding protein (TBP) and 13 different TBP-associated factors (TAFs) (1,2). Following TFIID binding, other general transcription factors (TFIIA, TFIIB, TFIIE, TFIIF and TFIH) and RNA polymerase II (Pol II) are recruited to the promoter (3). Nine of the TAFs contain a histone fold domain (HFD), and form five histone fold pairs. TAF8 is an HFD-containing TAF which forms a histone fold pair with TAF10 in TFIID (4,5). Biochemical studies revealed that the TFIID complex is assembled in a step-wise manner, first forming a stable core complex consisting of TAF5, TAF6-TAF9 and TAF4-TAF12. This core is then bound by TAF2 associated with the TAF8-TAF10 heterodimer, forming the 8TAF complex. Next, TBP and the remaining TAFs join the 8TAF complex, to form holo-TFIID (6,7).

Studies performed in yeast show comprehensively that TFIID is recruited to almost all promoters and is required for the transcription of all Pol II transcribed genes (8). In contrast, individual yeast TAFs are essential for viability. Similarly, germline knock-out of several TFIID subunits in mouse (Taf2, Taf8, Taf10 and Tbp) results in early embryonic lethality around E4.0 (9–12), suggesting that these mammalian TFIID subunits are absolutely essential for blastocyst viability and early development. Furthermore, conditional deletion of Taf4 or Taf10, in embryonic keratinocytes (13,14), deletion of Taf10 in embryonic liver (15), or ablation of Taf7 in CD4+CD8+ thymocytes (9) compromises the viability of the mutant cells, suggesting that these TFIID subunits play an essential role in transcription in these cell types.

Curiously, however, deletion of Taf10 in postnatal liver resulted in a less dramatic phenotype compared with the lack of Taf10 during embryogenesis, with mild effect on transcription of liver genes (15). Similarly, in adult mouse keratinocytes, deletion of Taf10 had no effect on epidermal function and no apparent effect on transcription (13). Also, mouse CD4+CD8+ T-cells lacking Taf7 appear to be viable with no apparent effect on transcription (9). Together these animal studies demonstrate that many cell types absolutely require the canonical TFIID complex for transcription and survival; however, certain cell types are seemingly unaffected by loss of TFIID subunits. How and why these transcription regulatory differences occur amongst different cell types of embryonic and/or adult origin is not understood.

Several different human TFIID subunit coding genes (TBP, MIM: 600775; TAF1, MIM: 313650; TAF2, MIM: 604912; TAF6, MIM: 602955; and TAF13, MIM: 600774) have been implicated in human diseases including many with neurological outcomes and intellectual disability (16–28). A polyglutamine expansion in TBP has a well-characterized gain of function effect resulting in a neurodegenerative disorder, spinocerebellar ataxia-17 (17,19,22,24,26,27,28). In contrast, most of the other reported mutations in genes encoding TFIID subunits are recessive missense mutations in different TAFs found in patients with intellectual disability and various degree of developmental delay and microcephaly (16,18,21,23,25,28). The outcome of these mutations on TFIID structure and function was not assessed in detail, but it can be expected that they cause, to a variable degree, a partial loss of function of the corresponding protein and potentially of TFIID.

In the present study, we identified a homozygous splice site mutation in TAF8 in a patient presenting with intellectual disability, developmental delay and mild microcephaly. The mutation resulted in a frame shift changing the C-terminus of TAF8. The mutant TAF8 protein was undetectable by either western blot analysis or mass spectrometry in cell extracts as well as in TAF-containing complexes purified from the patient fibroblasts. We show that TFIID assembly is impaired in the patient fibroblasts, but astonishingly, there was no significant reduction of the genome-wide RNA Pol II occupancy, elongation rates and pre-mRNA synthesis.

Results

Intellectually disabled child identified with a TAF8: c.781–1G > A mutation

A girl, first seen at 15 months of age, presented with severe global developmental delay and intellectual disability. She was born to non-consanguineous 15- and 17-year-old Hispanic parents after an uncomplicated term pregnancy. Growth parameters have been at –4–5 SD for height and weight and –2 SD for head circumference. Her development has been severely delayed with smiling by 3–4 months and rolling at a year. At 4.5 years she cannot sit or speak. She has developed progressive spasticity and equinovarus foot deformities. However, she does not have seizures and her general health is reasonably good without intercurrent hospitalizations. Her craniofacial findings include brachycephaly, a mildly short forehead, a tented mouth, mild left sided ptosis and a long philtrum (Fig. 1A). Her hands and feet are unremarkable except for the progressive deformities of her feet.

Extensive genetic evaluation revealed a normal chromosomal SNP microarray without excessive homozygosity, normal metabolic studies including glycosylated transferrin, lactate, acylcarnitine profile, very long chain fatty acids and urine organic and plasma amino acids. Two MRI’s have revealed mild microcephaly, reduced myelination, a small cerebellar vermis and partial absence of the corpus callosum (Fig. 1B).

Exome sequencing revealed a homozygous splice site variant in TAF8 (NC_000006.12: g.42077099G > A (NM_138572.2: c.781–1G > A)) (Fig. 1C), whereas no other mutation or variant in genes possibly associated with the phenotype could be identified. Both parents were found to be heterozygous for the same variant. This change was observed in the Genome Aggregation Database (gnomAD) in the heterozygous state in ten individuals, including five of Hispanic origin, out of 277264 alleles.

TAF8: c.781–1G > A variant generates a cryptic splice site resulting in a frame shift in the TAF8 mRNA

As the TAF8: c.781–1G > A mutation in the 3’ end of intron 7 of the TAF8 gene occurred in a predicted splice acceptor site
we analyzed the consequences of this mutation on TAF8 mRNA and protein. To determine the outcome of the mutation, reverse transcriptase-polymerase chain reaction (RT-PCR) analyses were performed on RNA extracted from control and patient fibroblasts using a combination of primers (Supplementary Material, Table S1). These analyses excluded skipping of exon 8 or intron 7 retention (Supplementary Material, Fig. S1A and B). Sequencing of a cDNA fragment generated by RT-PCR using primers annealing in exons 7 and 8 revealed that a cryptic splice acceptor site, one base pair downstream of the original one, was utilized instead of the original splice acceptor site, in the patient fibroblasts. As a result, exon 8 was missing the first G nucleotide (Fig. 1C and D), causing a frame shift. Identical results were observed when the samples were treated with emetine, a non-sense mediated decay inhibitor, confirming that this cryptic acceptor site is exclusively used in the mutant fibroblasts. Quantitative PCR (qPCR) was carried out on mutant and control cDNA, using primers amplifying the junction between exon 2 and 3, which revealed that the abundance of the mutant TAF8 mRNA was mildly reduced by about 30% in the patient cells. This mild reduction is likely caused by non-sense mediated decay, because treatment with emetine restored normal levels of TAF8 mRNA (Fig. 1E).

The TAF8: c.781–1G > A gene encodes a highly unstable protein lacking the C-terminal nuclear localization signal

The amino acid sequence produced from the mutant TAF8 cDNA was predicted using an in silico translation tool. Loss of the first G nucleotide in exon 8, results in a frame shift, affecting the final 50 amino acids of the ‘wild-type’ (WT) TAF8 protein, creating a different C-terminus in the mutant TAF8 protein and a premature stop codon encoded in the 8th exon (Fig. 2A). In the WT TAF8 sequence, the stop codon is in the last 9th exon (Fig. 1C). The region

**Figure 1.** Developmentally delayed child identified with c.781–1G > A mutation in TAF8 gene. (A) Photographs of patient at 3 years of age (top) and 4 years of age (bottom). (B) Brain MRI acquired at 2 years and 7 months of age shows (sagittal on top and two transversal levels on bottom) mildly prominent extra-axial spaces, diffuse thinning of white matter with delayed myelination, borderline enlarged lateral ventricles, short corpus callosum with narrow posterior body and absent splenium, normal brainstem, borderline small cerebellar vermis, and mildly small posterior fossa size. (C) (Top) Patient and control genomic sequences at the intron 7–exon 8 boundary of interest. Asterisk highlights the G > A splice site mutation (in red). Capital letters show exon 8 coding nucleotides. (Bottom) Schematic representation of the TAF8 gene (not to scale) with the position of the mutation indicated by the red asterisk. (D) Sequencing chromatogram highlighting the G nucleotide missing from the beginning of exon 8 in the patient cDNA. (E) RT-qPCR performed with primer annealing to the boundaries between exons 2 and 3, showing abundance of TAF8 mRNA in control and patient cells treated with and without emetine. Error bars represent SEM of three technical replicates. ***P < 0.001.
encompassing the TAF8 HFD, which is required for its pairing with TAF10, is upstream of the frame-shift and is therefore unaffected in the mutant TAF8 protein. The C-terminal region of the WT TAF8 contains a nuclear localization signal (NLS) [30], which is lost in the predicted mutant TAF8 protein (Fig. 2A). WT TAF8 protein has a predicted molecular weight of 34.3 kDa, but migrates around the 43 kDa position [5]. To detect the endogenous mutant TAF8 protein, whole cell extracts (WCEs) were prepared from control and mutant fibroblasts using two different extraction methods (see Materials and Methods) and were subjected to western blot analysis with two different polyclonal anti-TAF8 antibodies (pAb 3477 and 3478), which were generated against the entire TAF8 protein. Unexpectedly, mutant TAF8 protein was not detected in patient fibroblasts (Fig. 2B and C lanes 2 and 3), irrespective of mutant TAF8 mRNA levels (Fig. 1E). To exclude the possibility that the anti-TAF8 pAbs are unable to detect the mutant TAF8 protein because of its unique carboxyl terminus, we generated recombinant baculoviruses expressing the mutant TAF8 protein fused to an N-terminal FLAG tag. The anti-TAF8 pAb readily detected the mutant FLAG-TAF8 fusion protein when highly over-expressed in Sf9 insect cells (Fig. 3A).

These results suggest that the mutant TAF8 protein, as compared with WT, may be less stable and prone to degradation in patient cells. To verify the stability of the mutant TAF8 in a different cellular system, first insect Sf9 cells were infected with baculoviruses expressing either FLAG-tagged WT, or mutant TAF8. When identical total protein amounts from both WT and mutant TAF8 expressing cells were analyzed by western blot, the mutant TAF8 protein was hardly detectable, although high levels of WT TAF8 were detected (Fig. 3B, compare lanes 3 and 4 with 5 and 6). Importantly, however, both WT and mutant TAF8 expressing Sf9 cells contained comparable amounts of TAF8 mRNAs (Fig. 3C), suggesting that the altered final 50 C-terminal amino acids lead to instability of the mutant TAF8 protein. Secondly, to determine whether the mutant TAF8 mRNA is translated into protein at a similar rate as the WT TAF8 mRNA, we generated a single cassette system which co-expresses FLAG-tagged WT, or mutated TAF8 proteins, together with FLAG-green fluorescent protein (GFP), separated by the P2A ‘self-cleaving’ 22 amino acid peptide. Importantly, the P2A sequence mediates ribosome-skipping, enabling the generation of equal amounts of two separate peptide products from a single mRNA [31,32]. When the WT FLAG-TAF8 was expressed with FLAG-GFP from the same cassette (Fig. 3D), using a baculovirus expression system in insect cells, approximately equal amount of the two proteins were detected (Fig. 3E, lane 3). However, in good agreement with our earlier results, when the mutated FLAG-TAF8 was expressed with FLAG-GFP from the same cassette (Fig. 3D), the mutated FLAG-TAF8 was hardly detectable, although the FLAG-GFP, produced from the same mRNA, was readily detected (Fig. 3E, lane 4). These results unequivocally demonstrate that the mutated TAF8, containing the unrelated 50 C-terminal amino acids, is intrinsically unstable.

Degradation of the mutant TAF8 protein could be mediated by the proteasome. In order to test this hypothesis, patient and control fibroblasts were treated with the proteasome inhibitor MG132 for 12 or 18 h, and then western blot analysis was conducted. In untreated samples, the WT TAF8 protein was readily detected, although the mutant TAF8 protein was not. However, in the patient fibroblasts treated with MG132, a protein, smaller than the WT TAF8 protein was detected, likely corresponding to the mutant TAF8 protein (Fig. 2C). These results suggest that degradation of the mutant TAF8 protein is likely carried out, at least partially, by the proteasome.

Together these results show that the TAF8: c.781–1G > A mutation generates a protein with a unique C-terminus that is prone to degradation, causing an apparently complete loss of function of TAF8, further indicating that TAF8: c.781–1G > A, p.Glu261Argfs*38 is most likely the causative mutation in this patient.

Ablation of mouse TAF8 protein in embryonic stem cells results in cell death caused by transcriptional defects

The human patient fibroblasts lacking any detectable TAF8 protein were apparently healthy. Thus, we decided to investigate mouse cells in which TAF8 protein elimination can be induced by Taf8 knock-out. Mouse blastocysts lacking Taf8 die very early during development, at around embryonic day 4.0 (E4.0), because of apoptosis in the inner cell mass (ICM) [12]. To be able to monitor the direct consequences of inducible TAF8 protein ablation, we used Taf8lox/lox: Rosa26-CreERT2 mouse embryonic stem cell lines (mESCs), in which Taf8 deletion can be induced by the addition of 4-hydroxy Tamoxifen (4-OH) (Supplementary Material, Fig. S2A). Cells were seeded at a density of 8 x 10^5 cells per well and every second day, live and dead cells were counted, then 8 x 10^5 cells were reseeded. After 8 days of treatment, there was a massive reduction in the number of live cells, and an increase in dead cells, in the Taf8lox/lox: Rosa26-CreERT2 cultures treated with 4-OH tamoxifen, as compared with control cultures (Fig. 4A–C). The cell death observed in the tamoxifen treated Taf8lox/lox: Rosa26-CreERT2 mESCs correlated with a gradual increase in the deletion of the Taf8 allele (Supplementary Material, Fig. S2B) and a gradual depletion of TAF8 protein, with no TAF8 protein detected after day 6 (Fig. 4D).

Next, we wanted to determine if TAF8 depletion causes global transcriptional defects. As steady-state mRNA levels can be buffered by modulation of mRNA decay, especially in situations where Pol II transcription is globally impaired [8,33,34] and references therein], we measured newly synthesized pre-mRNA using primers that amplify an intron–exon boundary in unspliced pre-mRNA. Pre-mRNA transcript abundance was normalized to the RNA Pol I transcribed 45S pre-rRNA, which is not affected by loss of Taf8. For all tested genes, the abundance of pre-mRNAs was reduced by about 2-fold upon TAF8 depletion, suggesting a global defect in Pol II transcription in the absence of TAF8. Together, these experiments show that mESCs die in the absence of mouse TAF8, similarly to the in vivo Taf8−/− blastocyst phenotype reported by Voss et al. [12], and our preliminary transcriptional analysis suggests that a global decrease in PIC formation correlates with cell death.

The lack of detectable human TAF8 protein dramatically alters the assembly of the TFIID complex in patient fibroblasts

As the patient fibroblasts did not present any obvious phenotype when compared with control fibroblasts, we next assessed if TFIID integrity was affected in the cells harboring the TAF8: c.781–1G > A mutation. To this end, WCEs were prepared from control and TAF8 mutant fibroblasts and subjected to immunoprecipitations (IPs) using anti-TAF7, anti-TBP or anti-TAF10 antibodies to capture the TAF7−/−, TBP−/− or TAF10-containing complexes. The abundance of the TFIID subunits tested was similar in the WCEs from control and patient cells (Fig. 5A–C, lanes 1 and 2). Likewise, the amount of bait protein captured by the corresponding antibody was similar (Fig. 5A–C, lanes 3 and 4).
However, in complexes purified using either anti-TAF7, anti-TBP or anti-TAF10 antibodies, the levels of several of the tested TFIID subunits was markedly reduced in the patient fibroblasts when compared with intact TFIID complexes purified from control cells (Fig. 5A–C, compare lanes 3 and 4).

To detect potential very low levels of mutant TAF8 protein and additional TFIID subunits with a higher sensitivity and in a more quantitative way, anti-TAF10 immunopurified complexes were analyzed by matrix assisted laser desorption ionisation-time of flight mass spectrometry. To be able to compare the composition of complexes purified from control and Taf8 mutant cells, normalized spectral abundance factor (NSAF) values were calculated (35). Although all the known TFIID subunits were identified by mass spectrometry analysis of TAF10-containing complexes isolated from control fibroblasts, we did not detect any TAF8 peptides from the patient cells. We also did not detect TAF1, TAF3, TAF11, TAF13 and TBP, which are supposed to be present in a functional module of TFIID, which does not contain the TAF8/10 histone fold pair (6) (Fig. 5D, Supplementary Material, Table S2). Moreover, the bait-normalized anti-TAF10 immunopurification performed on patient cells identified eight TFIID subunits (TAF2, TAF4, TAF5, TAF6, TAF7, TAF9/9b and TAF12) with significantly less peptide counts than detected in the control sample (Fig. 5D). It should

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**Figure 2.** The TAF8 c.781–1G > A mutant protein lacks an NLS and is degraded by the proteasome. (A) Amino acid sequence of WT TAF8 protein aligned to the in silico predicated amino acid sequence generated by the TAF8: c.781–1G > A mutation, which possesses a unique carboxy-terminus (depicted in red), and lacks an NLS (depicted in grey highlight in the WT sequence). Histone fold domain in the WT sequence is highlighted in blue. (B) Two different protein WCE (WCE1 and WCE2) preparations (see Materials and Methods) were analyzed by western blot. Blots were probed with two different polyclonal anti-TAF8 antibodies (affinity purified pAb 3478 and crude pAb 3477 serum) targeting the entire TAF8 protein, as indicated. (C) Patient and control fibroblasts were treated with MG132 for 12 or 18 h, WCEs were prepared and probed with an anti-TAF8 antibody (affinity purified pAb 3477). The presence of a smaller protein, likely corresponding to the mutant TAF8 protein was detected in the presence of MG132.
be noted that the overall peptide counts were rather low, likely owing to the overall low level of TFIID subunits in fibroblasts relative to other cell types (Supplementary Material, Fig. S3). Therefore, we cannot exclude the possibility that low levels of mutant TAF8 protein are present in the TFIID complexes purified with anti-TAF10 antibodies, but the levels were below the detection limit.

Taken together, these results indicate that in the fibroblasts harboring the TAF8: c.781–1G > A mutation, the lack of detectable mutant TAF8 results in TFIID disassembly to partial TAF-containing complexes, in spite of the fact that the abundance of TFIID subunits (with the exception of TAF8 protein) in the input extracts seem to be unaffected (Fig. 5A–C). Because of the crucial role of TFIID in transcription initiation and during early embryonic mouse development, these observations together offer a likely causative explanation for the homozygous TAF8: c.781-1G > A mutation in the patient phenotype.

A defective TFIID complex does not cause a global reduction of the genome-wide RNA Pol II occupancy and elongation rates

Because TFIID was disassembled in fibroblasts harboring the TAF8: c.781–1G > A mutation (Fig. 5), we set out to verify whether the global recruitment of RNA polymerase II to active promoters may also be compromised. To test this possibility, control and patient fibroblasts were subjected to chromatin immunoprecipitation (ChIP) coupled to...
high throughput sequencing (ChIP-seq), using a mouse monoclonal antibody recognizing the C-terminal domain (CTD) of the largest subunit of Pol II (RPB1, hereafter called anti-Pol II ChIP-seq). Anticipating that subtle difference may be present between the Pol II occupancies at promoters in the control and patient fibroblasts, which may be difficult to quantify, we added a consistent trace amount of Drosophila chromatin to the human fibroblast chromatin at the beginning of the ChIP procedure. As the antibody used in this experiment recognizes the CTD of both human and Drosophila RPB1, the two distinct human chromatin preparations were normalized to the
Drosophila ‘spike-in.’ Surprisingly, Pol II occupancy at selected representative Pol II transcribed genes (Fig. 6A), and genome-wide, analyzed by either K-means clustering (Fig. 6B) or by metagene plots (Fig. 6C), did not change significantly when compared with control cells. Selecting a few representative Pol II-bound promoters (Fig. 6A), the ChIP-seq results were validated by ChIP-qPCR. This validation further confirmed the surprising observation that in patient fibroblasts the Pol II occupancy at the selected promoters was not significantly altered (Supplementary Material, Fig. S4A). Thus, in spite of the complete disorganization of holo-TFIID, Pol II occupancy across all transcribed genes did not change dramatically in patient fibroblasts harboring the TAF8: c.781–1G > A mutation (Fig. 6A–C).

Next, we wanted to determine if Pol II pausing at promoters and/or elongation rates in gene bodies was altered in the patient cells compared with control. To this end we measured the Pol II travelling ratio, which is defined as the ratio of the Pol II occupancy signal at the promoter [from −100 to +300 bp around the transcription start site (TSS)] relative to the Pol II signal in the gene body (from +300 to +2 kb) (36,37). Because the Pol II ChIP-seq signal in the gene body can be close to background, we clustered the genes on the basis of Pol II signal in their gene body (Supplementary Material, Fig. S4B), and calculated Pol II travelling ratio ([37] and see Materials and Methods) for two gene clusters having significant gene body signal. In the first cluster, consisting of 580 genes, the Pol II promoter pausing and gene body signals as well as Pol II travelling ratios were similar between patient and control fibroblasts (Fig. 6D and Supplementary Material, Fig. S4C). In cluster 2, consisting of 1102 genes (Supplementary Material, Fig. S4B), Pol II promoter pausing was slightly decreased and the Pol II signal in the gene body was somewhat higher in patient’s fibroblasts compared with control fibroblasts (Fig. 5E). As a consequence, the Pol II travelling ratio was shifted to the left in patient fibroblasts compared with control (Supplementary Material, Fig. S4D), suggesting that in cluster 2 genes, Pol II was slightly redistributed from promoters to gene bodies in the mutant fibroblasts.
Figure 6. Genome wide RNA polymerase II occupancy and transcription is not significantly effected in fibroblasts harboring the TAF8: c.781-1G > A mutation. (A) UCSC genome browser snapshots of randomly selected representative genes showing Pol II occupancy in control fibroblasts (HFF-1, purple) and TAF8: c.781-1G > A patient fibroblasts (red). Arrows at TSSs show the direction of transcription. (B) Heatmap of anti-Pol II ChIP-seq generated by K-means clustering in a window showing TSS to transcription termination site (TTS) of all genes in control and patient fibroblasts. (C) Metagene plots of all expressed Pol II transcribed genes, generated from spike-in normalized anti-Pol II ChIP-seq, on control fibroblasts and TAF8 mutant fibroblasts. Average Pol II profiles were calculated for every expressed refseq gene from -5 kb region upstream of every TSS, including the gene bodies (GBs), until +5 kb downstream from the TTS. The Y-axis shows mean tag/50 bp densities. (D and E) On the basis of the clustering of Pol II occupancy in the gene bodies (Supplementary Material, Fig. S3B) Cluster 1 (D), containing 580 genes (Supplementary Material, Table S3), and Cluster 2 (E), containing 1102 genes (Supplementary Material, Table S4), profiles from -2 to +2 kb relative to TSS are shown. From these profiles, the corresponding Pol II traveling ratios were calculated and are depicted in the Supplementary Material, Figure S3C and D. In C-E, control graphs are in blue and patient graphs in pink. (F) RT-qPCR of newly transcribed mRNA in control fibroblasts and TAF8: c.781-1G > A patient fibroblasts of 11 representative genes normalized to the Pol I transcribed 45S pre-rRNA. Primers for each gene amplify fragments either only in introns or overlapping intron–exon boundaries from the indicated pre-mRNA. See Supplementary Material, Table S1 for primer sequences.
Loss of TAF8 and holo-TFIID complex do not alter mRNA synthesis from selected genes

The lack of obvious changes in the overall distribution of RNA Pol II despite the undetectable mutant TAF8 protein and holo-TFIID (Figs 5 and 6), suggested that transcription may not be dramatically affected in the patient fibroblasts. To verify this hypothesis, the level of pre-mRNA transcription of randomly selected genes was measured by RT-qPCR. Because steady-state levels of mRNA, determined by assessing total mRNA abundance, are modulated by mRNA decay rates ([8,33,34] and references therein), we measured the abundance of newly synthesized unspliced pre-mRNA relative to the abundance of the newly synthesized RNA polymerase I transcribed 45S pre-rRNA (Fig. 6F). Consistent with the lack of significant Pol II occupancy changes (Fig. 6A–E), we found that the level of transcription of all the tested genes was not affected in the patient fibroblasts as compared with two control fibroblast cell lines (Fig. 6F). In addition, we tested the abundance of some Pol II transcribed histone mRNAs, snRNA and snoRNAs in patient fibroblasts compared with control fibroblasts (Supplementary Material, Fig. S4E) and found no significant changes. Together, these results indicate that in the fibroblasts harboring the TAF8: c.781–1G>A mutation, on average, genome-wide RNA pol II occupancy, Pol II elongation rates and pre-mRNA transcription are not dramatically impaired compared with control fibroblasts.

Discussion

In this study, we identified a patient with severe intellectual disability and developmental delay, who inherited a homozygous splice site mutation in TAF8 (c.781–1G>A), encoding a subunit of the general transcription factor TFIID. In the patient fibroblasts, TAF8 protein was undetectable by western blotting and/or mass spectrometry analyses in WCEs as well as in immunopurified TAF-containing complexes (Figs 2B and C and 5). The much lower levels of mutant TAF8 compared with WT TAF8 protein in the different recombinant expression systems used, clearly shows that the mutant protein is intrinsically unstable (Fig. 3). The identified mutation leads to a seemingly complete loss of TAF8 function, and subsequent disassembly of the TFIID complex. These observations are highly unexpected given the key role attributed to TFIID in transcription. Indeed, in the mouse, germline deletion of Taf8 results in embryonic lethality at E4.0, because of apoptosis of the pluripotent ICM cells of the blastocyst (12). This early lethality demonstrates that in the absence of TAF8 protein mouse blastocysts cannot undergo gastrulation, organogenesis and further embryonic development. Moreover, we also demonstrate here that the inducible deletion of Taf8 in mESCs leads to cell death and globally affects Pol II transcription (Fig. 4). Mouse and human TAF8 proteins are 95.5% identical, and the TFIID complex as a whole is highly conserved between mouse and human. Although suffering from severe neurological deficits, the child identified in this study is physically mostly healthy and has developed well beyond the stage that one would anticipate if this mutation resulted in complete elimination of the mutated TAF8 protein in all cell types during the early stages of development.

We speculate that the TAF8 (c.781–1G>A) mutation did not cause a complete loss of TAF8 function during early embryogenesis and that residual levels of mutated TAF8 could have been sufficient to maintain blastocyst survival and embryo development. In support of this hypothesis, we observed that mESCs with low residual levels of TAF8 protein can survive and divide similarly to controls (see day 4 of 4-OH treatment in Fig. 4B and D); however, when TAF8 protein is completely lost, mESCs rapidly undergo massive apoptosis. Thus, in the patient cells, low residual TAF8 protein incorporated in limiting amounts of holo-TFIID, and partial TAF-containing complexes lacking TAF8 (Fig. 5) are sufficient to maintain Pol II transcription (Fig. 7). Alternatively, in the patient the mutant TAF8 protein might have been present in higher quantities during embryonic development, and/or maybe are higher in different cell types of the patient, to which we had no access. Indeed, it has been recognized that proteins can have different stabilities in different cell types (38,39), likely owing to the interaction with different factors in the varying cellular environments. Nonetheless, from the experiments carried out on the human mutated TAF8 patient fibroblasts and by comparison with the effect of a complete loss of TAF8 function in mice, we conclude that the homozygous TAF8: c.781–1G>A mutation is the disease causing mutation.

In the patient fibroblasts described in this study, TAF8 protein is not detectable, and TFIID was disassembled. However, recruitment of RNA polymerase II to promoters was not significantly affected on a genome-wide scale. This result together with previous knowledge (9,13,15) suggests that the PIC may exhibit more diversity than commonly thought and warrants further mechanistic investigations to determine the heterogeneity of PIC/TFIID function in different cell types.

Patient fibroblasts were used in this study, and although they were useful for performing biochemical analyses, they fail to explain the patient’s neurological phenotype. Our analyses did not reveal a global defect in Pol II occupancy/transcription, which could have been expected considering the extent of TAF8 depletion and TFIID disassembly. However, it is likely that Pol II transcription is mildly impaired as a result of the TAF8 mutation (see Fig. 7) leading to subtle changes in gene expression, which may explain some aspects of the patient phenotype when occurring during neurogenesis and brain development. This is supported by our analyses suggesting putative minor changes in Pol II elongation rates (see for example cluster 2 in Fig. 6 and Supplementary Material, Fig. S4) which should be confirmed by ChIP-seq using antibodies against phosphorylated serine 5 and serine 2 Pol II CTD.

Although many links have been made between mutations in genes encoding TFIID subunits and intellectual disabilities, the outcome of the identified mutations appears different from that of the TAF8: c.781–1G>A mutation identified here. This present mutation appears to cause a nearly complete loss of TAF8 protein, whereas all other mutations identified in other TAF genes were missense mutations expected to result in only partial loss of function of the corresponding gene product. Homozygous missense mutation and compound heterozygous mutations in TAF2 were identified in two independent consanguineous families presenting with autosomal recessive intellectual disability associated with microcephaly and thin corpus callosum (18,40). The impact of the TAF2 mutations on TFIID function was not explored experimentally, but TAF2 is known to interact with TAF8 (7), which could explain the apparent similarities between the TAF2 and TAF8 patients. Recently, homozygous TAF13 missense mutations, identified in two families with mild intellectual disability, growth retardation and microcephaly, were also suggested to affect TFIID structure (25). Missense mutations in TAF6 were also found associated with intellectual disability (16) or an autism spectrum disorder (41). Because TAF6 forms part of
the TFIID core (6), it is plausible that small alterations in its structure could impact TFIID assembly. Of note, homozygous missense mutations in TAF6 were found to be associated with a Cornelia de Lange like syndrome and were reported to affect TFIID assembly (42).

Hemizygous missense mutations in the X-linked TAF1 gene were identified in male patients with a global developmental delay, intellectual disability, facial dysmorphic features, microcephaly and hypoplasia of the corpus callosum (23). The authors predicted that the TAF1 mutations likely disrupted the TAF1–TAF7 interaction, or the TAF1 bromodomain, likely affecting the structure and/or function of TFIID (23). As noted earlier, the clinical phenotypes observed in patients with mutations in several TAFs partially overlap, in particular sharing common features of variable intellectual disability and microcephaly. These similarities suggest that a common molecular mechanism, namely TFIID disruption, results in a secondary variable effect on Pol II transcription in certain, but not all cell types.

In conclusion, our detailed TAF8 patient study, including biochemical characterization of general transcription complexes by mass spectrometry and genome-wide high throughput ChIP analyses, together with previous less mechanistic reports of patients with mutations in TAF genes, demonstrates that perturbed TFIID function is less deleterious for Pol II transcription initiation than originally predicted. Additional studies using animal model systems are needed to further delineate the exact mechanism(s) responsible for the developmental, growth and neurological abnormalities seen in individuals with TAF mutations and to understand how cells can survive without holo-TFIID.

Figure 7. Schematic representation of the TFIID composition within WT, Taf8 KO and patient cells. In WT cells holo-TFIID, containing TBP and 13 TAFs, including WT TAF8, is abundant in cells and drives faithful transcription. In Taf8 knockout cells, such as the mESCs described in Figure 4 of this study, loss of TAF8 results in impaired TFIID and no transcription. In contrast, in the patient fibroblasts either only partial TAF assemblies form, or the tiny (undetectable) amount of mutant TAF8 protein integrates into a TFIID-like complex. Either, or both of these complexes could mediate Pol II transcription initiation in patient fibroblasts.

Materials and Methods

Subject

Written parents’ consent was obtained for conducting the study and showing MRIs and photographs of their affected child.

Exome sequencing

Exome sequencing performed by GeneDx revealed a homozygous splice site variant in TAF8 [NC_000006.12: g.42077099G > A (NM_138572.2: c.781–1G > A)] (Fig. 1C), whereas no other mutation or variant in genes possibly associated with the phenotype could be identified. Both parents were found to be heterozygous for the same variant. This change was observed in the gnomAD in the heterozygous state in 10 individuals, including 5 individuals of Hispanic origin. This variant was observed once out of 12 065 alleles in the NHBLI exome sequencing project.

Plasmids

The baculovirus expression vector for FLAG-tagged human TAF8 WT was PCR amplified from the pPBAC MultiBac vector, expressing TAF8 and TAF10 together, described in (7), and the cDNA was inserted in the pVL1393 baculovirus expression vector digested with Bam HI and Eco RI together with a primer coding for a FLAG-tag on the 5’ end of the hTAF8 cDNA. The mutated TAF8 (FLAG-TAF8 MUT) was generated by site-directed mutagenesis using pVL1393-FLAG-TAF8 WT vector as a template to delete the G
residue at the border of exon 7 and 8 (see Fig. 1D). The WT FLAG-TAF8 and the mutated FLAG-TAF8 cassettes encoding the P2A self-cleaving peptide (GSGATNFSLKQADVEENPG (31,32)) and the FLAG-GFP sequences were inserted into the pVL1393 baculovirus expression vector driven by the polyhedrin promoter. The constructs were verified by sequencing.

Isolation and culturing of human fibroblasts and Drosophila S2 cells

A biopsy was taken from the patient which was used to establish a fibroblast cell line. Control fibroblasts were either HFF1 fibroblasts (ATCC) or control FD137 (male 3 years old). Fibroblasts were cultured on 15 cm² plates in standard fibroblast medium (DMEM/F12 with GLUTAMAX, 10% fetal calf serum (FCS), 15 mM HEPES and pen/strep) at 37°C. Drosophila S2 cells, used for ChIP spike-in were cultured in standard medium (Schneider, 10% heat inactivated FCS and 0.5% pen/strep) at 28°C with no CO2.

Generation of conditional Taf8 mutant mice and isolation of mESCs

A targeting construct was produced using a 17.6-kb fragment from a C57BL/6-derived BAC (RP23–182C6) containing the entire Taf8 gene. A loxP site was introduced 2.5 kb 5’ of exon 1 and a neomycin phosphotransferase expression cassette flanked by FRT sites also containing a single loxP site at the 3’ end was introduced between exons 8 and 9 using the recombinering method (43). This construct was electroporated into C57BL/6 ESCs, and correctly targeted cells line were confirmed by Southern blot analysis. PCR genotyping using three oligonucleotides allowed the identification of the Taf8 exons 1 to 8-deleted allele (Taf8<sup>lox/lox</sup>), the undeleted allele (Taf8<sup>lox<sub>del</sub></sup>) and the WT allele (Taf8<sup>Wt</sup>) simultaneously. Amplification of the WT and undeleted alleles using primers (see Supplemental Material, Table S1) resulted in products of 222 and 345 bps, respectively. Amplification of the deleted allele resulted in a product of 429 bps.

Mice carrying the Taf8<sup>lox</sup> allele were bred to mice carrying the Rosa26Cre-ERT2 allele (44) to produce Taf8<sup>lox<sub>del</sub></sup>; Rosa26-CreERT2 mice. mESCs were isolated from E3.5 blastocysts of Taf8<sup>lox/lox</sup>), the undeleted allele (Taf8<sup>lox<sub>del</sub></sup>), and correctly targeted cell lines were confirmed by Southern blot analysis. PCR genotyping using three oligonucleotides allowed the identification of the Taf8 exons 1 to 8-deleted allele (Taf8<sup>lox/lox</sup>), the undeleted allele (Taf8<sup>lox<sub>del</sub></sup>) and the WT allele (Taf8<sup>Wt</sup>) simultaneously. Amplification of the WT and undeleted alleles using primers (see Supplemental Material, Table S1) resulted in products of 222 and 345 bps, respectively. Amplification of the deleted allele resulted in a product of 429 bps.

WCE preparations

Two methods were used for WCE isolation. In the first method (used for generating the WCE used in the co-IPs, and in the first lane of Fig. 2B), pelleted cells were resuspended in 1× volume of buffer containing 400 mM KCl, 20 mM Tris–HCl pH 7.5, 2 mM dithiothreitol (DTT), 20% glycerol and 1× complete protease inhibitors. The cells were frozen and thawed in liquid N₂ four times before being centrifuged at 13 000 RPM for 10 min to pellet debris and isolate WCE supernatant. In the second method, pelleted cells were first resuspended in 1× volume of 600 mM KCl, 50 mM Tris–HCl pH 7.9, 25% glycerol, 0.2 mM EDTA, 0.5 mM DTT, 5 mM MgCl₂, 0.5% Nonidet P-40 and 1× complete protease inhibitors, then incubated on ice for 10 min, then 3× volume of another buffer containing 25 mM Tris–HCl pH 7.9, 5% glycerol, 0.5 mM DTT, 5 mM MgCl₂, 0.1% Nonidet P-40, and 1× protease inhibitor cocktail was added and incubated on ice for a further 10 min, followed by centrifugation at 13 000 RPM for 10 min to pellet debris and isolate WCE supernatant.

Recombinant protein production

Recombinant baculoviruses were generated as described (47) and used for protein complex production in S9 insect cell culture. Infected insect cells were harvested 48 h after cell infection by centrifugation and stored at −80°C until further use.

Pellets of baculovirus-infected S9 insect cells expressing TAF8 constructs were resuspended in lysis buffer (400 mM KCl, 50 mM Tris–HCl pH 7.9, 10% glycerol, 0.2 mM EDTA, 0.5 mM DTT, containing 1× protease inhibitor cocktail) and crude extracts were prepared by three round freeze thawing cleared by centrifugation.

Coimmunoprecipitation experiments

Confluent plates of TAF8 mutant and control fibroblasts were harvested and lysed in WCE buffer no.1 (described earlier), and 3 mg of WCE from each genotype was used for each coimmunoprecipitation. Protein G or A magnetic Dynabeads (Invitrogen 10002D or 10004D) were incubated with 6 µl of anti-TAF10 antibody (1H8), anti-TAF7 antibody (3475) or anti-GST antibody (15TF21D10) for 2 h, then unbound antibody was removed with two washes in IP100 buffer (25 mM Tris–HCl pH 7.5, 5 mM MgCl₂, 10% glycerol, 0.1% Nonidet P-40, 100 mM KCl, 2 mM DTT and 1× complete protease inhibitors). Fibroblast WCE was made up to 600 µl in IP100 buffer, precleared then added to Dynabeads coupled to antibody, and incubated overnight at 4°C with gentle agitation. The following day the magnetic beads were isolated with a magnetic rack, the unload material was collected, and the beads were subjected to two rounds of washing for 10 min each with IP500 buffer (25 mM Tris–HCl pH 7.5, 5 mM MgCl₂, 10% glycerol, 0.1% Nonidet P-40, 500 mM KCl, 2 mM DTT and 1× complete protease inhibitors), followed by IP100 buffer. For the anti-TAF10 coimmunopurification, proteins bound to the beads were eluted by adding 60 µl of 2 mg/ml competing TAF10 peptide for 4 h, then again for 2 h (48). For the anti-TAF7 co-IP, proteins were eluted with 0.1 M pH 2.8 glycine, then neutralized with 2 M Tris pH 8.8. Co-IP-ed proteins were separated on an 4–12% sodium dodecyl sulphate-polyacrylamide gel electrophoresis (SDS-PAGE) gel, along with the input extract and flow through samples, and were probed with antibodies against TFIID subunits.

Antibodies

Mouse monoclonal (mAb) and rabbit polyclonal (pAb) antibodies raised against the following proteins have been described previously, or were purchased commercially: anti-TBP (mAb 3G3) (49), anti-TAF2 (pAb 3083) (7), anti-TAF4 (mAb 32TA2B9) (11), anti-TAF5 (mAb 1TA1C2) (50), anti-TAF6 (mAb 25TA2G7) (50), anti-TAF7 (mAb 19TA) (51) and (pAb 3475) (52), anti-TAF8 (pAb 3478 and 3477) (52), anti-TAF10 (mAb 6TA2B11) (48) and TAF12 (22TA) (53), anti-GST antibody (mAb 15TF21D10) (54) anti-GFP (Abcam Ab290), anti-γ-Tubulin (Sigma Aldrich T6557), and anti-FLAG (Sigma Aldrich M2).
Mass spectrometry

IP-ed protein samples were analyzed using an Ultimate 3000 nano-RSLC (Thermo Scientific, San Jose, California) coupled in line with a linear trap Quadrupole (LTQ)-Orbitrap ELITE mass spectrometer via a nano-electrospray ionization source (Thermo Scientific). Data were analyzed by calculation of the NSAFbait (52,55). The mass spectrometry proteomics data have been deposited to the ProteomeXchange Consortium via the PRIDE partner repository (56,57) with the dataset identifier PXD008084.

ChIP-seq and ChIP-qPCR

For each ChIP experiment, a minimum of 20 × 15 cm² plates were used per genotype. Approximately 2 × 10⁶ fibroblast cells were seeded onto each 15 cm² plate, then the following day the fibroblasts were fixed in 1% FDA for 10 min at room temperature, then the FDA was quenched with glycine at a final concentration of 125 mM for 5 min at room temperature. After two washes in cold 1× PBS, fibroblasts were scraped, and pelleted. Nuclei were isolated by incubating cells with nuclear isolation buffer (50 mM Tris–HCl pH 8.0, 2 mM EDTA pH 8.0, 0.5% Nonidet P-40, 10% glycerol and 1× protease inhibitors) for 10 min at 4°C with gentle agitation, followed by centrifugation at 13,000 RPM to pellet the nuclei. Nuclei were resuspended in sonication buffer (0.1% SDS, 10 mM EDTA, 50 mM Tris–HCl pH 8.0 and 1× protease inhibitors) then chromatin was sheared with the Covaris E220 sonicator and chromatin concentration was measured with the Qubit 3.0 (ThermoFischer). A minimum of 50 μg of chromatin was used for each IP, which was diluted in ChIP dilution buffer (0.5% Nonidet P-40, 16.7 mM Tris–HCl pH 8.0, 1.2 mM EDTA, 167 mM NaCl and 1× protease inhibitor) and each sample was spiked with 1/10 amount of Drosophila sheared chromatin for normalization. Antibodies used for the ChIP included anti-RPB1 CTD (1PB 7G5) (58), and mouse IgG (Jackson Laboratories) which were incubated with the chromatin overnight with gentle agitation at 4°C. The following day, magnetic beads were isolated and washed for 5 min at 4°C, once with low salt wash buffer (0.1% SDS, 0.5% Nonidet P-40, 2 mM EDTA, 150 mM NaCl, 20 mM and Tris–HCl pH 8.0), high salt wash buffer (0.1% SDS, 0.5% Nonidet P-40, 2 mM EDTA, 500 mM NaCl, 20 mM and Tris–HCl pH 8.0), and LiCl wash buffer (0.2 M LiCl, 0.5% Nonidet P-40, 0.5% sodium deoxycholate, 1 mM EDTA, 10 mM Tris–HCl pH 8.0), then twice with TE buffer, then the beads were incubated in freshly prepared elution buffer (1% SDS, 0.1 M NaHCO₃) at 65°C with shaking to elute complexes. Crosslinks were reversed with by adding NaCl at a final concentration of 0.2 M overnight as well as 50 μg/ml RNase A at 65°C and the following day the samples were treated with 20 μg Proteinase K, 26.6 of 1 M Tris–HCl pH 7.9, and 13.3 μl of 0.5 M EDTA, and DNA was isolated with phenol chloroform. The precipitated DNA was used to generate libraries with the Diagenode MicroPlex Library Preparation kit v2 for ChIP-seq according to the manufacturer’s instructions, with read lengths of 1 × 50 bp, then the samples were sequenced on Hiseq 4000, reads were mapped to the human hg38 and Drosophila dm3 assemblies. Samples were normalized on the basis of Drosophila spike-in as described in (59) then peak calling was performed using the MACS2 software (60). Heatmaps and Metagene plots were generated by the seqMINER software (61). The ChIP-seq data were deposited on the Gene Expression Omnibus (GEO) database with the accession number GSE106252. ChIP-qPCR primers were designed to span a region directly under the RNA Pol II peaks identified by the Pol II ChIP-seq, for randomly selected genes including GAPDH, HSP90AB1, SPARC, RPL13a, ADAMST1 and ID3 as well as a negative region at chromosome 9. Primer sequences are detailed in Supplementary Material, Table S1. The clustering of the genes according their Pol II density in the –2 to +2 kb regions relative to the TSS was carried out by the seqMINER software (61). Pol II traveling ratio was calculated as described in (37), except that the TSS tag density values in +100 to +300 bp regions per gene in clusters 1 and 2 were divided by the corresponding gene body tag density values in +300 to 2 kb regions.

Newly synthesized mRNA analysis

Cells were harvested, and RNA was extracted followed by DNase I treatment, according to the manufacturer’s instructions (Macherey-Nagel). cDNA was generated with Super Script II (Thermo Fischer Scientific) using random hexamer primers, according to the manufacturer’s instructions. To measure genomic DNA contamination in the cDNA preparations, cDNA synthesis reactions were conducted without reverse transcriptase enzyme (~RT). The CP values generated from RT-qPCR with the ~RT samples were always >35 cycles, indicating that genomic DNA contamination contribution was negligible. RT-qPCR primers were designed to amplify intron-exon boundaries (Supplementary Material, Table S1).

Accession numbers

The ChIP-seq data were deposited on the GEO database with the accession number GSE106252. The mass spectrometry proteomics data have been deposited to the ProteomeXchange Consortium via the PRIDE partner repository (56,57) with the dataset identifier PXD008084.

Supplementary Material

Supplementary Material is available at HMG online.

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Conflict of Interest statement. None declared.

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