Splicing factors control *C. elegans* behavioural learning in a single neuron by producing DAF-2c receptor

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Alternative splicing generates protein diversity essential for neuronal properties. However, the precise mechanisms underlying this process and its relevance to physiological and behavioural functions are poorly understood. To address these issues, we focused on a cassette exon of the *Caenorhabditis elegans* insulin receptor gene *daf-2*, whose proper variant expression in the taste receptor neuron ASER is critical for taste-avoidance learning. We show that inclusion of *daf-2* exon 11.5 is restricted to specific neuron types, including ASER, and is controlled by a combinatorial action of evolutionarily conserved alternative splicing factors, RBFOX, CELF and PTB families of proteins. Mutations of these factors cause a learning defect, and this defect is relieved by DAF-2c (exon 11.5+) isoform expression only in a single neuron ASER. Our results provide evidence that alternative splicing regulation of a single critical gene in a single critical neuron is essential for learning ability in an organism.
diverse neuronal populations are essential for physiological functions of the brain. It is widely accepted that neuronal identities are determined by combinatorial actions of transcription factors that are controlled by regulatory networks of microRNAs and signal transduction pathways. It has also been suggested that post-transcriptional regulation by alternative splicing plays critical roles in many steps of neuronal development. Dysregulation of alternative splicing events in the brain has been linked to several neurodegenerative and neuropsychiatric disorders, highlighting the physiological importance of alternative splicing in the brain. Several brain-enriched RNA-binding proteins (RBPs), such as NOVA, RBFOX and polypyrimidine tract-binding (PTB) families of proteins, have been found to regulate alternative splicing events in the nervous system. Altered expression of these splicing factors causes substantial defects in survival and/or function of neurons. Extensive studies using genome-wide approaches and computational methods have unveiled the neural splicing networks controlled by these proteins and some biologically relevant targets in brain function. However, functional roles of such a considerable number of splicing isoforms in the brain are yet poorly understood.

Cell-type specificity of alternative splicing patterns can be ensured by combinatorial actions of alternative splicing factors. Bioinformatic analysis using Bayesian networks reveals that cis-regulatory elements of neural splicing factors, NOVA and RBFOX, coexist in a number of sequences surrounding cassette exons, which implies their combinatorial splicing regulations of a set of neuronal genes. Recent findings suggest that this type of regulation is used to determine neuronal subtype identities in vivo. Combinatorial regulation of the CELF family protein, UNC-75, and the Hu/EuAV family protein, EXC-7, is required for alternative splicing of a JIP3 homologue unc-16 in C. elegans. Predominant inclusion of a cassette exon of unc-16 occurs in cholinergic neurons that co-express UNC-75 and EXC-7, but not GABAergic neurons that express only UNC-75, suggesting that predominant inclusion of the cassette exon is achieved by combinatorial expression of a pair of splicing factors. The in vivo mechanism to generate diverse neuronal properties via alternative splicing is just beginning to be understood.

To analyse alternative splicing events and their relevance to physiological and behavioural functions, the C. elegans nervous system is useful. C. elegans hermaphrodites have only 302 neurons, which are classified into 118 classes based on their morphology and neuronal connectivity. Among 60 sensory neurons, amphid sensory neurons comprising 12 neuron classes are the major sensors of environmental chemicals and temperature. C. elegans uses the amphid sensory neurons for innate and learned behavioural responses to seek suitable conditions and to avoid unfavourable conditions. We have reported that insulin/IGF receptor signalling in the taste receptor neuron ASE right (ASER), the right-sided ASE class of amphid sensory neurons, plays a pivotal role in taste-avoidance learning during which worms learn to avoid salt concentrations encountered under starvation conditions. C. elegans has two isoforms of the insulin/IGF receptor homologue, DAF-2α and DAF-2γ, which are produced by skipping and inclusion of exon 11.5, respectively. Only DAF-2α (exon 11.5+)/DAF-2γ (exon 11.5−), increases after food deprivation and can support taste-avoidance learning. In contrast, DAF-2α is more effective than DAF-2γ in otherdaf-2-regulated processes, including dauer formation and longevity. Thus, DAF-2 isoforms play distinct roles in the biological processes, and DAF-2c has a critical role in the ASER neuron in taste-avoidance learning.

Here we report that alternative splicing ofdaf-2 exon 11.5 is regulated in a cell-type-specific manner. Using fluorescence-based splicing reporters and a messenger RNA (mRNA)-tagging method, we demonstrate thatdaf-2 exon 11.5 inclusion occurs in restricted neuron classes, including ASE. We investigate the molecular mechanism that underlies this neuron-class-specific alternative splicing event. We find thatdaf-2 exon 11.5 inclusion is regulated by RBFOX, CELF and PTB families of proteins, whose mammalian homologues act in the brain as alternative splicing factors. Furthermore, our analysis reveals that these splicing factors are co-expressed only in the restricted neuron classes, wheredaf-2 exon 11.5 inclusion predominantly occurs. Ectopic expression ofPTB-1/PTB was sufficient to induce predominant inclusion ofdaf-2 exon 11.5 in the nervous system, suggesting that the PTB-1 expression is a critical determinant of the neuron-class specificity of exon 11.5 inclusion. We also show that mutations of these factors cause defects in taste-avoidance learning. Remarkably, expression ofDAF-2c (exon 11.5+) in a single neuron, ASER relieved the learning defect of the splicing factor mutant. Thus, combinatorial action of the splicing factors in ASER determines the neuronal property required for learning ability by controlling alternative splicing ofdaf-2.

Results

Cell-type-specific alternative splicing ofdaf-2 exon 11.5. To visualize the alternative splicing patterns ofdaf-2 exon 11.5, we created splicing reporters, in which either green fluorescent protein (GFP) or red fluorescent protein (RFP) complementary DNA (cDNA) was fused with the modifieddaf-2 genomic fragments that contain exon 11.5, the flanking introns and parts of exons 11 and 12 to monitor exon skipping or exon inclusion, respectively (Fig. 1a; Supplementary Fig. 1a). When both reporters were expressed under a common ubiquitous promoter, GFP signals, which reflect exon skipping, were widely observed. In contrast, RFP signals, which reflect exon inclusion, were observed only in restricted cell types, including neurons, the somatic gonad and hypodermal cells in the tail region (Fig. 1b). When these reporters were exclusively expressed in the nervous system, GFP and RFP expressions were observed in distinct neurons in a mutually exclusive manner (Fig. 1c; Supplementary Movie 1), suggesting that exon 11.5 selection occurs in a neuron-class-specific manner.

To further investigate the reporter expression patterns in each neuron class, these reporters were expressed under several neuron-class-selective promoters. All the amphid sensory neurons, including ASER, predominantly expressed E11.5+::RFP (Fig. 1d; Supplementary Fig. 2a). The predominant RFP signals were also observed in some other neuron classes, including gas-sensing BAG and mechanosensory OLL neurons, and a small subset of inter- and motor-neurons (Fig. 1d; Supplementary Fig. 2b). By comparison, predominant GFP signals were observed in some other neuron classes, including IL2 sensory neurons, and inter- and motor-neurons (Fig. 1d; Supplementary Fig. 2b and c). Swapping of GFP and RFP cDNAs in the exon 11.5-skipping/inclusion reporters reversed the expression patterns of GFP and RFP (Supplementary Fig. 2d), which indicated that neuron-class-specific expression of the exon 11.5-skipping/inclusion reporters was not due to the different characteristics of these fluorescent proteins, but because of the differential splicing patterns.

To confirm neuron-class-specific alternative splicing of exon 11.5 in the endogenousdaf-2 transcripts, we isolated mRNAs specifically from the amphid sensory neurons or other classes of
neurons, using an mRNA-tagging method\textsuperscript{20}, and compared enrichment levels of total daf-2 and the daf-2 (exon 11.5 +) isoform. We utilized the gpc-1 and the glr-1 promoters for expression of FLAG-tagged poly(A)-binding protein in 10 neuron classes, including 7 amphid sensory neurons, and in 15 classes of inter- and motor-neurons, respectively (Fig. 2a–d). Total daf-2 mRNAs were enriched at comparable levels in the gpc-1- and glr-1-expressing neurons, whereas daf-2 (exon 11.5 +) mRNA was enriched at a significantly higher level in the gpc-1-expressing neurons (Fig. 2e). Reverse transcription–PCR (RT–PCR) analysis confirmed the significantly higher inclusion level of daf-2 exon 11.5 in the gpc-1-expressing neurons than in the glr-1-expressing neurons (Fig. 2f). The expression patterns of the exon 11.5-skipping/inclusion reporters in a variety of neurons, including all the amphid sensory neurons and some inter- and motor-neurons, under the casy-1 promoter\textsuperscript{21} (hereafter this transgene is referred to as ‘neuronal exon 11.5-skipping/inclusion

**Figure 1** | Cell-type-specific alternative splicing of daf-2 exon 11.5. (a) Schematic of daf-2 alternative splicing reporters for monitoring skipping (top) and inclusion (bottom) of exon 11.5. GFP and RFP fusion proteins are expressed by exon skipping and exon inclusion, respectively. (b–d) Expression patterns of the exon 11.5-skipping/inclusion reporters driven by a ubiquitous promoter, the eaf-1A.1 promoter (b), a pan-neuronal promoter, the H20 promoter (c) and the casy-1 promoter (d). Maximum intensity projection images of L1/L2 larvae. Asterisks in c represent neurons with predominant RFP expression. Lateral and medial positions of the head region of a worm carrying the neuronal exon 11.5-skipping/inclusion reporter (d). The predominant RFP signals are observed in the amphid sensory neurons (dotted region) as well as in other neuron classes, including BAG sensory neurons, RIC and RIG interneurons (d). The predominant GFP signals are observed in neurons, including IL2 sensory neurons and AVG interneurons (d). BF, bright field. The utilized worm strains were JN1737 (b), JN1736 (c) and JN785 (d), whose genotypes are shown in Supplementary Table 3. Scale bars, 50 \( \mu \)m (b, whole body); 30 \( \mu \)m (c); 10 \( \mu \)m (b, mid-body region and tail region, and d).

Multiple RBPs regulate exon 11.5 selection. To understand the mechanism underlying neuron-class-specific inclusion of daf-2 exon 11.5, we investigated the effects of mutations of genes whose mammalian orthologues were known to encode alternative splicing factors in the brain\textsuperscript{3} (Supplementary Table 1). We compared the fluorescence signals between wild-type and mutant worms expressing the exon 11.5-skipping/inclusion reporters in a variety of neurons, including all the amphid sensory neurons and some inter- and motor-neurons, under the casy-1 promoter\textsuperscript{21} (hereafter this transgene is referred to as ‘neuronal exon 11.5-skipping/inclusion
Error bars represent s.e.m. among experiments using three biological replicates, and P values were determined by two-tailed t-test (d-f). The utilized worm strains were JN1709 (b,d,f) and JN1710 (c,d,f), whose genotypes are shown in Supplementary Table 3.

Figure 2 | Neuron-class-specific exon 11.5 selection of endogenous daf-2. (a) Schematic of DNA constructs in transgenes used for cell-type-selective expression of FLAG-tagged poly(A)-binding protein (FLAG-PAB-1). An intergenic region between flag::pab-1 and GFP contains an acceptor site of spliced leader (SL2) trans-splicing. The minigenes produce two mature mRNAs from a polycistronic pre-mRNA by trans-splicing, and GFP is co-expressed with FLAG-PAB-1 as a marker. (b,c) GFP expression patterns of a worm expressing the flag::pab-1::SL2::GFP minigene under the gpc-1 (b) or the glr-1 (c) promoter, respectively. The gpc-1 promoter drives expression in 10 neuron classes, including 7 amphid sensory neurons (ADL, ASI, ASE, AFD, AWC, ASH and ASJ). The head and tail regions of L1 larvae are shown. Scale bars, 10 μm. (d,e) Enrichment levels of odr-3 and glr-2 mRNAs (d), and total daf-2 and daf-2 (exon 11.5+) mRNAs (e) relative to a ubiquitous gene, eef-1A.1, mRNA in gpc-1- and glr-1-expressing neurons. Relative mRNA levels were quantified by quantitative RT–PCR, and enrichment levels were calculated as ratios of the mRNA levels in neuron-type-specific poly(A)⁺ RNAs to those in total RNAs from whole worms. odr-3 is exclusively expressed in some amphid sensory neurons; glr-2 is co-expressed with glr-1 in many neurons; glr-1 mRNA to the total. Error bars represent s.e.m. among experiments using three biological replicates, and P values were determined by two-tailed t-test (d-f). The utilized worm strains were JN1709 (b,d,f) and JN1710 (c,d,f), whose genotypes are shown in Supplementary Table 3.

reporter; Fig. 1d). The fluorescence signals in the head ganglia were evaluated based on intensity ratios of RFP to GFP and fractions of GFP and RFP expression areas (Fig. 3a). Some of the mutants showed significant changes in the reporter expression compared with the wild-type worms. E11.5(+)::RFP (exon 11.5 inclusion) signals were significantly reduced in the mutants of a CELF family gene unc-75, a PTB protein orthologue pth-1, an RBFOX family gene asd-1 and an hnRNP A1 homologue hrp-1 (Fig. 3b,c). On the contrary, E11.5(+)::RFP signals were increased in the mutants of an hnRNP F/H homologue hrp-1, a Hu/ELAV family gene exc-7 and a TRA2β orthologue rsp-8 (Fig. 3b,c). These data indicated that a number of evolutionarily conserved alternative splicing factors were involved in precise control of alternative splicing of exon 11.5 in the nervous system. Except for a ubiquitously expressed hrp-1 gene, these genes were reported to be expressed in restricted cell types, raising the possibility that neuron-class-specific alternative splicing of exon 11.5 is ensured by a combinatorial action of the splicing factors. To explore this idea, we further investigated the roles of the RBFOX
Figure 3 | Multiple RBPs are required for exon 11.5 selection. (a) A flow chart for quantification of the neuronal exon 11.5-skipping/inclusion reporter expression. The top right graph shows the averaged RFP-to-GFP intensity ratio in the wild-type reporter worms with s.e.m. (n = 514 worms). The bottom right graph shows the averaged fractions of GFP- and/or RFP-positive pixels in the wild-type reporter worms with s.e.m. (n = 459 worms). See details in the Methods section. (b, c) Analysis of alternative splicing patterns of the neuronal exon 11.5-skipping/inclusion reporter in the mutant backgrounds of RBPs. Normalized fluorescence intensity ratios of RFP (exon inclusion) to GFP (exon skipping) (b) and fractions of GFP and RFP expression (c) in the head ganglia are shown. Data were normalized by the averaged values in the wild-type worms (n ≥ 25; b). The n values are shown in each bar. Error bars represent s.e.m. *P < 0.05; **P < 0.01, different from wild type, two-tailed t-test with Bonferroni correction.
genes \( \text{asd-1} \) and \( \text{fox-1} \), \( \text{unc-75} \) and \( \text{ptb-1} \), which are required for the inclusion of exon 11.5 in the nervous system.

**RBFOX proteins directly promote exon 11.5 inclusion.** Exon inclusion of the neuronal exon 11.5-skipping/inclusion reporter was reduced in two \( \text{asd-1} \) mutants (Fig. 4a,b). A mutation in the paralogous gene \( \text{fox-1} \) enhanced the inclusion defect of \( \text{asd-1} \) mutants, and neurons with predominant exon 11.5 inclusion were severely reduced in the \( \text{asd-1} \); \( \text{fox-1} \) double mutants (Fig. 4a–c; Supplementary Fig. 3a,b). Together with their common property

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**Figure 4 | RBFOX family proteins directly promote exon 11.5 inclusion.** (a,b) Normalized fluorescence intensity ratios of RFP to GFP (a) and fractions of GFP and RFP expression (b) of the neuronal exon 11.5-skipping/inclusion reporter in the mutants of the RBFOX family genes. Data were normalized by the averaged values in the wild-type worms \((n≥22)\). The same data as in Fig. 3b,c are presented for \( \text{fox-1} \)\((e2643)\) and \( \text{asd-1} \)\((ok2299)\). (c) Maximum intensity projection images of the head region of a worm carrying the neuronal exon 11.5-skipping/inclusion reporter in the \( \text{asd-1} \)\((ok2299) \); \( \text{fox-1} \)\((e2643)\) background. (d) Schematic of modified \( \text{daf-2} \) exon 11.5-inclusion reporters. An intergenic region between RFP and GFP contains an acceptor site of SL2 trans-splicing, by which a polycistronic pre-mRNA is converted into monocistronic \( \text{daf-2} \) mini-gene-fused RFP mRNA and GFP mRNA. The RFP fusion protein is expressed upon exon 11.5 inclusion. Sites and sequences of the mutations \( \text{M1} \) and \( \text{M2} \) are indicated with lower case above the consensus RBFOX target stretches in red. (e) Expression patterns of the wild-type (control) and mutant form \((\text{M1}/\text{M2})\) of the modified exon 11.5-inclusion reporter driven by the \( \text{gpc-1} \) promoter. (f) Normalized fluorescence intensity ratios of RFP (exon inclusion) to GFP (promoter activity) in the amphid sensory neurons. Data were normalized by the averaged values in the wild-type worms \((n≥20)\). The \( n \) values are shown in each bar (a,b,f). Error bars represent s.e.m. **P < 0.01, different from wild-type (a) and control (f) worms; #P < 0.05, different from each other, two-tailed t-test with Bonferroni correction. L1/L2 larvae were used for the analyses. Scale bars, 10 μm.
as the RBFOX family splicing factor23, these data suggest functional redundancy of these genes for exon 11.5 inclusion. These RBFOX family proteins have been shown to regulate splicing through an evolutionarily conserved binding motif UGCAUG23,24. We found that the flanking intron downstream of daf-2 exon 11.5 contains two putative RBFOX target stretches (Supplementary Fig. 1a). To test the involvement of these stretches in exon 11.5 inclusion, we created modified versions of the E11.5(+)::RFP reporter to monitor exon-inclusion levels relative to the reporter expression levels (Fig. 4d). Mutations in either of the UGCAUG stretches caused a significant reduction, and a double mutation of these stretches caused a strong
Bonferroni correction. L1/L2 larvae were used for the analyses.

proteins for neuron-class-specific alternative splicing of the next examined requirements of UNC-75, PTB-1 and the RBFOX proteins. Whole worms revealed that inclusion of daf-2 (exon 11.5) mutant worms for the analyses (Fig. 6b,c). Amount of the scripts in the amphid sensory neurons from the wild-type and the method. We used RNA fractions comparably enriched for transcripts in the amphid sensory neurons by the mRNA-tagging method. We used RNA fractions comparably enriched for transcripts in the amphid sensory neurons from the wild-type and the mutant worms for the analyses (Fig. 6a,c). Amount of the daf-2 (exon 11.5 +) isoform relative to the daf-2 (exon 11.5 −) isoform was significantly reduced in the amphid sensory neurons of the unc-75, ptb-1 and asd-1 single mutants, and a ptb-1; asd-1 double mutant (Fig. 6d). These results confirmed that these splicing factors promote inclusion of exon 11.5 of endogenous daf-2 in the amphid sensory neurons.

Abnormal splicing of endogenous daf-2 in RBP mutants. We next examined requirements of UNC-75, PTB-1 and the RBFOX proteins for neuron-class-specific alternative splicing of endogenous daf-2. RT–PCR analysis using total RNAs from whole worms revealed that inclusion of daf-2 exon 11.5 was significantly reduced in the splicing factor mutants compared with the wild type (Fig. 6a). We then analysed daf-2 isoform expression in the amphid sensory neurons by the mRNA-tagging method. We used RNA fractions comparably enriched for transcripts in the amphid sensory neurons from the wild-type and the mutant worms for the analyses (Fig. 6a,c). Amount of the daf-2 (exon 11.5 +) isoform relative to the daf-2 (exon 11.5 −) isoform was significantly reduced in the amphid sensory neurons of the unc-75, ptb-1 and asd-1 single mutants, and a ptb-1; asd-1 double mutant (Fig. 6d). These results confirmed that these splicing factors promote inclusion of exon 11.5 of endogenous daf-2 in the amphid sensory neurons.

The RBFOX genes are expressed in various neuron classes. To test the hypothesis that neuron-class-specific inclusion of exon 11.5 is achieved by combinatorial expression of the splicing factors, we examined expression patterns of the RBFOX genes (asd-1 and fox-1), unc-75 and ptb-1. As previously reported,23, the RBFOX genes were expressed in various tissues including the pharynx, the intestine, hypodermis, neurons, muscles and the somatic gonad (Supplementary Fig. 6a–g). As the neuronal asd-1 expression was difficult to observe because of its expression in various tissues in the head region, we utilized split GFP26 to observe neuronal expression pattern of the RBFOX genes (Fig. 7a). One of the two split-GFP fragments, spGFP1-10, which contains 10 of the 11 strands of the β-barrel structure of GFP, was expressed under the asd-1 and the fox-1 promoters. mCherry-fused spGFP11, another split-GFP fragment that contains the 11th strand of the GFP β-barrel, was expressed by the pan-neuronal H2O promoter. GFP signals, which reflect co-expression of spGFP1-10 and spGFP11:mCherry (Fig. 7a), were observed in many but not all neuron classes, suggesting that the expression of the RBFOX genes covers a variety of neuron classes (Fig. 7b).

ptb-1 is expressed in subsets of tissues. Large-scale expression analyses suggested that ptb-1 was expressed in a subset of neurons as well as non-neuronal cells27,28. We now precisely determined the expression pattern of ptb-1. We first confirmed expression of two ptb-1 variants, ptb-1a and ptb-1b, driven by alternative promoters (Fig. 8a), as annotated in WormBase (http://www.wormbase.org). PTB-1a has four evolutionarily conserved RNA recognition motifs (RRM1 – 4)29, while PTB-1b lacks the first RRM (RRM1; Fig. 8a).

We also found a 34-nucleotide (nt) cassette exon, exon 11 (Fig. 8a,b), skipping of which causes a frameshift and a premature termination codon. RT–PCR analysis of the PTB-1 isoforms in the nonsense-mediated mRNA decay (NMD)-deficient smg-2 mutant indicated that both ptb-1a and ptb-1b have the exon 11 (−) isoforms destabilized by NMD (Fig. 8b). The exon 11 (−) ptb-1 variants were not detected in the ptb-1 mutant (Fig. 8c), indicating that PTB-1 proteins repress splicing of exon 11 in their own transcripts to produce unproductive mRNA isoforms. The overall domain structure of PTB-1a and the mechanism of negative autoregulation of its own expression level via alternative splicing of the 34-nt cassette exon encoding a portion of the third RRM coupled with the NMD pathway are quite similar to those of its mammalian orthologues PTBP1 and PTBP2 (refs 30,31), suggesting their conserved functions as alternative splicing regulators.

We next investigated the expression patterns of transcriptional reporter genes containing the upstream regulatory elements for either of the first exons (Fig. 8a). The ptb-1a promoter drove the expression in a small subset of neurons, including all the amphid sensory neurons, muscles and somatic gonadal cells, while the ptb-1b promoter drove strong expression in neurons (Fig. 8d–f). In the nervous system, the ptb-1a and ptb-1b promoters drove the expression in distinct but partially overlapping neuron classes, including ASE (Fig. 8d).

Neural PTB-1 expression correlates with exon 11.5 inclusion. We then examined cell types that express ptb-1 along with the RBFOX genes using split GFP (Fig. 7a). spGFP1-10 and
spGFP11::mCherry were expressed under the two ptb-1 (ptb-1a and ptb-1b) promoters and the two RBFOX gene (asd-1 and fox-1) promoters, respectively. GFP signals, which reflect co-expression of ptb-1 and either of the RBFOX genes, were observed in subsets of neuronal, muscle and somatic gonadal cells (Fig. 7c). This expression pattern was highly similar to that of the ptb-1 transcriptional reporter genes (Fig. 8d–f), suggesting that RBFOX was expressed in most ptb-1-expressing cell types. Pan-neuronal expression of unc-75 (ref. 32) further restricted cell types that co-express the RBFOX, unc-75 and ptb-1 genes to a subset of neurons (Fig. 7d; Supplementary Fig. 6h).

A series of expression analyses raised the possibility that the ptb-1-expressing neurons (that is, co-expression sites of the RBFOX, unc-75 and ptb-1 genes) might correlate with predominant inclusion of daf-2 exon 11.5, which requires combinatorial action of these splicing factors. Therefore, we compared expression patterns of the ptb-1 transcriptional reporters with that of the daf-2 exon 11.5-inclusion or -skipping reporter in the nervous system. All neurons that expressed ptb-1a::Venus and/or ptb-1b::Venus also expressed E11.5(+)::RFP (Fig. 8g). In contrast, expression of Venus and E11.5(−)::RFP was mutually exclusive in many neuron classes (Fig. 8h). When the E11.5(+):RFP and E11.5(−):GFP reporters were co-expressed under the ptb-1a or ptb-1b promoter, the nervous system preferentially expressed E11.5(+):RFP, with a few exception such as in AVA and RIF neurons (Supplementary Fig. 2e,f), which further confirmed preferential inclusion of exon 11.5 in the PTB-1-expressing neurons. These analyses revealed strong correlation between PTB-1 expression and inclusion of daf-2 exon 11.5 in the nervous system.

PTB-1 confers neuron-type specificity on exon 11.5 selection. To test whether PTB-1 expression is sufficient for preferential inclusion of daf-2 exon 11.5 in the nervous system, either PTB-1a or PTB-1b was ectopically expressed in a variety of neurons of the ptb-1 mutant carrying the neuronal exon 11.5-skipping/inclusion reporter. Expression of either of the PTB-1 isoforms significantly
enhanced exon 11.5 inclusion in the nervous system of the ptb-1 mutant (Fig. 9a,b). The effect of ectopic PTB-1a expression was prominent compared with that of PTB-1b (Fig. 9b). Lack of the RRM1 domain in the PTB-1b isoform implies the requirement of RRM1 for efficient PTB-1 action. Most PTB-1a-expressing neurons, even the neurons where the E11.5(−)::GFP reporter was predominantly expressed in the wild type, such as IL2 sensory neurons (Fig. 1d), exhibited preferential expression of the E11.5(−)::RFP reporter, whereas neurons that failed to express PTB-1a did not (Fig. 9c). PTB-1b expression also caused preferential E11.5(−)::RFP expression in many neurons, although some neurons showed preferential E11.5(−)::GFP expression despite PTB-1b expression (Fig. 9d). These data indicated that the PTB-1 expression is a primary determinant of the neuron-class specificity of exon 11.5 inclusion in the nervous system.

**Impaired learning in RBP mutants.** We explored the behavioural consequences of defective inclusion of daf-2 exon 11.5 in the splicing factor mutants. *C. elegans* worms alter their preferences for NaCl concentrations after conditioning with different NaCl concentrations. They are attracted to NaCl concentrations at which they were fed, whereas they avoid NaCl concentrations at which they were starved (Fig. 10a)33. Insulin-like signalling mutants exhibit defects in salt concentration avoidance after starvation conditioning, which we call taste-avoidance learning20,33. As only the DAF-2c (exon 11.5+) isoform can mediate signalling for taste-avoidance learning20,33, we tested salt concentration learning of the splicing factor mutants. As the unc-75 null mutant exhibits a severe locomotion defect24, we examined reduction-of-function mutants of unc-75. The unc-75(yb1714) mutant showed defects in migration to higher salt concentrations (Supplementary Fig. 7), suggesting that unc-75 is required for proper behavioural responses to salt as well as for locomotion. ptb-1 and asd-1 mutants had no obvious defect in behavioural responses to salt after conditioning under fed conditions, whereas they showed defects in taste-avoidance learning (Fig. 10b; Supplementary Fig. 8a,b). The ptb-1; asd-1 double mutants showed further impaired taste-avoidance learning (Fig. 10b). These data indicated that ptb-1 and asd-1 are required for taste-avoidance learning.

To further characterize the behavioural phenotype of the splicing factor mutant, we tested for genetic interactions between the splicing factor genes and daf-16. DAF-2 signalling controls the action of the DAF-16/FOXO transcription factor, by inhibiting its nuclear translocation34. Most phenotypes of daf-2 mutants are suppressed by daf-16 mutations34, whereas we...
reported that daf-16 mutations strongly enhanced the taste-avoidance learning defect of daf-2, implying that, unlike the canonical DAF-2 signalling, the insulin-like signalling mediated by DAF-2c (exon 11.5 +) may regulate taste-avoidance learning in parallel with DAF-16 (ref. 19). daf-16; ptb-1; asd-1 triple mutants showed largely normal behavioural responses to salt after conditioning under fed conditions, whereas they showed a strong defect in taste-avoidance learning as if they were not starved during conditioning (Fig. 10c; Supplementary Fig. 8c,d). This behavioural phenotype resembled that of the daf-16; daf-2 mutant (Fig. 10c; Supplementary Fig. 8c,d)19. The similar genetic interactions of the daf-16 gene with ptb-1; asd-1 and daf-2 support the idea that the splicing factors regulate taste-avoidance learning through the production of DAF-2c (exon 11.5 +) isoform.

**DAF-2c expression in ASER relieves the impaired learning.** To assess impacts of altered expression of daf-2 isoforms on the taste-avoidance learning, we generated wild-type and mutant worms ectopically expressing either of the DAF-2 isoforms. Neuronal expression of either DAF-2a (exon 11.5 –) or DAF-2c...
In this study, we showed cell-type-specific alternative splicing of daf-2 exon 11.5. Within the nervous system, daf-2 exon 11.5 inclusion preferentially occurs in a restricted subset of neurons, including the amphid sensory neurons. This neuron-class-specific alternative splicing is determined by the combinatorial action of the evolutionarily conserved RBPs, RBFOX proteins (ASD-1 and FOX-1), UNC-75/CELF and PTB-1/PTB. We found that PTB-1, which is co-expressed with the RBFOX proteins and UNC-75 in the restricted neuron classes (Fig. 7d), has a key role in conferring neuron-class specificity of exon 11.5 inclusion. The combinatorial mutations of these splicing factors result in a pronounced defect in taste-avoidance learning. DAF-2c (exon 11.5+) expression only in a single neuron ASER relieves the learning defect of the splicing factor mutant. Therefore, the precisely regulated daf-2 (exon 11.5+) isoform in ASER.

### Discussion

In this study, we showed cell-type-specific alternative splicing of daf-2 exon 11.5. Within the nervous system, daf-2 exon 11.5 inclusion preferentially occurs in a restricted subset of neurons, including the amphid sensory neurons. This neuron-class-specific alternative splicing is determined by the combinatorial action of the evolutionarily conserved RBPs, RBFOX proteins (ASD-1 and FOX-1), UNC-75/CELF and PTB-1/PTB. We found that PTB-1, which is co-expressed with the RBFOX proteins and UNC-75 in the restricted neuron classes (Fig. 7d), has a key role in conferring neuron-class specificity of exon 11.5 inclusion. The combinatorial mutations of these splicing factors result in a pronounced defect in taste-avoidance learning. DAF-2c (exon 11.5+) expression only in a single neuron ASER relieves the learning defect of the splicing factor mutant. Therefore, the precisely regulated daf-2

(exon 11.5+) in the wild-type resulted in impaired learning (Fig. 10d), suggesting that appropriate expression levels of DAF-2a and DAF-2c or their balance in the nervous system is important for proper learning ability. On the other hand, neuronal expression of DAF-2c, but not that of DAF-2a, rescued the learning defect of the daf-2 mutant (Fig. 10d), indicating that the critical neuron for the splicing factors to control taste-avoidance learning by producing DAF-2c. The DAF-2c expression in ASER also rescued the learning defect of the daf-16; ptb-1; asd-1 mutant (Fig. 10e), suggesting that DAF-2c (exon 11.5+) signalling regulates taste-avoidance learning in a DAF-16-independent manner. Collectively, these

Figure 9 | Ectopic PTB-1 expression promotes exon 11.5 inclusion in the nervous system. (a,b) Normalized fluorescence intensity ratios of RFP to GFP (a) and fractions of GFP and RFP expression (b) of the neuronal exon 11.5-skipping/inclusion reporter in the ptb-1(gk347274) mutant and in those expressing either PTB-1a or PTB-1b under the casy-1 promoter. Data were normalized by the averaged values in the wild-type worms (n ≥ 33). (a) Error bars represent s.e.m. **P < 0.01, different from each other, +P < 0.01, comparison of RFP-positive fractions, two-tailed t-test with Bonferroni correction. (c,d) Maximum intensity projection expression images of the neuronal exon 11.5-skipping/inclusion reporter in the ptb-1(gk347274) mutant expressing PTB-1a (c) or PTB-1b (d) under the casy-1 promoter. CFP is co-expressed along with PTB-1 from a polycistronic pre-mRNA. As PTB-1 was expressed from extrachromosomal arrays (Supplementary Table 4), they exhibited mosaic expression patterns.
Figure 10 | DAF-2c expression in ASER relieves impaired learning of the RBP mutant. (a) A flow chart of salt concentration learning assay. For conditioning, worms were exposed to low or high NaCl concentrations (low [NaCl] or high [NaCl], respectively) under fed or starvation conditions (food (+) conditioning or food (−) conditioning, respectively). After conditioning, chemotaxis to NaCl was tested on an agar plate with a NaCl gradient, and then chemotaxis and learning indices were calculated. See details in the Methods section. Chemotaxis and learning indices of wild-type worms are shown (n = 21). (b) Salt concentration learning of the wild type, daf-2(pe1230), unc-75(yb1714), ptb-1(gk347274), asd-1(ok2299) and ptb-1(gk347274); asd-1(ok2299) mutants. (c) Salt concentration learning of the wild type, daf-16(mgDf50), daf-16(mgDf50); daf-2(e1370) and daf-16(mgDf50); ptb-1(gk347274); asd-1(ok2299). (d) Salt concentration learning of wild-type, daf-2(pe1230) and ptb-1(gk347274); asd-1(ok2299) worms expressing none (−) or either of the DAF-2 isoforms under the H20 promoter. Salt conditioning was performed under starvation conditions. (e) Salt concentration learning of wild-type, ptb-1(gk347274); asd-1(ok2299) and daf-16(mgDf50); ptb-1(gk347274); asd-1(ok2299) worms expressing none (−) or DAF-2c only in ASER under the gcy-5 promoter. Salt conditioning was performed under starvation conditions. n ≥ 4 assays (b, c), n ≥ 6 assays (d, e). Error bars represent s.e.m. *P < 0.05; **P < 0.01, different from wild-type or control worms, one-way analysis of variance (ANOVA) followed by Dunnett post hoc test (b–d) or two-tailed t-test with Bonferroni correction (e).
exon 11.5 inclusion in ASER is essential for taste-avoidance learning (Supplementary Fig. 9). DAF-2 signalling is also required for starvation-induced odour chemotaxis learning in AWC olfactory neurons\(^{23,35}\), although a DAF-2 isoform that regulates olfactory learning has not been determined. The predominant inclusion of exon 11.5 in the amphid sensory neurons, including AWC (Supplementary Fig. 2a), implies a requirement of the DAF-2c isoform in a range of sensory plasticity, including those of taste and smell responses.

PTB functions as both a repressor and an activator in many alternative splicing events in mammalian cells\(^{31,36}\). The neuronal PTB knockout mice show severe defects in neurogenesis and result in death soon after birth\(^{8,12}\). In this study, we unveiled the role of C. elegans PTB-1 in determination of the neuronal property essential for behavioural learning. The expression pattern and the ectopic expression analyses suggest that PTB-1 is a strong determinant of daf-2 exon 11.5 inclusion in the nervous system. We characterized the two PTB-1 isoforms, PTB-1a and PTB-1b, which are produced by alternative promoters. PTB-1a strongly activates exon 11.5 inclusion compared with PTB-1b. Mammalian PTB proteins, PTBP1 and PTBP2, also have different effects on alternative splicing. It was reported that a switch between expressions of PTBP1 and its paralogue PTBP2 is required for reprogramming of protein isoforms during neuronal maturation\(^{12,31}\). This switch in expression is due in part to PTBP1-induced alternative splicing of a 34-nt cassette exon of PTBP2, which leads to mRNA degradation by NMD. We found the cassette exon (exon 11) in the ptb-1 gene, skipping of which leads to NMD-dependent mRNA degradation. It is surprising that its role in negative autoregulation as well as its position and size is highly conserved between mammals and C. elegans. The two PTB-1 isoforms are expressed in distinct but partially overlapping neurons, including ASER. It would be interesting to investigate how PTB-1 isoform expressions are controlled at transcription and post-transcription to determine the neuronal properties essential for learning ability.

Except for daf-2 exon 11.5 and ptb-1 exon 11, we identified in this study, a target exon of PTB-1 has not been reported. The RBFOX family proteins regulate several alternative splicing events in various tissues, including the muscles and the nervous system\(^{23,24}\). UNC-75 is exclusively expressed in the nervous system and determines protein isoforms of various neuronal genes\(^{16,37}\). Each of these splicing factors might control a different subset of neuronal genes. Because DAF-2c (exon 11.5+) expression only partially restores the taste-avoidance learning defect of the splicing factor mutant, other genes related to behavioural learning may be affected in the mutant. It would be interesting to identify exons cooperatively regulated by these splicing factors, which may be involved in learning ability. And this would provide further insights into how the neuron-class-specific alternative splicing contributes to the complex neuronal functions underlying learning.

In addition to these splicing factors, we showed that mutations in exc-7 and rep-8 increase expression of the neuronal exon 11.5-inclusion reporter. EXC-7/Hu/ELAV and RSP-8/TRAF2 might repress the exon 11.5 inclusion and further fine-tune the neuron-type specificity of alternative splicing of daf-2. Further investigation is needed to clarify the functional role of the candidate repressors of exon 11.5 inclusion in the nervous system. On the other hand, the hpr-1 mutation reduces expression of the neuronal exon 11.5-inclusion reporter. HRP-1/hmRNP A1 is ubiquitously expressed and essential for larval development\(^{22,39}\). RNA interference knockdown of hpr-1 during the adult stage extends lifespan in a daf-16-dependent manner similar to that of daf-2 (ref. \(^{39}\)). It will be interesting to determine whether and how HRP-1 contributes to selection of daf-2 exon 11.5 and longevity extension.

We demonstrated that the RBFOX family proteins directly act through the binding motif UGCAUG in the intron downstream of exon 11.5. We identified a cassette exon homologous to daf-2 exon 11.5 in Cbr-daf-2, the daf-2 orthologue of the closely related nematode species C. briggsae\(^{20}\). Cbr-daf-2 also has an RBFOX-binding motif near the 5′ splice site in the intron downstream of the cassette exon (Supplementary Fig. 1b). Exon 11.5 of daf-2 resides in a similar position to a cassette exon, exon 11, of mammalian insulin receptor (IR) genes\(^{30}\). The RBFOX-binding motif also resides near the 5′ splice site in the intron downstream of exon 11 of the human and mouse IR genes. It was predicted that inclusion of Insr exon 11 is directly promoted by RBfox proteins in the mouse brain\(^{3}\). The conserved regulation by the RBFOX family proteins in the nematodes and mammals suggests the biological importance of RBFOX-regulated alternative splicing of insulin receptors in the nervous systems.

Mammalian CELF family proteins have been shown to repress exon 11 of the IR genes\(^{30,41}\). These CELF proteins are widely expressed in numerous tissues, whereas alternative splicing of exon 11 of the IR genes occurs in a tissue-specific manner\(^{22}\). It was postulated that the coordination of CELF proteins and other factors, such as muscleblind-like 1, confer tissue-type specificity to the alternative splicing regulation of IR\(^{43}\). All six human CELF family genes are strongly expressed in the nervous system\(^{6}\). The CELF family proteins regulate cell-type-specific alternative splicing in concert with other splicing factors, including PTB\(^{44,45}\). These reports raise the possibility that the CELF family proteins and other splicing factors, such as RBFOX and PTB, might cooperatively control neuron-type-specific alternative splicing of the mammalian IR genes in a manner analogous to that in the C. elegans nervous system. Dysfunction of neuronal IR signalling is linked to neurological diseases such as Alzheimer’s disease\(^{46}\). However, an isoform-specific function of IR in the brain is still unclear. It will be interesting to investigate alternative splicing patterns of the IR genes in distinct neuron types and an isoform-specific function of IR in the brain.

**Methods**

**C. elegans strains.** C. elegans Bristol strain N2 was used as the wild type. The strains were grown and maintained at 20°C. Standard genetics methods were used to generate multiple mutants by crosses\(^{47}\). hpr-1(ok9635) IV and ptb-1(kk347274) II; hpr-1(ok9635) IV and ptb-1(kk347274) II; asd-1(ok2299) III, fex-1(c2645) X were maintained with the genetic balancers, C176 (htr-1[ts]/[ts]) and q3C1[ok2260] I, respectively. daf-2(pe1230) was previously isolated by forward genetic screening for mutants defective in taste-avoidance learning\(^{22}\). No daf-2(pe1230) worm shows a dauer constitutive (Daf-c) phenotype at 20°C. The mutation site of pe1230 is shown in Supplementary Table 1. The mutants and transgenic worms used in this study are listed in Supplementary Tables 2–4.

**DNA constructs and transgenesis.** To generate daf-2 cDNA expression plasmids, a PCR-amplified H20 promoter (2.5 kb) was inserted upstream of daf-2/c cDNAs in pBS-daf-2a/c plasmids\(^{26}\). Plasmids except for daf-2 cDNA constructs were generated using the Gateway system (Invitrogen) as previously described\(^{26}\). To
generate daf-2 exon 11.5-splicing reporter constructs, a PCR-amplified genomic region of daf-2 between parts of exons 11 and 12 was inserted upstream of mRFP, EYFP or EGFP fused mRFP separated by a sequence with an acceptor site of a spliced leader (SL2), in a Gateway destination vector, pPD-DEST2. Mutations were introduced into exon 11.5, exon 12 and/or RBFOX-binding motifs by PCR reactions. Sequences of the daf-2 genomic region and mutation sites in the splicing reporters are shown in Supplementary Fig. 1a. To generate Flag-tagged expression constructs, Flag-tagged cDNA was inserted upstream of an acceptor site of SL2 followed by GFP in pPD-DEST2. To generate plb-1 cDNA expression constructs, PCR-amplified plb-1a and plb-1b cDNAs were inserted upstream of an acceptor site of SL2 followed by GFP in pPD-DEST2. To generate a spGP11::mCherry expression construct, spGP11 cDNA was inserted in-frame upstream of mCherry cDNA in pPD-DEST2. spGP11 and spGP11 cDNAs are a generous gift from Dr Cori Bargmann. Gateway entry vectors with promoters for H2O (ref. 21), casy-1 (ref. 20), gpc-1 (ref. 48), gr-1 (ref. 19) and odr-2 (ref. 19) were previously described. PCR-amplified asd-1 (2.3 kb), ptb-1a (5.1 kb), ptb-1b (7.4 kb) and plb-1 (5.3 kb) cDNA were inserted downstream of a Gateway entry vector. Pfox-1::spGP11::mCherry and Pfox-1::spGP11::mCherry were expressed as a vector-free construct generated by a PCR fusion-based method49. A 4.0-kb upstream regulatory sequence of fox-1 was fused to spGP11::mCherry followed by an untranslated region using PCR fusion reactions. Germine transformations were performed using standard microinjection methods50. DNA constructs were injected at concentrations between 20 and 100 ng μl−1 and with or without a co-injection marker, unc-122p::mCherry (20 ng μl−1), PR4a carrying rol-6(su100) (50 ng μl−1) or myo-3p::Venus (10 or 30 ng μl−1), and a carrier DNA, pPD49.26. Injection mixtures were prepared at a final concentration of 100 ng μl−1.

**Cloning of plb-1 and Cbr-daf-2 cDNAs.** Full-length plb-1 and partial Cbr-daf-2 cDNAs were amplified by RT-PCR from C. elegans and C. briggsae total RNAs, respectively. The plb-1 and Cbr-daf-2 cDNAs were cloned and sequenced to examine the genome structures. The plb-1a and plb-1b cDNA sequences were submitted to DNA Data Bank of Japan and were given accession numbers, LC026472 and LC026473, respectively. Primers used for PCR reactions are listed in Supplementary Table 5.

**mRNA tagging.** Neuron-type-specific poly(A) + RNA was isolated from transgenic adult worms with FLAG-PAB-1 expressed under a neuron-type-selective promoter according to our previously published protocol51 with some modifications. We used transgenic worms expressing FLAG-PAB-1 and GFP under the gpc-1 or gr-1 promoter (Fig. 2a–c, Supplementary Tables 3 and 4). To co-link poly(A) + RNA with FLAG-PAB-1, the worms grown on nematode growth medium (NGM) plates were collected and washed three times with M9 buffer, and then treated with 5% formaldehyde in M9 for 20 min at room temperature. To immunoprecipitate the RNA/FLAG-PAB-1 complexes, the supernatant of worm lysates was incubated with anti-FLAG M2 affinity gel beads (Sigma-Aldrich) for 4 h at 4 °C. The precipitated materials including RNA/FLAG-PAB-1 complexes were incubated for 30 min at 65 °C to remove the crosslink.

**RT-PCR and quantitative RT-PCR assays.** Neuron-type-specific RNA and total RNA were reverse-transcribed, and then subjected to PCR reactions with gene- or exon-specific primers to compare transcript levels between samples. The RT-PCR and quantitative RT-PCR assays were performed according to previously reported procedures33 with some modifications. For conditioning, adult worms were transferred to NGM plates with 25 or 100 mM NaCl for 5 h. After conditioning, the worms were placed at the centre of a test plate with a NaCl gradient and were allowed to crawl for 45 min. The RT-PCR and learning indices were determined as shown in Fig. 10a. A learning index after conditioning under fed conditions was determined by subtracting the chemotaxis index after conditioning on an NGM plate with 25 mM NaCl from that with 100 mM NaCl. A learning index after conditioning under starvation conditions was determined by subtracting the chemotaxis index after conditioning on an NGM plate with 100 mM NaCl from that with 25 mM NaCl.

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