Brown adipose tissue (BAT) plays an essential role in metabolic homeostasis by dissipating energy via thermogenesis through uncoupling protein 1 (UCP1). Previously, we reported that the TATA-binding protein associated factor 7L (TAF7L) is an important regulator of white adipose tissue (WAT) differentiation. Here, we show that TAF7L also serves as a molecular switch between brown fat and muscle lineages \textit{in vivo} and \textit{in vitro}.

In adipose tissue, TAF7L-containing TFIID complexes associate with PPAR\(\gamma\) to mediate DNA looping between distal enhancers and core promoter elements. Our findings suggest that presence of the tissue-specific TAF7L subunit in TFIID functions to promote long-range chromatin interactions during BAT lineage specification.

Recent studies found that adult humans retain active BAT depots capable of sustaining elevated basal metabolic rates compared to WAT and those higher levels of BAT correlate with lower body mass index\(^1\text{-}^4\). These findings inspired heightened efforts to better understand the formation of BAT during mammalian development. Previously, we showed that TAF7L, an orphan TBP-associated factor cooperates with PPAR\(\gamma\) in directing WAT gene regulation\(^5\). Given the central role of PPAR\(\gamma\) in both WAT and BAT development\(^6\text{-}^7\), we wondered whether TAF7L might also serve as a co-activator regulating the formation of BAT.

To test this hypothesis and assess the functional requirement for TAF7L during BAT development, we performed haematoxylin \& eosin (H&E) staining as well as immune-staining using FABP4 and UCP1 antibodies to delineate regions of BAT in wild type (WT) and \textit{Taf7l} knockout (KO) embryos. As expected, FABP4 stains both WAT and BAT, while UCP1 only stains BAT\(^8\), which is filled with multilocular lipid droplets (Fig. 1A). We found that UCP1\(^+\) BAT in \textit{Taf7l} KO mice become largely disorganized, is significantly reduced in size and contains decreased lipid levels (Fig. 1A and
Measurement of dissected BAT pads showed a ~40% weight reduction of BAT in Taf7l KO animals compared to WT embryos (Fig.1-S1B). Intriguingly, skeletal muscle-like tissue emerges and invades regions that normally contain BAT in Taf7l KO mice (Fig. 1A and Fig. 1-S1D), as revealed by a more pronounced red color of Taf7l KO BAT and the staining of pan-skeletal muscle marker myosin heavy chain (MYHC)9 (Fig. 1-S1 C, E). Our data suggest that loss of TAF7L substantially alters the relative proportion of BAT and muscle lineages in vivo.

Next, we analyzed global gene expression profiles in WT and Taf7l KO BAT by RNA-sequencing (mRNA-seq) (Fig.1B). Over a thousand genes exhibited altered expression levels by >3 fold upon Taf7l KO. In particular, BAT-selective genes such as Ucp1, Pgc1α, Cidea, and Scd1 involved in brown fat differentiation, thermogenesis, and mitochondrial function became significantly down-regulated (Fig. 1C and Fig.1-S2)4,10. Consistent with the observed enhanced skeletal muscle morphological phenotype (Fig. 1A), we observed a concomitant up-regulation of skeletal muscle genes including Myh1-8, Myf5, and Pax3 (Fig. 1C and Fig.1-S2)11. Loss of TAF7L also led to activation of select genes in the formation of cartilage and bone development, but the majority of up-regulated genes appear to be involved in skeletal muscle development and function.

We speculate that because BAT and muscle share common MYF5+ dermotomal precursors12-14, this relationship might favor the switch from BAT to skeletal muscle as shown in Figure 1A. Our findings suggest that TAF7L tips the balance in favor of BAT development at the expense of skeletal muscle.

To examine the effects of TAF7L loss on brown adipocyte differentiation in vitro, we used C3H10T1/2 mesenchymal stem cells to form multiple cell lineages including adipocytes, muscle, cartilage, and bone. We first depleted Taf7l levels in C3H10T1/2 cells by RNA interference using previously described shTAF7L and control shGFP constructs followed by induction of BAT differentiation5. As expected, we observed efficient formation of round fat cells peaking at day 2 post-induction in C3H10T1/2 cells treated with shGFP controls (Fig.2A). By contrast, Taf7l-depleted C3H10T1/2 cells formed elongated muscle-like cells rather than round fat laden adipocytes (Fig. 2A). mRNA analysis of post-induction cells (day2) revealed that specific muscle markers (Myf5, Myod1, and Mef2c) become activated in cells treated with shTAF7L, in some
cases reaching up to ~30% of their expression level in myotubes (Fig. 2B). At the same time, protein (PPARγ and UCP1) and mRNA (Pgc1α, Ucp1, and Cidea) levels of brown adipocyte markers become significantly reduced 5 days post-induction, suggesting an efficient blockade of brown fat differentiation (Fig. 2C and D). We confirmed these results using primary WT and Taf7l KO brown adipocytes (Figure 2-S1). Our findings suggest that loss of TAF7L shifts mesenchymal stem cells in culture from adopting an adipocyte fate toward formation of muscle–like cells.

We next examined TAF7L gain-of-function in vitro using C2C12 myoblasts that can efficiently form myotubes in vitro. Instead of treating C2C12 cells with the normal muscle-inducing protocol, we applied the brown fat differentiation regime to either control C2C12 cells (C2C12.CNTL) or myoblasts ectopically expressing TAF7L (C2C12.TAF7L) and found that only C2C12.TAF7L cells form multilocular brown fat cells (Fig. 2E). RNA analysis showed that TAF7L in C2C12 cells represses the expression of myoblast genes Myf5 and Myod1 while significantly increasing the expression of brown fat-selective genes (Ucp1, Cidea, Pgc1α, and Pparα) post-differentiation (Fig.2F, G and Fig.2-S2), suggesting that TAF7L can drive myoblasts toward the brown fat lineage. These gain of function in vitro results are consistent with the switch from BAT to muscle we observed in Taf7l KO animals in vivo.

Previously, we showed that TAF7L associates with PPARγ when over expressed in 293T cells. Here we wanted to confirm this interaction in C3H10T1/2 cells upon brown fat induction. By using sequential immune-precipitations of doubly tagged TAF7L (FLAG&V5), followed by liquid chromatography coupled with tandem mass spectrometry (LC-MS/MS) and western blotting, we found that TAF7L pulls down a subset of TAFs normally associated with the prototypical TFIID complex. Although this coIPed complex contained a number of canonical TFIID TAF subunits, we nevertheless suspect that it may represent a fat-specific and structurally distinct complex (Fat-TFIID) because it behaved in a manner distinct from canonical TFIID by size exclusion chromatography (data not shown). These immunoprecipitation assays also revealed that PPARγ co-purifies with TAF7L in the Fat-TFIID complex from differentiated C3H10T1/2 cells but not from control FLAG-V5-GFP cells (Fig.3A and B) nor with canonical TFIID lacking TAF7L (data not shown). Although we failed to detect...
PRDM16 in these affinity purification experiments, we did find that ectopically expressed TAF7L and PRDM16 associate with each other in 293T cells (Fig. 3C). These data suggest that TAF7L-containing Fat-TFIID has gained the ability to bind endogenous PPARγ, which we speculate might facilitate its association with distinct cofactors such as PRDM16 in BAT or TLE3 in WAT to differentially regulate brown and white adipocyte formation\textsuperscript{16,17}. To probe the potential functional relevance of these interactions, we next examined the genome-wide co-occupancy of TAF7L and PPARγ at BAT-selective genes in differentiated C3H101/2 adipocytes. Due to the presence of rosiglitazone in the differentiation regime, our previous ChIP-seq analysis of C3H10T1/2 derived fat cells represented a mixture of white and brown fat cells, as manifested by enhanced \textit{Cidea}, \textit{Elov3}, and \textit{Ucp1} expression\textsuperscript{4}. Here we have re-analyzed the binding site data for TAF7L, TBP, and PPARγ focusing on loci of activated BAT-selective genes (\textit{Cidea}, \textit{Prdm16}, and \textit{Ucp1}) (Figure 3D and Figure 3-S1). As shown in Figure 3-S1A and B, TAF7L and PPARγ co-occupy overlapping regions at the \textit{Prdm16} and \textit{Ucp1} loci in post-differentiated (-post) cells, but not at control genes (\textit{Ajap1} and \textit{Bmod2}) nor to the same genes in pre-differentiated cells (-pre). These findings suggest that PPARγ and TAF7L likely function as a coupled activator-coactivator pair that regulates BAT-specific gene transcription much as we previously described for WAT differentiation.

To extend our analysis of TAF7L mechanisms in regulating fat-specific gene transcription, we next employed Chromatin Conformation Capture (3C) to assess its participation in transcription factor-mediated long distance DNA looping at two \textit{Taf7l}-activated genes \textit{Cidea} and \textit{Scd1} (Fig. 3D and Fig. 3-S1)\textsuperscript{18}. These two target genes are bound by TAF7L and PPARγ both at their core promoters as well as at their distal enhancers located ~10kb away. In contrast, TBP only binds at the core promoters of these genes. Our 3C results revealed the likely formation of DNA looping between core promoters and distal enhancers in WT BAT for both loci while significantly reduced looping was seen in BAT lacking TAF7L (Fig.3D and Fig. 3-S1C). These data suggest that the presence/incorporation of TAF7L into the putative Fat-TFIID complex likely mediates or enhances long-range chromatin transactions that influence BAT cell fate. Taken in aggregate, our findings strongly suggest that TAF7L may serve as a key...
component of an alternative TFIID complex that favors brown fat formation versus muscle lineages. We do not fully understand the molecular mechanisms driving the enhanced formation of skeletal muscle upon loss of TAF7L, but we speculate that TAF7L depletion decreases the transcription of BAT-selective genes, which might indirectly de-repress muscle-selective genes thereby switching precursor cells toward a skeletal muscle lineage. Moreover, the presence of TAF7L potentiates the efficient participation of a specialized Fat-TFIID complex in long-range chromatin interactions. Such a DNA looping mechanism is likely mediated through association with the fat-specific transcription factor PPAR\(\gamma\) (Fig.3E). These new findings and our previous studies support the notion that TAF7L functions as a common cofactor regulator required for both WAT and BAT development.

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Methods and materials
Vectors and plasmids
Full length Green Fluorescent Protein (GFP) or Taf7l cDNAs were inserted into pCMV-FlagX10 vector to construct pCMV-Flag-V5-GFP/TAF7L or into pCS2+ vector to construct pCS2+HA-TAF7L; Full length PRDM16 cDNA was inserted into pCMV-FlagX10 vector to construct pCMV-Flag-PRDM16.

Antibodies
TAF4 (BD 612054), TBP (abcam 62126), V5 (Invitrogen,R960-25), HA(abcam 9110), FLAG(Sigma, F31655) MYHC (Millipore 05-716 ), FABP4 (abcam 66682),
UCP1(abcam10983), TAF7L (prepared in-house), Pol II (monoclonal 8GW16, protein-A purified), PPARγ (sc-7196), ANTI-FLAG M2 Affinity Gel (Sigma, A2220), Anti-V5-agarose affinity gel (Sigma, A7345).

**Cells culture, stable cell line establishment**

C3H10T1/2 and C2C12 cells were cultured in high glucose DMEM with 10% fetal bovine serum at 5% CO2.

C3H10T1/2 cells stably expressing Flag-V5-GFP/TAF7L were established by transfection of the pCMV-FLAG-V5-GFP/TAF7L plasmids followed by two weeks of 1ug/ul neomycin selection.

**Brown adipocyte differentiation, Oil red O staining, and C2C12 myogenesis**

For adipogenesis, C3H10T1/2, control and FLAG-TAF7L-expressing C2C12, and Flag-V5-GFP/TAF7L-expressing C3H10T1/2 cells were grown in high glucose DMEM supplemented with 10% fetal bovine serum. At confluence, cells were exposed to induction medium containing dexamethasone (1 μM), isobutylmethylxanthine (IBMX, 0.1 mM), T3 (1uM), insulin (5 μg/ml), rosiglitazone (5 μM), and 10% FBS. 3 days later, cells were further cultured in high glucose DMEM containing insulin (5 μg/ml), T3 (1uM), and rosiglitazone (5 μM) until they were ready for harvest.

For Oil red O staining, pre- and post-differentiated C3H10T1/2 cells, shGFP and shTAF7L-treated C3H10T1/2 cells, C2C12.CNTL and C2C12.TAF7L cells were washed once in PBS and fixed with freshly prepared 4% formaldehyde in 1×PBS for 30 min, followed by standard Oil red O staining method described previously5.

C2C12 myogenesis procedure followed a previous study5.

**RNA isolation and real-time PCR analysis**

Total RNA from cultured cells or mouse tissues was isolated using QIAGEN RNaseq Plus mini columns according to the manufacturer's instructions (Qiagen). For RT-qPCR analysis, 1 μg total RNA was reverse transcribed using cDNA reverse transcription kit (Invitrogen). SYBR green reactions using the SYBR Green PCR Master Mix (Applied Biosystems) were performed according to the manufacturer's instruction using an ABI 7300 real time PCR machine (Applied Biosystems). Relative expression of mRNA was
determined after normalization to cyclophilin gene. Student's t-test was used to evaluate statistical significance.19

**Western blot analysis, immunoprecipitation, and silver staining**

Whole cell extracts were prepared from cells by homogenization in lysis buffer containing 50 mM Tris–Cl, pH 8.0, 500 mM NaCl, and 0.1% Triton X-100, 10% glycerol and 1 mM EDTA, supplemented with protease inhibitor cocktail (Roche) and phenylmethylsulphonyl fluoride (PMSF). 15 μg of whole-cell lysates were separated by SDS-PAGE and transferred to nitrocellulose membrane. For immunoblotting, membranes were blocked in 10% milk, 0.1% Tween-20 in TBS for 30 min, and then incubated with TAF7L, FLAG, β-actin, PPARγ and TBP antibodies for 2 hr at room temperature; detailed Western blotting procedure was performed as previously described.20

3mg whole-cell extracts from FLAG-V5-GFP/TAF7L post-differentiated adipocytes were immunoprecipitated with 100ul of ANTI-FLAG M2 Affinity Gel under the conditions of 0.3 M NaCl, 0.2% NP-40 at 4°C overnight. After extensive washing with buffer containing 0.15 M NaCl and 0.1% NP-40, proteins were eluted from the affinity gel with 100ug/ml FLAG peptide in 0.3 M NaCl Tris buffer. Elutions from both samples were subsequently immunoprecipitated with 40ul Anti-V5-agarose affinity gel antibody with a similar procedure as above. After extensive washes, proteins were eluted with 40ul pH2.5 glycine for three times and immediately neutralized with 12ul of pH9.0 Tris-Cl. 10ul of eluted samples were subjected to 10% SDS-PAGE and followed by standard silver staining or by western blotting analysis with V5, PPARγ, TAF4, and TBP antibodies to detect tagged-proteins in the inputs and associated proteins as previously described.4 The remaining samples were precipitated by 20% trichloroacetic acid and the precipitates were sent to liquid chromatography-tandem mass spectrometry (LC-MS/MS) to detect peptides derived from proteins pulled-down by FLAG-V5-GFP/TAF7L.

500 μg whole-cell extracts from 293T cells transfected with HA-TAF7L and FLAG-PRDM16 were immunoprecipitated with FLAG or HA antibodies at 4°C for overnight under the conditions of 0.3 M NaCl and 0.2% NP-40, 30 μl protein A/G beads were added and incubated for additional 2 hr at 4°C, after extensive washing with buffer containing 0.15 M NaCl and 0.1% NP-40, remaining beads were subjected to 10% SDS-PAGE and followed by western blotting analysis with FLAG and HA antibodies to
detect tagged-proteins in the inputs and IPs as described previously\textsuperscript{5}.

Animals and genotype analysis

The derivation of \textit{Taf7l} KO mice has been previously described\textsuperscript{14}. All animal experiments were performed in strict accordance with the recommendations in the Guide for the Care and Use of Laboratory Animals of the National Institutes of Health. All of the animals were handled according to animal use protocols (#R007) approved by the Institutional Animal Care and Use Committee (IACUC) of the University of California, Berkeley. Mice were maintained on a standard rodent chow diet with 12 hr light and dark cycles. \textit{Taf7l} KO mouse line was maintained on a C57BL/6 background. Genotyping was performed by PCR as previously described\textsuperscript{21}.

Immunohistochemistry

For histological analysis on interscapular BAT of E18.5 embryos from WT and \textit{Taf7l} KO mice, freshly-harvested mouse embryos were genotyped and the interscapular regions of embryos were transversally dissected and then fixed in 10\% formaldehyde for 24 hr at 4°C; tissue was embedded in paraffin using the microwave method as described and then sectioned into 8–10 \( \mu \)m to mount on slides\textsuperscript{19}. The following immunohistochemistry by haematoxylin and eosin (H&E) staining and FABP4, UCP1, and MYHC immunostaining were performed using the method described previously\textsuperscript{5}.

Preparation of primary brown adipocytes and brown fat differentiation

Fresh interscapular brown adipose tissues were removed from 3 week old euthanized WT and \textit{Taf7l} KO mice and finely minced, digested with 0.25\% trypsin for 30 min at 37°C, and centrifuged for 5 min at 2,000g to get rid of debris. The pellet was resuspended in culture media before plated on gelatin coated plates. Cells were cultured at 37°C in high glucose DMEM supplemented with 20\% FBS. Brown adipocyte differentiation and staining were followed the same procedure as C3H10T1/2 cells.

mRNA-seq libraries preparation and deep sequencing

Total RNA was extracted from BAT, carefully excised to get rid of surrounding tissue based on texture and color, of 6 WT and 6 \textit{Taf7l} KO mice. RNA was extracted separately for each mouse by RNeasy Plus Mini Kit (Qiagen) and then pooled for WT or \textit{Taf7l} KO samples; 4 \( \mu \)g of each RNA pool was used to purify mRNA using oligo (dT) and
subsequently converted into multiplexed mRNA-seq libraries using mRNA-Seq Trueseq Kit (Illumina). Samples were multiplexed and sequenced in one lane on an Illumina HiSeq 2000 sequencer. 50 bp single-end reads were used for both samples; each sample produced over 30 million reads.

**Digital gene expression of mRNA-seq and gene ontology analysis**

Reads were mapped to the mouse transcriptome (mm10), using TopHat\textsuperscript{22,23}, version v1.4.0., with default parameters. We then applied cufflinks\textsuperscript{24}, version v1.3.0, using the default parameters except: --max-mle-iterations 1, to estimate the digital expression levels at each transcript. Gene ontology analysis was done using DAVID Bioinformatics Resources 6.7.

**Chromosome Conformation Capture (3C)**

3C analysis was performed as previously described on WT and Taf7l KO BAT\textsuperscript{18}, which was carefully excised to get rid of all other possible tissue based on the brown color and the tissue texture. 2mg of freshly dissected interscapular BAT from 6 WT or 6 Taf7l KO mice were minced, homogenized extensively to nearly single cells and washed, crosslinked with 1% formaldehyde for 15mins at 4°C and then quenched with 0.125M glycine for 5 mins.

Crosslinked BATs were lysed and the chromatin was digested with 8 U HaeIII (NEB) for the Cidea and Scd1 loci. Digested fragments were cleaned and subsequently ligated with 60 units T4 DNA ligase (Invitrogen) for 4 hours at 16°C. Following proteinase K digestion and reverse-crosslinking at 65°C overnight, DNA fragments was recovered by phenol-chloroform extraction.

Control templates were generated using individual BAC clones covering Cidea or Scd1 locus (Bacpac). 10ug of BAC DNA was digested with 20U HaeIII and then randomly ligated with 10 units T4 DNA ligase in 50ul volume.

3C primers were designed upstream and downstream of the core promoter site (P). Primers annealing to distal enhancers (D) corresponding to TAF7L and PPARγ binding sites on either Cidea or Scd1 were used as anchor points. 3C analysis was done by qPCR using as a primer pair the anchor point primer and one annealing to region under investigation. Each data point in WT and Taf7l KO BAT was normalized by the BAC control template and presented as interaction frequency.
Data availability

Raw and mapped sequencing reads are available from the National Center for Biotechnology Information's GEO database (http://www.ncbi.nlm.nih.gov/geo/) under accession number GSE55797. Primer sequences are listed in Supplementary File 1.

Figure legends

Figure 1, TAF7L is required for proper mouse brown adipose tissue (BAT) formation. A, Wild type (WT) and Taf7l knockout (Taf7l KO) interscapular BAT from E18.5 embryos was stained with haematoxylin and eosin (H&E), FABP4, and UCP1 (25X magnification shown). Rightmost panel shows FABP4 staining at 200X magnification. B, Left panel: scatterplot shows gene expression profile in WT and Taf7l KO BAT tissue; red dots represent genes up-regulated in Taf7l KO, green dots represent down-regulated genes in Taf7l KO. Right panel: gene ontology analysis shows functional groups of genes changed by at least 3 fold. C, Difference in expression of either muscle-specific (top) or brown fat (bottom) genes between WT and Taf7l KO BAT with red and green shades showing degrees of up- and down-regulation, respectively.

Figure 1-S1, Depletion of TAF7L shifts BAT to muscle lineage. A, UCP1-stained BAT samples as in Figure 1A for 3 additional animals (2WT and 1 TAF7L KO). B, BAT weight from WT (n=10) and Taf7l KO embryos (n=9). C, Photographs of whole BAT from 1-month and 4-month old WT and Taf7l KO mice. D, High-magnification images (200X) of similar areas of WT and Taf7l KO embryonic BAT stained with FABP4 antibody. E, Haematoxylin and eosin (H&E) and myosin heavy chain (MYHC) staining of Taf7l KO embryos shows muscle-tissue structures along the BAT at high magnification (200X).

Figure 1-S2, RT-qPCR comparison of gene expression in WT and Taf7l KO BAT. Expression levels of BAT-specific genes Ucp1, Cidea, Pgc1a, and Scd1, muscle-specific genes Myhc, Myh1, Myh4, and Myot. Common fat marker gene Adipoq, adipocyte progenitor marker Dlk1, Taf7l and Cyclophilin were used as control. mRNA levels in WT BAT was assigned to 1, and mRNA levels of each gene in Taf7l KO BAT was compared
to WT BAT. RT-qPCR was normalized to the levels of *Cyclophilin* (L). *p<0.05, data is mean and s.e.m is from triplicates.

Figure 2, TAF7L bidirectionally regulates BAT/muscle lineages. A, 2 days (2D) and 5 days (5D) post-induced control (shCNTL) and TAF7L knockdown (shTAF7L) C3H10T1/2 cells stained with Oil Red O. B, mRNA levels of myoblast genes *Myf5*, *Myod1*, *Mef2c*, *Myhc*, and *Mlc* in 2D post-induced shCNTL and shTAF7L C3H10T1/2 cells, compared to C2C12-induced myotubes. C, Protein levels of TAF7L, PPARγ, and UCP1 in 5D post-induced shCNTL and shTAF7L cells; Pol II was used as loading control. D, mRNA levels of brown fat genes (*Pgc1a, Ucp1, and Cidea*) in 5D post-induced control and shTAF7L cells. E, 5D post-induced control (C2C12.CNTL) and TAF7l-expressing C2C12 (C2C12.TAF7L) cells were stained with Oil Red O. F, mRNA levels of myoblast genes *Myf5* and *Myod1* in pre-induction cells. G, mRNA levels of BAT-specific genes (*Ucp1, Cidea, Cox8b, Pgc1a, and Ppara*) 5D post-differentiation. *p<0.05, data is mean and s.e.m is from triplicates.

Figure 2-S1, TAF7L is required for brown fat cell differentiation from primary brown adipocytes. A, isolated brown adipose progenitor cells from WT and *Taf7l* KO mice was induced with brown adipocyte differentiation regime for 5 days and then stained with Oil Red O. B and C, mRNA levels of fat-selective genes (B) and muscle-selective genes (C) pre-differentiation. *p<0.05, data is mean and s.e.m is from triplicates. D, Expression levels of *Taf7l* and brown adipocyte marker genes on cells from A, the expression levels of genes in 5D post-induced *Taf7l* knockout cells were compared to WT cells, whose levels were assigned to 1.

Figure 2-S2, Gene expression analysis of control (CNTL) and TAF7L-expressing (TAF7L) C2C12 cells from 0 (0D) to 5 days (5D) post BAT-induction. mRNA levels of *Ucp1, Prdm16, Cidea, Pgc1a, Taf7l*, and *Myod1* are plotted as relative value to 0D CNTL cells, data is mean and s.e.m is from triplicates.

Figure 3, TAF7L within a Fat-TFIID associates with PPARγ and facilitates DNA looping
formation. A, Silver staining shows co-immunoprecipitated proteins in FLAG-V5-GFP-expressing (GFP, lane 1) and FLAG-V5-TAF7L-expressing (TAF7L, lane 2) C3H10T1/2 differentiated fat cells. Comparative LC-MS/MS analysis identified peptides matching with TFIID subunits (TAF1, TAF4, TBP, TAF5, TAF6, and TAF10) and PPARγ in TAF7L-expressing but not in GFP-expressing cells. FKBP15 and ACSL1 are representative non-specific associated proteins. B, Western blot analyzing input and immunoprecipitated protein levels of PPARγ, TAF4, TAF7L, and TBP in samples from A. C, HA tagged TAF7L and FLAG tagged PRDM16 were overexpressed in 293T cells, immunoprecipitations were performed on both FLAG and HA antibodies and followed by Western blotting with FLAG and HA antibodies. D, upper panel, schematic picture shown the distance between distal enhancer (D) and core promoter (P) of Cidea gene; middle panel, read accumulation of TAF7L, TBP, and PPARγ on Cidea locus in differentiated fat cells from ChIP-seq analysis; bottom panel, 3C experiments assess long-range DNA interactions between the TAF7L/PPARγ binding distal enhancer (D) and core promoter (P) sites of Cidea in WT and Taf7l KO BAT. ▲, anchor point. Also see Figure 3-S1C. E, Model shows TAF7L mediating regulatory DNA looping to specify BAT differentiation from mesenchymal stem cells (MSC).

Figure 3-S1, TAF7L colocalizes with PPARγ on core promoters and enhancers of BAT-specific genes. ChIP-seq read accumulation for TAF7L and PPARγ in both pre- and post-differentiated C3H10T1/2 cells on Ajap1/Prdm16 locus (A) and Ucp1/Bmod2 locus (B). C, 3C experiments assess long-range interactions between the TAF7L/PPARγ binding distal enhancer (D) and core promoter (P) sites of Scd1 in WT and Taf7l KO BAT. ▲, anchor point.

Supplementary File 1, Primer sequences for RT-qPCR experiments (upper panel) and 3C experiments (lower panels).

Reference


Figure 1

A

H&E  25X  α-FABP4  α-UCP1  BAT

WT  TGFβ KO

B

- Genes up-regulated by 3 fold (n=1743)
  - GO terms:
    - Muscle development (p=17.56)
    - Muscle tissue development (p=2.17)
    - Cartilage development (p=8.71)
    - Bone development (p=2.81)

- Genes down-regulated by 3 fold (n=1239)
  - GO terms:
    - Brown fat cell differentiation (p=8.71)
    - Thermogenesis (p=5.73)
    - Mitochondria (p=1.63)
    - Metabolism (p=3.03)

C

[Heatmap or similar data representation]
Figure 2
Figure 3

A

B

C

D

E

![Graphical representation of Figure 3](image)