Short Communication

Polypyrimididine tract binding protein 1 regulates the activation of mouse CD8 T cells

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The RNA-binding protein polypyrimididine tract binding protein 1 (PTBP1) has been found to have roles in CD4 T-cell activation, but its function in CD8 T cells remains untested. We show it is dispensable for the development of naïve mouse CD8 T cells, but is necessary for the optimal expansion and production of effector molecules by antigen-specific CD8 T cells in vivo. PTBP1 has an essential role in regulating the early events following activation of the naïve CD8 T cell leading to IL-2 and TNF production. It is also required to protect activated CD8 T cells from apoptosis. PTBP1 controls alternative splicing of over 400 genes in naïve CD8 T cells in addition to regulating the abundance of ∼200 mRNAs. PTBP1 is required for the nuclear accumulation of c-Fos, NFATc2, and NFATc3, but not NFATc1. This selective effect on NFAT proteins correlates with PTBP1-promoted expression of the shorter Aβ1 isoform and exon 13 skipped Aβ2 isoform of the catalytic A-subunit of calcineurin phosphatase. These findings reveal a crucial role for PTBP1 in regulating CD8 T-cell activation.

Keywords: CD8 T cells · cellular activation · gene regulation · RNA binding proteins

Introduction

Upon antigen recognition through peptide presentation by MHC I and co-stimulation, a series of signaling pathways are triggered in CD8 T cells which ultimately lead to cytoskeleton remodeling, metabolic reprogramming, and transcription factor activation [1]. The calcium, MAP kinase, and PKC signaling pathways function by inducing nuclear translocation of the nuclear factor of activated T cells (NFAT), activator protein 1 (AP-1), and nuclear factor kappa-light-chain-enhancer of activated B cells (NFκB) transcription factors [2]. These coordinate the expression of multiple genes involved in a programme of proliferation and differentiation. In
addition, they induce expression of pro-inflammatory cytokines, including tumor necrosis factor (TNF), interferon gamma (IFN-γ), and interleukin-2 (IL-2) [3, 4]. Cytokine expression by CD8 T cells is also regulated posttranscriptionally; this has been shown to be context-dependent and is orchestrated by calcium, MAPK, and PKC pathways, which may coordinate the activities of RNA binding proteins that regulate the stability and translation of mRNA [5].

Polypyrimidine tract binding protein 1 (PTBP1) is a ubiquitously expressed RNA binding protein, which regulates alternative splicing (AS), polyadenylation, mRNA abundance, and internal ribosome entry site-mediated translation [6, 7]. Previous work has shown that Ptbp1 is important in B lymphocytes for the germinal centre reaction [8, 9] and that it is redundant with its paralogs Ptbp2 in early B-cell development [10] and with Ptbp3 in mature B cells [11]. PTBP1 also promotes CD154 (CD40-ligand) expression by mouse CD4+ T cells by binding the 3'UTR of Cd40lg mRNA [12]. In human CD4 T cells, PTBP1 has been found to be necessary for the optimal signaling via the NFκB and ERK1/2 pathways [13]. Upon knockdown of PTBP1, CD4 T cells showed a reduced proliferation, survival, and expression of effector molecules [13]. PTBP1 also regulated 3' end formation of the human CD5 transcript in T cells [14]. Whether PTBP1 regulates CD8 T cell responses is unknown.

Here, we identify PTBP1 as a regulator of the early signaling events that occur upon CD8 T-cell activation required for the production of IL-2 and TNF. In the absence of PTBP1, the accumulation of c-Fos and NFATC3 in nuclei is reduced. We show that PTBP1 regulates the AS of many genes including the Aβ catalytic subunit of calcineurin, which may contribute to the activation of CD8 T cells.

Results and discussion

PTBP1 is essential for the expansion and differentiation of CD8 T cells in vivo

We generated mice with conditional deletion of Ptbp1 in T cells (Cd4CreTgPtbp1fl/fl—abbreviated to P1KO). PTBP1 protein was absent in P1KO CD8 cells from P1KO mice and, as expected, PTBP1-KO CD8 T cells expressed PTBP2 (Supporting information Fig. S1A) due to loss of the repressive effect of PTBP1 on PTBP2 expression [6]. The numbers of naïve CD8 cells in the spleen and LNs of P1KO mice were similar to control Cd4CreTg mice (Supporting information Fig. S1B). Following introduction of the OT1 transgenic T-cell receptor, we found that ~90% of naïve P1KO OT1 cells lacked PTBP1 protein (Supporting information Fig. S1C). Taken together, these data indicate that the accumulation of naïve CD8 T cells shows no absolute requirement for PTBP1.

To identify cell-intrinsic roles for PTBP1 in CD8 T-cell activation, we transferred naïve OT1 cells into recipient mice and subsequently infected them with Listeria monocytogenes expressing ovalbumin (attLm-OVA). P1KO OT1 cells showed a severe expansion defect compared to PTBP1-sufficient OT1 cells (Fig. 1A). Upon co-transfer of P1KO and control OT1 cells and infection with attLm-OVA, the expansion of PTBP1-deficient OT1 cells was also greatly reduced (Fig. 1B and C). Transferred P1KO OT1 cells underwent fewer cell divisions than control OT1 cells (Fig. 1D) and more frequently stained positive for active-Caspase-3 (Fig. 1E and Supporting Information S1D) showing that PTBP1 was required for proliferation and protected cells from apoptosis in vivo.

Among the few P1KO cells present both the amounts and the proportions of cells producing TNF, IFN-γ and IL-2 were reduced (Fig. 1F and G). Moreover, while all PTBP1-deficient cells contained granzyme B (Fig. 1F), there was an approximately twofold reduction (Fig. 1H). P1KO OT1 cells present at days 4 and 5 postinfection also had diminished expression of KLRG1 (Fig. 1I and Supporting Information S1E). Thus, PTBP1 contributes to the differentiation of effector CD8 T cells and is necessary for the production of effector molecules in vivo.

PTBP1 is necessary for optimal activation of CD8 T cells

Following in vitro stimulation of naïve P1KO CD8 cells with anti-CD3 and anti-CD28 antibodies, a number of cells that have undergone two or more cell divisions were reduced compared to control CD8 cells (Supporting Information Fig. S2A). This defect was accompanied by reduced amounts of IL-2 in the culture supernatants (Supporting information Fig. S2B). PTBP1-deficient CD8 T cells responded to the addition of recombinant IL-2 showing increased proliferation (Supporting Information Fig. S2C and D), but the %−divided cells and the proliferation index were always less than control cells (Supporting Information Fig. S2D). The frequencies of active-Caspase-3+ cells were similar between naïve or 24-h stimulated P1KO and control CD8 cells (Supporting Information Fig. S2E). However, at 48 h, both nondivided and proliferating PTBP1-deficient CD8 T cells had increased frequencies of active-Caspase-3+ cells at all division numbers compared to controls (Supporting Information Fig. S2E). Thus, even when supplemented with IL-2, PTBP1 is necessary for the optimal proliferation and survival of activated CD8 T cells in vitro. The reduced proliferation, IL-2 production, and increased apoptosis of antibody-stimulated nontransgenic cells reflect the in vivo behavior of the P1KO OT1 cells. The results are consistent with findings in human CD4+ T cells following retroviral delivery of PTBP1 RNAi and point to a requirement for PTBP1 function long-after stimulation in maintaining cell survival [13]. Moreover, as reported in human CD4+ T cells, we find in mouse CD8+ T cells that PTBP1 is required for the optimal production of IL-2.

PTBP1 regulates early signaling events in CD8 T cells

To investigate further the role of PTBP1 in the early events following CD8 T-cell activation, we measured TCR-induced expression
Figure 1. PTBP1 is essential for the expansion and differentiation of CD8 T cells in vivo. (A) Proportions of CD45.2+ cells (among live eFluor780− lymphocytes) identified by flow cytometry in the blood of B6.SJL (CD45.1+) mice which received 1000 naïve CD8 T cells from OT1 control (CD45.2+ OT1Tg) or OT1 P1KO (CD45.2+ OT1Tg Cd4CreTg Ptbp1fl/fl) mice and were infected with attLm-OVA 1 day after OT1 cell transfer. Each symbol shows the mean ± SD of four mice per group. Data shown are from one experiment. (B) Note that 75,000 naïve CD8 T cells from CD45.1+ OT1Tg Cd4CreTg and 75,000 naïve CD8 T cells from CD45.2+ OT1Tg Cd4CreTg Ptbp1fl/fl P1KO mice were adoptively co-transferred into CD45.1 and CD45.2 double-positive mice, which were infected with attLm-OVA 1 day after cell transfer. Shown are the proportions of CD45.1+ or CD45.2+ cells among live (eFluor780−) CD8a+ splenocytes at different days postinfection. (C) Numbers of CD45.1+ or CD45.2+ cells among live (eFluor780−) CD8a+ splenocytes per mouse at different days postinfection treated as shown in (B). padj values shown are from Sidak’s multiple correction test after two-way ANOVA from the log-transformed data. (D) Cell Trace Violet profiles of CD45.1+ or CD45.2+ CD8 OT1 cells (identified as live eFluor780− and CD8a+; splenocytes of mice treated as shown in (B) and re-stimulated with 50 nM SIINFEKL peptide for 3 h in the
of the transcription factor NR4A1. NR4A1 was reduced in PTBP1-deficient CD8 T cells compared to control (Fig. 2A). Additionally, the production of IL-2 and TNF by P1KO OT1 cells was impaired when cells were stimulated with peptide (Fig. 2B). The production of TNF and IL-2 following treatments of P1KO CD8 cells with PMA+Ionomycin was also defective pointing to a defect in T-cell activation that cannot be overcome through bypass of the receptor proximal transduction pathways (Fig. 2B). Previous work in activated human CD4+ T cells indicated a requirement for PTBP1 for optimal IkBa degradation [13]. The nuclear translocation of p65-RelA is a sensitive measure of TCR signal transduction [2], yet we found this to be similar in control and P1KO CD8 cells (Fig. 2C). By contrast, the amounts of c-Fos, a component of AP-1, was reduced in PTBP1-deficient CD8 T nuclei (Fig. 2D). The reduced nuclear accumulation of c-Fos might be a consequence of a reduced amount of c-Fos protein. These findings suggest a role for PTBP1 in regulating the early events upon TCR signaling with an impact in the optimal expression of cytokines.

**PTBP1 controls AS of calcineurin Aβ in CD8 T cells**

We performed RNA-seq on the transcriptome of naive CD8 T cells from P1KO and control mice. In the absence of PTBP1, approximately 100 mRNAs were both increased and decreased in abundance (FDR < 0.05, Supporting Information Fig.S3A—Source Data 1); as this number was calculated with no thresholds on expression values, we conclude PTBP1 has a modest effect on transcript levels in resting CD8+ T cells. We found changes in AS among 453 genes (Supporting Information Fig. S3B—Source data 2) indicating a role for PTBP1 in splicing regulation. Among these changes, we focused on the catalytic subunits of calcineurin, a serine-phosphatase with an important role in TCR signaling.

Resting CD8 T cells express all three paralogs of calcineurin A with calcineurin-Aα (encoded by Ppp3ca) and Aβ (encoded by Ppp3cb), expressed at much higher protein copies/cell than Aγ (encoded by Ppp3cc) (Fig. 3A) [15]. In P1KO CD8 T cells, no differences were found in the total abundance of the mRNAs encoding the paralogs (Fig. 3B). However, Ppp3cb had increased inclusion of exon-13 in PTBP1-deficient compared to control cells (Fig. 3C and Supporting Information Fig. S3C). PTBP1 binds to the introns flanking exon-13 (Fig. 3C), where it is predicted to promote exon skipping. This is supported by the FPKMs estimated for each isoform by cuffnorm, which indicates a decrease in the exon 13-skipped Ppp3cb-201 isoform in PTBP1-deficient cells (Fig. 3D). In addition, Ppp3cb has an alternative polyadenylation site in intron-12. This generates an extended exon-12 (transcript Ppp3cb-206) encoding a shorter protein termed Aβ1 with an alternative C-terminus with distinctive localization and function and the absence of an inhibitory domain present in the Aβ2 variant [16, 17]. Previous work had indicated this polyadenylation event can be regulated by muscleblind like splicing regulator 1 [18]. PTBP1 binds to the extended exon-12 upstream of the intronic polyadenylation site (Fig. 3C) where it is predicted to promote the use of this alternative polyadenylation site [14]. The FPKM for this isoform is decreased in PTBP1-deficient cells (Fig. 3D).

We performed RT-PCR to validate the AS and polyadenylation of Ppp3cb in P1KO naïve CD8 T cells. This confirmed a significant increase in the proportion of transcripts that included exon 13, concomitant with decreased skipping of this exon, and a trend toward decreased usage of the alternative polyadenylation site, although this did not reach significance (Supporting Information Fig. S3D-F). Decreased usage of this alternative polyadenylation site in P1KO cells was further supported by long read Oxford Nanopore Technologies RNA sequencing in naïve and activated CD8 T cells (Supporting Information Fig. S3G-H), albeit many reads were truncated and the number of reads that could be unambiguously assigned to isoforms were low. Therefore, PTBP1 promotes the expression of the shorter Ppp3cb-206 isoform encoding Aβ1 and a variant of Aβ2 which lacks 10 amino acids encoded by exon 13.

Calcineurin acts on many substrates and is essential for the dephosphorylation and nuclear translocation of NFAT. In the mouse C2C12 myogenic cell line, the inclusion of Ppp3cb exon 13 has been linked specifically to reduced NFATc3 nuclear translocation [19]. In activated CD8 T cells, we found that NFATc1 translocation was normal in the absence of PTBP1 (Fig. 3E and Supporting Information Fig. S4A). This suggested that TCR-activated calcium signaling was not globally impaired. However, NFATc2 fluorescence intensity in the nucleus of PTBP1-deficient CD8 T cells was slightly reduced compared to control CD8 T cells indicating reduced translocation (Fig. 3F and Supporting Information Fig. S4B). For NFATc3-staining, stimulation resulted in the rapid loss of nuclear staining and a gradual increase in the frequency NFATc3 between 2 and 4 h that was diminished in nuclei from PTBP1-deficient cells (Fig. 3G and Supporting Information Fig. S4C). As there were no differences in the NFATc1, NFATc2, and NFATc3 transcript isoforms [20–22] arising from AS and polyadenylation in PTBP1-deficient CD8 T cells compared to control cells.

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Figure 2. PTBP1 regulates early signaling events in CD8 T cells. (A) Histograms showing NR4A1 staining of control (black histograms) or PTBP1-deficient (blue histograms) naïve CD8 T cells activated ex vivo with SIINFEKL peptide (10 nM) for the time indicated. Gates show positive populations. Below, graphs show proportions of NR4A1-expressing cells and geometric mean fluorescence intensity (gMFI) of the staining within cells identified as positive. Each dot represents a biological replicate. Data shown are from one representative experiment out of three. (B) Histograms (left) show IL-2 and TNF-α staining in unstimulated (unst) CD8 T cells (eFluor780–CD8α+ TCRβ+) and stimulated with SIINFEKL peptide (OVApep) or PMA/Ino. (C) Histograms show p65-RelA staining. (D) Histograms show c-Fos staining.
(data not shown), our data implicate PTBP1 in controlling the selective nuclear translocation of NFAT family members indirectly via an intermediate. We highlight the control of alternative isoforms of calcineurin Aβ as a candidate mechanism to mediate this effect.

**Concluding remarks**

Our data from CD8 cells is broadly consistent with the findings in human CD4 T cells and indicate an important role of PTBP1 in the early activation events following TCR stimulation. These events require the nuclear translocation of a collection of transcription factors that together stimulate the gene expression programme of the activated T cell [23, 24]. It is likely that PTBP1 will regulate gene expression at times late after activation and this may include genes that protect activated T cells from apoptosis. In the present work, we validate Ppp3cb as a direct target of PTBP1 and its isoforms as candidates for selective regulators of early events in T-cell activation; we anticipate that further regulation mediated by PTBP1 of other transcripts will contribute to T-cell activation via AS, polyadenylation, and control of their abundance.

**Materials and methods**

**Mice**

Mice were derived from crossing the following transgenic strains: *Ptbp1* 

\[ \text{Ptbp1}^{\text{fl/fl}} (\text{Ptbp1}^{\text{tm1Msol}}) \] [25], *Cd4CreTg* (*Tg(Cd4-cre)1Cwi*) [26], and *OT1Tg* (*Tg(TcraTcrb)1100Mjb*) [27]. All mice used were between 7 and 20 weeks old and were age- and sex-matched within experiments.

**Cell and nuclei isolation**

Single cell suspensions were prepared from spleen and LNs by passing the organs through 70 and 40 μm cell strainers consecutively. Naïve CD8 T cell isolation was achieved by depletion of cells bound by biotinylated antibodies: CD4 (GK1.5), CD11b (M1/70), CD11c (N418), CD19 (1D3), B220 (A3-6B3), CD105 (M17/18), TER-119, gamma/delta TCR (GL3), Gr1 (RB6-8C5), NK1.1 (PK136), F4/80 (BM8), CD44 (1M7), and magnetic Dynabeads M-280 Streptavidin (ThermoFisher, cat#:11206D).

**Naïve CD8 OT1 cell transfer and att-Lm-OVA infection**

For analysis of naïve OT1 cell expansion in the blood (Fig. 1A), 1000 naïve OT1 cells isolated by fluorescence activated cell sorting (eFluor780 –, CD8a (53-6.7)-APC +, CD44(IM7)-BB515 –) and mixed with 10^6 carrier splenocytes from B6.SJL mice were adoptively transferred into B6.SJL recipient mice. One day later, mice were infected with 5 × 10^6 C.E.U. attLm-OVA lacking actA [28] intravenously. For analysis of naïve OT1 cell expansion in the spleens of CD45.1/CD45.2 recipient mice (F1 of B6. SJLxC57BL/6 mice) analyzed at days 3, 4, and 5 postinfection with 5 × 10^6 C.E.U. attLm-OVA (Fig. 2B), naïve OT1 cells were isolated by magnetic negative depletion as described above. The numbers of naïve OT1 transferred before attLm-OVA infection are specified in the figure legends.

**Ex vivo T-cell stimulation**

Note that 10^5 naïve CD8 T cells isolated by magnetic negative depletion as described above were cultured in RPMI 1640 medium (Dutch modification) with 20 mM HEPES (ThermoFisher, cat#:22409-015), 50 μM β-mercaptoethanol, 100 PMA/ionomycin (PMA/Iono) for 4 h from control (black histograms) or PTBP1-deficient (blue histograms) in the presence of brefeldin A. Gates show positive populations. Graphs (right) show proportions of IL-2 or TNFα-expressing cells and geometric mean fluorescence intensity (gMFI) of the staining within cells identified as positive. Each dot represents a biological replicate. Data shown are from one representative experiment out of two. OT-1 nuclei were identified as CellTrace Violet™ CD3εlow.

Note that 10^5 naïve CD8 T cells isolated by magnetic negative depletion as described above were cultured in RPMI 1640 medium (Dutch modification) with 20 mM HEPES (ThermoFisher, cat#:22409-015), 50 μM β-mercaptoethanol, 100
Figure 3. PTBP1 regulated gene expression programs in CD8 T cells. (A) Ppp3ca, Ppp3cb, and Ppp3cc protein levels analyzed by mass spectrometry by Marchingo et al. [15] in naïve CD8 T cells. (B) Abundance of Ppp3ca, Ppp3cb, and Ppp3cc mRNA (Normalized Reads Counts from DESeq2 analysis) in naïve CD8 T cells from control (Ptbp1fl/fl) and P1KO mice (Ptbp1fl/fl Cd4CreTg). (C) Sashimi plot of Ppp3cb transcript in naïve CD8 T cells from merged replicate datasets for control (Ptbp1fl/fl) and P1KO mice (Ptbp1fl/fl Cd4CreTg), showing mRNA-seq read coverage and reads that map to exon–exon junctions (arrows; numbers indicate reads that map to that junction), and cross-link sites detected by iCLIP for PTBP1 (below). (D) Graph showing FPKM of three different isoforms of Ppp3cb (shown in the genome reference Fig. 3C) in naïve CD8 T cells from control (Ptbp1fl/fl) and P1KO mice (Ptbp1fl/fl Cd4CreTg). (E–G) Graphs showing proportions of nuclear translocation of NFATc1 (D), NFATc2 (E), and NFATc3 (F) and geometric mean fluorescence intensity (gMFI) of the staining within cells identified as positive. OT-I nuclei were identified as CellTrace Violet+ CD3elow. Data shown for NFATc1 and NFATc3 are from one experiment with four biological replicates out of two. Data shown for NFATc2 are from one experiment out of three (n = 3).

units/ml penicillin and streptomycin, GlutaMAX, and 10% FCS on 96 flat-bottom well plates precoated with anti-CD3 (2C11, BioXcell) and CD28 (37.51, BioXcell) antibodies. For tracking cell division, naïve CD8 T cells were labeled with either Cell-Trace Violet (ThermoFisher, cat#:C34557) or Cell-Trace Yellow (ThermoFisher, cat#:C34567), following the manufacturer’s recommendations but labeling cells for only 10 min. Also 20 ng/mL (20 units in 200 μL) recombinant murine IL-2 (PeproTECH, cat#:212-12-100UG) was added to cultures when indicated. For measuring cytokine production, splenocytes from OT-I transgenic mice were stimulated with SIINFEKL peptide (N4) (ProImmune, Think Peptides) at the concentration of 10^{-7}M.
or with PMA and ionomycin. For nuclear staining, naïve OT-I transgenic CD8 T cells were labeled with CellTrace Violet and stimulated by SIINFEKL loaded splenocytes in a 96-u-bottom plate. Nuclei were isolated using previously described methods [29].

IL-2 amount from supernatant of naïve CD8 T cells activated for 24 h with 5 μg/mL anti-CD3 with 1 μg/mL anti-CD28 antibodies was analyzed by Meso Scale Discovery assay at the Core Biochemical Assay Laboratory (Cambridge).

Flow cytometry

Single cell suspensions from the spleen and LNs were prepared by passing the tissues through a 70 μm and subsequently a 40 μm cell strainer. Antibody binding to Fc receptors was blocked with the anti-FcγIII 2.4G2 monoclonal antibody. Cell surface staining and the Fixable Viability Dye eFlour780 (eBioscience, cat#: 65-0865-18) was carried out at 4°C. The antibody used were as follows: CD45.2-BV785 (104, BioLegend, cat # 109839), CD45.2-PE (104, eBioscience, cat # 12-0454-82), CD45.1-BUV395 (A20, BD, cat # 565212), CD8α-BUV737 (53-6.7, BD, cat # 563786), and CD8α.BUV737 (53-6.7, BD, cat # 564297).

For intracellular staining, cells were fixed with BD Cytofix/Cytoperm (BD, cat#: 554722) and either washed with BD Perm/Wash buffer (BD, cat#: 554723) or with PBS, frozen at –80°C in 10% DMSO and 90% FCS at least overnight, washed with PBS, re-fixed with BD BD Cytofix/Cytoperm for 5 min on ice, and washed with BD Perm/Wash buffer. Permeabilized cells were incubated for 20 min at room temperature with anti-FcγIII 2.4G2 antibody in BD Perm/Wash buffer. Intracellular staining was carried out with the following antibodies diluted in BD Perm/Wash buffer:

- Active-caspase-3-GranzymeB-PE (C92-605, BD, cat #561011), Active-caspase-3-BV650 (C92-605, BD, cat #564096), GranzymeB-PE-CF594 (GB11, BD, cat #562462), IFNγ-ITC (XM1G12, BioLegend, cat #505806), IL-2-PE (JES6-5H4, BioLegend, cat #503808), KLRG1-BV421 (2F1, BD, cat #562897), NR4A1-PE (12.14, eBioscience, cat #12-5965), PTBP1-AF647 (clone 1, ThermoFisher, cat #32-4800), PTBP2-AF488 (S43, from Michele Solimena), and TNFa-BV650 (MP6-XT22, BioLegend, cat #506333).
- PTBPs anti-bodies were conjugated to different fluorophores with anti-body labeling kits from ThermoFisher (AF488, cat #A20181; AF647, cat #A20186). Intracellular staining for PTBPs was carried out at least for 4 h at room temperature or overnight at 4°C. Cytokine staining of splenocytes after attLm-OVA infection was carried out after re-stimulation of CD8 T-cell enriched splenocytes with 50 nM SIINFEKL peptide for 3 h with brefeldin A (Biolegend, cat#: 420601).

Staining of nuclei was carried out with the following antibodies diluted in 1× PBS and 2% FCS and 0.3% Triton X-100 overnight at 4°C: anti-cFos (9F6)-PE (Cell Signalling), anti-p65-RelA (D14E12)-PE (Cell Signalling), anti-NFATc1 (7A6)-AF488 (Biolegend), anti-NFATc2 (D43B1)-AF488 (Cell Signalling), and anti-NFATc3 (Cell Signalling).

mRNASeq libraries

RNA was extracted from naïve CD8 T cells using the RNeasy Micro Kit (cat# 74004, Qiagen). Naïve T cells were isolated from four female control (Ptbp1fl/fl) and four P1KO (Cd4CrePtbp1fl/fl) mice as described in the CD8 T-cell isolation section. Note that 10 ng RNA was used and seven PCR cycles were carried out to synthesize cDNA with the SMART-Seq® v4 Ultra® Low Input RNA Kit (cat # 634891, Takara). RNAseq libraries were prepared with the Nextera XT DNA Library Preparation Kit (cat# FC-131–1096, Illumina) and carrying out eight PCR cycles. mRNAseq libraries were sequenced on an Illumina HiSeq2500 on a 2×125 bp paired-end run.

RNA sequencing

mRNA-seq libraries were run across two lanes of a HiSeq 2500 RapidRun. After demultiplexing, data from the two lanes were merged for each sample before trimming TrimGalore (v0.6.1; https://www.bioinformatics.babraham.ac.uk/projects/trim_galore/), and quality control FastQC (https://www.bioinformatics.babraham.ac.uk/projects/fastqc/) and FastQ Screen (https://www.bioinformatics.babraham.ac.uk/projects/fastq_screen/). Reads were mapped to the GRCh38 using HISAT2 (v2.1.0) [30], suppressing unpaired or discordant alignments and taking into account known splice sites from the Ensembl Mus_musculus GRCh38.90 annotation. FPKM were calculated using cuffnorm from Cufflinks (v2.2.1) [31], with geometric normalization, using the Ensembl Mus_musculus. GRCh38.90 annotation and genes with average FPKM < 1 or that encode immunoglobulin or T-cell receptor genes were excluded from subsequent analysis.

Differential mRNA abundance analysis

Raw read counts were generated using Seqmonk (v1.45.0; https://www.bioinformatics.babraham.ac.uk/projects/seqmonk/) over mRNA features with merged isoforms, using the Ensembl Mus_musculus GRCh38.96 annotation, assuming unstranded, paired reads. Downstream analysis was performed using R v3.5.3. Genes encoding immunoglobulin or T-cell receptor gene segments were excluded from analysis; genes with average FPKM < 1 in both conditions were also excluded. DESeq2 analysis (v1.20.0) [32] comparing naïve control and Ptbp1 KO was performed using default parameters, with "normal" log2 fold change shrinkage. Significantly differentially expressed genes were defined as those with FDR-adjusted p value < 0.05 and mean normalized read counts > 5.

Transcriptome-wide AS analysis

Fastq files that had been trimmed using Trim Galore as above were further trimmed to 123 bases, discarding any reads below
this length and any resulting unpaired reads, using Trimmmomatic [33]. The resulting reads were mapped with HISAT2 as above, but only retaining uniquely mapped reads in the BAM files (samtools view -bS -F 4 -F 8 -F 256 -q 20). BAM files were sorted and indexed (samtools v1.9) before running rMATs (v4.0.2, turbo) to compare control and PTBP1 KO cells, specifying a read length of 123 and that libraries are paired end, using the Ensembl Mus_musculus.GRCm38.96 annotation. rMATS results that only counted reads spanning junctions were used for subsequent analysis. Significantly altered splicing events that were significantly altered for each comparison were defined as those with FDR < 0.05, absolute inclusion level difference > 0.1, and supported by a total of at least 35 reads mapping to the included or skipped form in control or P1KO (sum of four biological replicates), and that are from genes with FPKM > 1 in at least one condition (mean of four biological replicates). Duplicated events where the coordinates of the alternatively spliced exon were identical were filtered to retain only the event with the lowest FDR or, where tied, with the greatest number of supporting reads. Sashimi plots to visualize read coverage and junction-spanning reads were generated using the IGV genome browser, after merging replicate datasets into a single BAM file using samtools.

PTBP1 iCLIP library preparation and sequencing

CD8 T cells isolated by negative magnetic depletion as described above but omitting the anti-CD44 biotinylated antibody from control (Ptbp1fl/fl) or P1KO mice were stimulated ex vivo with plate-bound anti-CD3 (2C11, 5 μg/mL), anti-CD28 (37.51, 1 μg/mL), and 20 ng/mL murine IL-2 for 24 h. Note that 9 × 10^5 CD8 T cells were stimulated per well in 48-well plates. After 24 h, cells were irradiated 1 × with 254 nm UV light (150 mJ/cm^2). iCLIP libraries were carried out following an improved iCLIP protocol [34]. Cells were lysed with 50 mM Tris-HCl, pH 7.4, 100 mM NaCl, 1% Igepal CA-630 (Sigma 8896), 0.1% SDS, and 0.5% sodium deoxycholate lysis buffer. After sonication between 0.5 mg and 0.2 mg, protein lysates in 1 mL were incubated with low amounts of RNase I (0.05–0.01 units, ThermoFisher, cat#: EN0602) and 4 units Turbo DNase (Ambion, cat#: AM2238) for 3 min at 37°C. PTBP1-RNA complexes were precipitated with 2 μg of anti-PTBP1 antibody (ThermoFisher clone 1, cat#: 32-4800) and 20 μL of Protein A/G beads (Pierce, cat#: 88802) overnight rotating at 4°C. RNA isolation and cDNA library preparation was carried out as described in Lee et al. [34] using the L3-IR-App adapter to visualize and retro-transcribe the RNA cross-linked to PTBP1. We carried out four biological replicates from control (Ptbp1fl/fl) and two biological replicates from P1KO activated CD8 T cells.

iCLIP computational processing

Data from CD8 T cells and from mouse B cells [8] were processed on the iMaps platform (https://imaps.genialis.com/iclip). Reads were deduplicated based on their random barcodes, trimmed using Cutadapt (v2.4) [35] and mapped to the GRCm38 mouse genome build using STAR (v2.7.0f) [36]. Significant cross-link sites and clusters were identified for each replicate separately, except for mouse B cells where replicates were merged, using the iCount pipeline (v2.0.1dev; https://icount.readthedocs.io/en/latest/index.html). The gene and feature type to which each cross-link site or cluster was assigned was determined by considering all transcript isoforms from the Ensembl Mus_musculus. GRCm38.96 annotation. If multiple genes or feature types were prioritized, the most likely was chosen in a hierarchical way: first, protein coding isoforms were prioritized other transcript biotypes; second, prioritizing high confidence isoforms with transcript support level < 4; third, by a hierarchy of feature types (3'UTR > CDS > 5'UTR > intron > non-coding exon > non-coding intron); fourth, higher confidence isoforms (based first on transcript isoform level and second on whether they have a CCDS) were prioritized. iCLIP data were visualized using a shiny app (https://github.com/LouiseMatheson/iclip_visualisation_shiny); all cross-links merged across replicates are displayed, without filtering for significant sites.

Long read RNA sequencing library preparation

Libraries were generated essentially according to the CELLO-seq protocol [37] with the following modifications: The addition of UMIs by splint ligation was omitted. Reverse transcription was performed using 1 ng total RNA isolated from a pool of cells. PCR reactions were performed using standard Oxford Nanopore Technologies barcoded primers under the following conditions: 90°C for 45 s; 90°C for 15 s, 64°C for 30 s, 72°C for 10 min; 72°C for 5 min. PCR products were purified using 0.6X AMPure XP beads (Beckman).

MinION sequencing of long read cDNA libraries

Equal quantities of each cDNA sequencing library were pooled using a total of 200 fmol, before sequencing on a MinION R9.4.1 flow cell using the SQK-LSK109 kit (1D sequencing). The standard MinKNOW procedure was used for a 72-h run, according to manufacturers’ instructions.

Long read data processing and analysis

Raw reads were base called using Guppy (4.0.11) in fast mode. Sequence reads with a mean sequence quality of 7 were classified as passed reads; these were first demultiplexed with the Guppy debarcoder option and then processed with pychopper (2.5.0) in order to identify, orient, and trim the full-length cDNA reads. Next, the pychopped reads were aligned with minimap2 (2.17) with splice aware setting (-ax splice), which is compatible with pipelines for further isoform identification analysis. The
sequences were aligned to mouse genome reference GRCm38. BAM files for replicate datasets were merged using samtools, before filtering for reads overlapping at least one exon of Ppp3cb, and that contained at least one splice junction. These were visualized and a sashimi plot generated in IGV, and the number of reads that could be unambiguously assigned to isoforms including exon 13, skipping exon 13, or utilizing the alternative 3'UTR and polyadenylation site within the extended exon 12 was counted manually.

**RT-PCR**

cDNA using 0.5 μg total RNA, 50 ng of oligo(dT), 250 ng of random hexamers, and five units of Transcriptor Reverse Transcriptase (Roche) was prepared as described in manufacturer's protocol. PCRs with 50 ng of prepared cDNAs were carried out using Jumpstart Taq polymerase (Sigma) according to manufacturer's protocol. Primers binding to exons 11 (forward; 5'-TGCAGCGCCGAAGAAAATCA), 14 (reverse; 5'-GTATGGCCGGTGTTCAGGGA), and to the alternative 3'UTR in the extended exon 12 (reverse; 5'-GAGCTGCTCCCCATTCACCC) of Ppp3cb were included in each reaction. Cycling parameters were 94°C for 30 s, 60°C for 30 s, and 72°C for 60 s for 30 cycles. Minus RT cDNA for each sample were included. For visualization and quantification of the PSI values, PCR products were resolved in a QIAxcel Advanced System (QIAGEN) and PSI calculated within the QIAxcel ScreenGel software.

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Abbreviations: AS: alternative splicing · PTBP1: polypyrimidine tract binding protein 1

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