An ensemble of regulatory elements controls Runx3 spatiotemporal expression in subsets of dorsal root ganglia proprioceptive neurons

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The Runx3 transcription factor is essential for development and diversification of the dorsal root ganglia (DRGs) TrkC sensory neurons. In Runx3-deficient mice, developing TrkC neurons fail to extend central and peripheral afferents, leading to cell death and disruption of the stretch reflex circuit, resulting in severe limb ataxia. Despite its central role, the mechanisms underlying the spatiotemporal expression specificities of Runx3 in TrkC neurons were largely unknown. Here we first defined the genomic transcription unit encompassing regulatory elements (REs) that mediate the tissue-specific expression of Runx3. Using transgenic mice expressing BAC reporters spanning the Runx3 locus, we discovered three REs—dubbed R1, R2, and R3—that cross-talk with promoter-2 (P2) to drive TrkC neuron-specific Runx3 transcription. Deletion of single or multiple elements either in the BAC transgenics or by CRISPR/Cas9-mediated endogenous ablation established the REs’ ability to promote and/or repress Runx3 expression in developing sensory neurons. Our analysis reveals that an intricate combinatorial interplay among the three REs governs Runx3 expression in distinct subtypes of TrkC neurons while concomitantly extinguishing its expression in non-TrkC neurons. These findings provide insights into the mechanism regulating cell type-specific expression and subtype diversification of TrkC neurons in developing DRGs.

[Keywords: Runx3 transcription factor; DRG TrkC proprioceptive neurons; long-range transcription regulatory elements; BAC transgenic mice; CRISPR/Cas9 enhancer mutants]

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Runx3 is a member of the mammalian RUNX family of transcription factors (TFs), which are key gene expression regulators in several important developmental processes (Levanon and Groner 2004). During embryonic development, Runx3 expression is first detected at around embryonic day 11 (E11) in the dorsal root ganglia (DRGs) and at later stages in developing bones, whiskers, hair follicles, and hematopoietic cells (Levanon et al. 2001, 2011). Loss of Runx3 in these cell types impaired their function, leading to phenotypic defects (Inoue et al. 2002; Levanon et al. 2002, 2014; Yamashiro et al. 2002; Woolf et al. 2003; Brenner et al. 2004; Fainaru et al. 2004; Raveh et al. 2005; Djuretic et al. 2007; Cruz-Guilloty et al. 2009; Naito and Taniuchi 2010; Dicken et al. 2013; Lotem et al. 2013; Bauer et al. 2015).

The DRGs include three main subclasses of sensory neurons distinguishable by their neurotrophin receptors: the nociceptive TrkA neurons, the mecanoeptive TrkB neurons, and the proprioceptive TrkC neurons. RUNX TFs play key roles in the post-mitotic diversification of these neurons into distinct sensory modalities (Lallemand and Ernfors 2012). Runx1 and Runx3 are differentially expressed in TrkA and TrkC neurons, respectively (Levanon et al. 2001, 2002; Inoue et al. 2002; Chen et al. 2006; Kramer et al. 2006; Nakamura et al. 2008). Interestingly, Runx3, the phylogenetically most
ancient mammalian \textit{RUNX} [Bangsow et al. 2001; Levanon et al. 2003], regulates the neurogenesis of TrkC neurons [Inoue et al. 2002; Levanon et al. 2002; Chen et al. 2006a; Kramer et al. 2006] that are a major constituent of the simplest and most ancient neuronal circuit: the stretch reflex arc [Levanon et al. 2003; Sullivan et al. 2008]. In the absence of Runx3, TrkC neurons are initially formed but fail to extend peripheral afferents and undergo apoptosis, leading to congenital ataxia [Levanon et al. 2002]. The strict specificity to TrkC neurons implies that Runx3 expression is tightly regulated. However, little was known about the molecular mechanisms regulating the spatiotemporal expression of \textit{Runx3} in developing TrkC neurons. Here, we used reporter BAC transgenics and CRISPR/Cas9-mediated gene editing to demonstrate that TrkC neuron-specific Runx3 expression is tightly regulated. However, little was known about the molecular mechanisms regulating the spatiotemporal expression of \textit{Runx3} in developing TrkC neurons. We found that the overall expression pattern of the six BACs, only those that extend 5’ upstream of \textit{Runx3} [namely, BAC-E, BAC-C, and, to a lesser extent, BAC-A] conferred LacZ expression in BAC-B (boxed in Fig. 1A) contains DRG REs. Next, to identify conserved REs, we conducted a multiple species alignment using mVista [Frazer et al. 2004]. The genomic sequences surrounding mouse \textit{Runx3} were compared with sequences from the orthologous regions in five vertebrates: humans, rats, chickens, frogs, and pufferfish [i.e., fugu] [Fig. 1A, bottom panel]. Frogs and fish were included because Runx3 function in the sensory stretch reflex arc is phylogenetically ancient [Levanon et al. 2003; Sullivan et al. 2008], and therefore the regulatory sequences controlling its expression in TrkC neurons have likely been conserved over the course of evolution. The Vista analysis revealed 10 highly conserved noncoding elements [CNEs] within and surrounding \textit{Runx3} [Supplemental Table S2]. Of these elements, four CNEs, marked R1–R4, were located in the presumable DRG regulatory region.

\textbf{Conserved elements R1, R2, and R3 are essential for Runx3 expression in TrkC neurons}

BAC-C-GFP that contains R1, R2, and R3 conferred reporter expression to most of the Runx3-positive cells along the DRGs and was therefore selected for further studies focusing on brachial DRGs [Fig. 1D, left]. Deletion of these REs, yielding BAC-C-delR1,2,3-GFP mice [Supplemental Table S3], abolished reporter expression in developing TrkC neurons as early as E11.5 [Fig. 1D, right]. This result indicates that these three REs are required for Runx3 expression in TrkC neurons. The essential role of these REs was further verified in mice carrying homozygous CRISPR/Cas9-mediated [CR] deletion [Δ] of the 25-kb genomic region spanning R1–R3 [CRΔ1,2,3;CRΔ1,2,3] [Supplemental Fig. S3; Supplemental Table S4], which developed severe ataxia, similar to Runx3\textsuperscript{−/−} mice [Levanon et al. 2002]. In the compound heterozygous mice CRΔ1,2,3/P2\textsuperscript{GFP}
Figure 1. The Runx3 transcriptional unit: gene structure, REs, and DRG expression. (A, top panels) Schematic presentation of six BAC reporters marked as A, C, and E [green bars] and B, D, and F [red bars] (chromosome 4: 134,953,991–135,328,237; University of California at Santa Cruz [UCSC], mm10) spanning the Runx3 transcription unit. The blue box below the BACs represents the LacZ/GFP reporter inserted into the Runx3 coding region (see also Supplemental Table S1, Supplemental Fig. S1). (bottom panel) Vista comparative analysis demonstrating the evolutionary conservation of the Runx3 transcriptional unit. The four REs—R1, R2, R3, and R4—are highlighted. (B) Runx3-P2 drives expression in developing DRG neurons. Immunofluorescence with anti-Runx3, anti-TrkC, and anti-GFP in E11 brachial DRGs of P2GFP/+ (top left) and P1AFP/+ (top right). Expression of Runx3 and TrkC in P2GFP/GFP at E11 (middle left) and E14.5 (middle right). DRG is marked by a dashed line. (Middle right) Yellow dots represent blood cells. Runx3 expression in TrkC neurons of P1AFP/AFP at E11 (bottom left) and E14.5 (bottom right). (C) Expression of LacZ in BAC-C and BAC-A transgenics and Runx3LacZ/+ embryos at E14.5 (shown is the embryos’ inner side). Boxed regions are enlarged in the bottom panels, demonstrating staining in DRGs. (D) BAC-C spans the three REs: R1, R2, and R3 (shown schematically above the top panels). GFP coexpression with TrkC and endogenous Runx3 in brachial DRG neurons at E11.5 (top left) and E12.5 (bottom left). Deletion of BAC-C REs (R1, R2, and R3) suspends GFP expression in TrkC/Runx3 neurons at E11.5 (top right) and E12.5 (bottom right). (E) CRISPR/Cas9-mediated deletion of the endogenous 25-kb region spanning R1, R2, and R3 (shown schematically above the panels). Expression of Runx3 and TrkC in ΔR1,2,3/P2GFP E11.5 embryos (top left) and same panel stained also with anti-GFP antibodies (bottom left). GFP derived from the P2-GFP allele serves as an internal control for TrkC neurons that lost Runx3. Expression of TrkC, Runx3, and GFP at E11.5 (top middle) and E14.5 (bottom middle). Expression of TrkB, Runx3, and GFP at E12.5 (top right) and E14.5 (bottom right). Note the expression of Runx3 at E14.5 in rib cartilage but not in DRGs. Bar, 50 μm.
Runx3 expression in TrkC/GFP neurons was markedly lower at E11.5 (Fig. 1E, left). At E12.5, this Runx3 reduction was accompanied by a pronounced decrease in the number of TrkC neurons (Fig. 1E, top middle), with only a few TrkC/GFP neurons detected at E14.5 (Fig. 1E, bottom middle).

The loss of Runx3 in TrkC neurons was associated with a gain in TrkB expression. Consequently, at E12.5, the number of TrkB-expressing neurons in CRAR1,2,3/P2GFP DRGs increased and consisted of two populations (Fig. 1E, top right). One population represented the conventional TrkB neurons, and the second, an unusual one, coexpressed TrkB and GFP. This latter population constituted TrkC neurons in which the loss of Runx3 led to TrkB expression. These observations correspond with previous findings that Runx3 represses TrkB and promotes TrkC expression (Kramer et al. 2006; Inoue et al. 2007). The nearly complete loss of TrkB expression in E14.5 in CRAR1,2,3/P2GFP DRGs (Fig. 1E, bottom middle) was also evidenced by the marked reduction in neurons coexpressing TrkB and GFP, whereas the conventional TrkB neurons were maintained at a level comparable with that of wild-type DRGs (Fig. 1E, bottom right; data not shown). These results imply that TrkB could not functionally replace TrkC and thus did not rescue TrkC neurons from death due to loss of Runx3. The data also imply that the segregation between TrkB and TrkC neurons is not dependent on Runx3. Put together, the data demonstrate unequivocally that the three REs are essential for Runx3 expression, commencing from the earliest stages of TrkC neurogenesis.

Cooperation between R1, R2, and R3 promotes Runx3 expression at E11.5

Next, we assessed the contribution of each of the three REs to TrkC neuron development through simultaneous analysis of BAC transgenic and compound CRISPR/P2GFP mouse lines bearing single or double RE deletions (Supplemental Tables S3, S4). At E11.5, each single RE conferred Runx3 expression to most TrkC neurons (Fig. 2A–C). The intensity of Runx3 expression endowed by each single RE was significantly higher than in the absence of all three REs but lower compared with control (Fig. 2D). Similar results were obtained by analyzing lines of transgenic mice bearing RE-deleted BACs (Fig. 2E).

Interestingly, the loss of just R1 (ΔR1) markedly reduced Runx3 expression level in TrkC neurons as compared with control, whereas deletion of R3 (ΔR3) had no effect (Fig. 2B,D). These results indicate that, while at E11.5, the three REs are active and act cooperatively, R1 predominates. Taken together, these analyses establish that the onset of Runx3 expression in TrkC neurons at E11 is mediated by a synergism between the three REs.

Developmental switch in RE usage at E12.5

The unique contribution of each RE to Runx3 expression in TrkC neurons became evident at E12.5. Among single RE deletions, removal of R1 (CRAR1/P2GFP) led to the most pronounced decrease in the percentage of Runx3-expressing TrkC neurons (Fig. 2F,G). Moreover, the remaining ~50% of TrkC/Runx3 neurons displayed marked reduction in Runx3 intensity (Fig. 2H). The critical role of R1 in mediating Runx3 expression was further emphasized by the observation that R1 alone drove Runx3 expression in >80% of TrkC neurons (Fig. 2G). Similar results were obtained using BAC-C-delR1 transgenic embryos (Supplemental Fig. S4A,B). Conversely, deletion of R2 in either BAC transgenic-modified or CRISPR/Cas9-modified embryos had almost no effect on Runx3 expression (Fig. 2F,G, Supplemental Fig. S4A,B). Deletion of R3 modestly affected both the percentage of reporter-expressing or Runx3-expressing cells and Runx3 expression intensity (Fig. 2F–H, Supplemental Fig. S4A,B). Of note, the activity of R3 alone at E12.5 was lower than at E11.5, as evidenced by the reduction in percentage and intensity of Runx3 expression in CRAR1,2/P2GFP mutant embryos (Fig. 2). Interestingly, deletion of both R1 and R3 completely abolished reporter or endogenous Runx3 expression in TrkC neurons (Fig. 2F–H), indicating that, unlike at E11.5, R2 does not function alone at E12.5. Thus, both BAC transgenic-modified and CRISPR/Cas9-modified embryos showed that, at E12.5, R1 is the dominant RE, R3 exhibits weak activity, and R2 does not confer Runx3 expression in TrkC neurons. Of note, at this developmental stage, Runx3 was detected in wild-type and various CRΔRE mutants as well as in non-TrkC neurons (Fig. 2F), as discussed below.

Loss of Runx3 at E12.5 results in depletion of TrkC neurons at later developmental stages

Cell count of TrkC/GFP neurons in serial sections of various RE-deleted/P2GFP compound embryos revealed that, in most cases, neurons that had lost Runx3 expression at E12.5 disappeared at E15.5. This conclusion was supported by several lines of evidence: First, at E15.5 and E17.5, in all brachial DRGs, the presence of R1 was sufficient for maintaining a number of TrkC/Runx3 neurons similar to that in wild-type/P2GFP control (Fig. 3A,B, Supplemental Figs. S5, S6). This result corresponds with the observation that R1 alone conferred Runx3 expression to most TrkC/Runx3 neurons even at E12.5. In contrast, in the absence of R1, ~40% of TrkC neurons were lost at E15.5, similar to the percentage of neurons lacking Runx3 at E12.5 (Figs. 2G, 3A). Of note, a fraction of the remaining TrkC/GFP neurons in R1-deleted DRGs did not express Runx3 (Fig. 3A,B, Supplemental Figs. S5, S6), indicating that these cells can survive without Runx3. These Runx3-lacking neurons persisted also at E17.5 (data not shown). Second, at E15.5, in embryos bearing only R2, no Runx3-expressing TrkC/GFP neurons survived (Fig. 3B), as anticipated from the lack of Runx3 expression in these neurons already at E12.5 (Fig. 2G). Last, in the absence of R3, ~80% of TrkC/GFP neurons survived, and most of them expressed Runx3 (Fig. 3A,B), while, in the presence of R3 alone, most TrkC/GFP neurons that retained low levels of Runx3 at E12.5 (Fig. 2G ~40%) survived at E15.5 (Fig. 3A).
Figure 2. RE activity at E11.5–E12.5. (A–E) At E11.5, the three REs synergize in promoting Runx3 expression. (A) Expression of Runx3, TrkC, and GFP in DRG sections of various CRΔRE mutants. (B) Runx3 expression in the same sections as in A. (C) The percentage of TrkC/GFP neurons that express Runx3 in RE-deleted embryos. (Ct) Wild-type/P2GFP; (ΔR1) CRΔR1/P2GFP; (ΔR3) CRΔR3/P2GFP; (ΔR1,2,3) CRΔR1,2,3/P2GFP. R1, R2, and R3 represent the remaining single RE following removal of the two other REs. (*) P < 0.005; (**) P < 0.0001. (D) Runx3 expression intensity. (*) P < 0.0001 compared with Ct; (**) P < 0.0001 compared with ΔR1,2,3. (E) Percentage of TrkC/Runx3 neurons expressing GFP from BAC-C-GFP (Ct) and its RE-deleted (d) variants. (*) P < 0.005; (**) P < 0.0001. (F–H) Differential RE activity at E12.5. (F) Runx3, TrkC, and GFP expression at E12.5 in various CRΔRE genotypes. (G) The percentage of Runx3/TrkC/GFP neurons out of TrkC/GFP neurons. (*) P < 0.005; (**) P < 0.0001. (H) Intensity of endogenous Runx3 expression in TrkC neurons at E12.5. (*) P < 0.0002; (**) P < 0.0001 compared with Ct; (**) P < 0.0002; (**) P < 0.0001 compared with ΔR1,2,3. Bar, 50 μm.
Remarkably, at E15.5, the intensity of Runx3 in neurons bearing only R3 was similar to that in the control and much higher than at E12.5 (Fig. 3B,C). This finding indicates that R3 efficacy increased with embryonal age. Interestingly, RE mutants exhibited a similar pattern of Runx3 loss in the trigeminal ganglion (TG). Like in DRGs, deletion of either all three REs or only the duo R1 and R3 led to the complete loss of Runx3 expression in TG neurons, whereas removal of R1 alone resulted in the loss of Runx3 in many, but not all, TG TrkC neurons. However, unlike in DRGs, a substantial number of TG TrkC neurons still survived in the absence of Runx3 (Supplemental Fig. S7).

Mice lacking both R1 and R3 develop severe limb ataxia

As shown before, mice lacking the entire 25-kb genomic region harboring the three REs developed severe limb ataxia. Importantly, a precise deletion of R1 and R3 had a similar effect despite the retention of R2 and flanking sequence [Fig. 3D; Supplemental Movie S1]. Of note, although Runx3 was expressed at E11.5 in DRGs lacking both R1 and R3 (Fig. 2A–E), this early R2-mediated Runx3 expression did not rescue TrkC neurons at later stages. In fact, just 1 d later (E12.5), the entire TrkC neuron population nearly disappeared in mice bearing R2 alone [Fig. 2F–H]. The early loss of TrkC neurons in the embryo is compatible with the severe ataxia observed in these newborn mice. Conversely, the presence of R3 alone [which, at E12.5, mediated low levels of Runx3/TrkC expression] was sufficient for rescuing a subpopulation of TrkC neurons. Moreover, these R3-only (CRΔR1,2) mice did not develop ataxia. The increase in R3 activity during later embryonic development may explain the seemingly normal gait of R3-only mutant mice [Fig. 3B,C]. These results demonstrate that loss of R1 + R3 per se is sufficient for ataxia development.

Figure 3. RE deletion causes a differential loss of TrkC/Runx3 neurons at E15.5. (A) The percentage of Runx3/TrkC/GFP [dark gray] or total TrkC/GFP neurons [light gray] surviving in various ΔRE embryos relative to the corresponding neuron numbers in Ct ganglia. The percentage was determined in serial sections of C6–T1 ganglia [Supplemental Fig. S5]. (P) P < 0.05; (**) P < 0.001; (***)) P < 0.0001. (B) Expression of Runx3, TrkC, and GFP at E15.5 in C7–C8 ganglia. The last panel at the right in the bottom row shows Runx3 expression in ΔR1,2 in the same DRG section as the one adjacent at its left. (C) As detailed in B for ΔR1,2 at E12.5 [left panel] and Runx3 expression in the same section [right panel]. (D) ΔR1,3 mice encounter severe congenital ataxia. Picture of a 3-wk-old ΔR1,3 mouse [see Supplemental Movie S1]. Bar, 50 μm.
Loss of the R1-dependent TrkC neuronal subpopulation is associated with reduced central afferents and defects in mouse locomotor activity

To further investigate the function of the three REs, we immunostained the spinal cords of various CRΔRE/P2GFP mouse mutants for the Ca++-binding protein parvalbumin (Pvalb), which is expressed in most TrkC/Runx3 neurons starting at E14.5. This enabled us to monitor spinal cord proprioceptor-central afferents at late embryonic stages. Deletion of all three REs resulted in the complete loss of central afferents corresponding to the aforementioned early loss of TrkC neurons in CRΔR1,2,3 mice. A solitary R3 deletion caused only a minor effect, whereas R1 deletion led to a substantial reduction in the number of central afferents. Deletion of both R1 and R2, retaining only R3, exacerbated the loss of ventral afferents as compared with R1 deletion [Fig. 4A]. This result reflects the relatively small size of the R3-dependent subpopulation and the late onset of Runx3 expression in this population [Fig. 3B,C], which is essential for afferent extension [Lallemend et al. 2012]. Nevertheless, as noted before, this small number of afferents sufficed to support an outwardly normal gait.

To associate neuronal loss with functional consequences, we subjected the mice to several basic behavioral tests. The home cage locomotion test, which quantifies voluntary spontaneous activity in the home cage, revealed almost no difference between CRΔR1/P2GFP and CRΔR3/P2GFP mice as compared with their respective wild-type/P2GFP controls. On the other hand, CRΔR1,2/P2GFP mice displayed significantly lower mean hourly activity [Fig. 4B]. Performance in the horizontal beam-walking test required precise foot placing and assessed forced or challenged motor balance and coordination [Luong et al. 2011]. In this test, both CRΔR1,2/P2GFP and CRΔR1/P2GFP mice displayed a significantly higher rate of slippage from the beam relative to control wild-type/P2GFP mice. Effect size analyses revealed that the absence of both R1 and R2 was associated with more frequent foot slippage compared with the loss of only R1 (Fig. 4C). The time lapse required for beam crossing and the number of steps on the beam also tended to be higher in the CRΔR1,2/P2GFP group [data not shown]. The poorer performance of CRΔR1/P2GFP and CRΔR1,2/P2GFP mice relative to CRΔR3/P2GFP mice in the beam-walking test indicates that retention of the R1-dependent population is mandatory for foot-placing accuracy. The increased effect of ΔR1,2 compared with ΔR1 in the behavioral tests uncovered a functional role for R2, the physiological ramification of which is described below.

Figure 4. Projection of central afferents and functional analysis of RE-dependent TrkC/Runx3 subpopulations. (A) Central afferents of E17.5 C7 DRGs in sections of various RE-deleted embryos visualized by Pvalb staining. The bottom panels depict the corresponding enlarged insets. (B) Home cage locomotion: mean hourly activity assay during dark/active and light/dormant phases. ΔR3 [n = 9], ΔR1 [n = 25], ΔR1,2 [n = 16] male mice were compared with age-matched wild-type/P2GFP (Ct) mice [n = 10, n = 12, and n = 27, respectively]. [*] P < 0.05, [**] P < 0.01 compared with Ct, Student t-test. [**] P < 0.01 ΔR1,2 > ΔR3 and ΔR1, ANOVA followed by Scheffe post-hoc comparisons. (C) Beam-walking assay. The same mice as in B were examined. Falling ratio is shown, Mann-Whitney comparisons indicate ΔR1 > Ct (P = 0.003) and ΔR1,2 > Ct (P = 0.000). [**] P < 0.01. There is a difference in the effect sizes (r) between ΔR1 versus Ct [0.462] and ΔR1,2 versus Ct (0.700); ΔR1 is considered “medium” (0.3 < r < 0.5), while ΔR1,2 is considered “large” (>0.5). Kruskal-Wallis followed by Dunn’s corrected pair-wise comparison. [**] P = 0.003 ΔR1,2 > ΔR3. Bar, 50 μm.
**Differential activity of R1–R3 determines Runx3 expression in distinct subpopulations of DRG TrkC neurons**

Our observation that loss of both R1 and R3, but not of each of them alone, resulted in severe ataxia indicates that the function of these two REs in TrkC neurons is additive. To further evaluate the interrelationship between the three REs in different ganglia, we determined the number of surviving TrkC/Runx3 neurons in C5–T1 DRGs of CRAPP2GFP mutants and wild-type/P2GFP control mice. In DRGs at spinal level C5, deletion of R1 or R3 had little or no effect, respectively (Fig. 5A; Supplemental Figs. S5, S6), suggesting that the activity of R1 and R3 in this ganglion is redundant. Further analysis revealed that R1 on its own mediated Runx3 expression in all TrkC/GFP neurons (Supplemental Figs. S5, S6), whereas R3 conferred expression in ∼55% of the neurons, implying that the apparent redundancy between them is limited to that population (Fig. 5A, A2). Another R1-dependent subpopulation, comprising 15% of C5 TrkC/Runx3 neurons, consisted of neurons that lost Runx3 upon R1 deletion (Fig. 5A, A1). The remaining 30% of C5 TrkC/Runx3 neurons endured either R1 or R3 deletion due to redundant activity of R1 and R2 + R3 (Fig. 5A, A3). This last conclusion is based on the observation that while R3 alone drove Runx3 expression in 55% of TrkC/Runx3 neurons, the combination of R2 + R3 conferred expression in 85% of them. The data indicate that R2, which is not active alone, supports R3 activity in the A3 population [Fig. 5A, A3]. In summary, counting all TrkC/Runx3 neurons in the C5 ganglion of the various CRAPP2GFP mutant mice defined three neuronal populations: A1, an R1-dependent population comprising 15% of C5 TrkC/Runx3 neurons; A2, an R1/R3-dependent population comprising 55% of C5 TrkC/Runx3 neurons; and A3, an R1/R2 and R3-dependent population comprising 30% of these neurons.

In C6–C7 ganglia, unlike in C5, R1 deletion caused a pronounced (50%) loss of TrkC/Runx3 neurons (Fig. 5B; Supplemental Figs. S5, S6). Thus, 50% of TrkC neurons in these ganglia are R1-dependent (Fig. 5B, B1). R3 deletion showed that an additional 30% of C6–C7 neurons are R3-dependent, as its deletion eliminated 30% of TrkC/Runx3 neurons, and, vice versa, R3 alone mediated...
Runx3 expression in ~30% of TrkC neurons [Fig. 5B,R2]. In the remaining 20% of TrkC neurons, Runx3 expression was redundantly regulated by R1 plus R2 and R3 [Fig. 5B, B3]. This B3 population in C6–C7 ganglia is similar to the A3 subpopulation in the C5 ganglion. We also noted that the R1 plus R2 [ΔR3] population, comprising 70% of TrkC/Runx3 neurons, was significantly smaller compared with the R1-only [ΔR2,3] population, which comprised ~100% of TrkC/Runx3 neurons [Fig. 5B]. This result indicates that R2 represses R1 activity in 30% of TrkC/Runx3 neurons [Fig. 5B,B2]. Accordingly, the R1-repressed population corresponds to the R3-dependent (AR1,2) population.

In the C8–T1 ganglia, a similar pattern of R1 and R3 segregation generated three subpopulations, of which C1 is analogous to A1 and B1, and C2 is analogous to B2 [Fig. 5C,C1,C2]. The third C8–T1 subpopulation [Fig. 5C,C3] was inferred by combined R1 activity and R2-mediated repression of R3. This interpretation was borne out of the observation that, in the R3-alone [AR1,2] subpopulation, Runx3 was expressed in ~50% of the neurons, whereas the presence of R3 and R2 (i.e., Fig. 5C, AR1 or R2,3) reduced the number of Runx3-expressing TrkC neurons to 30%. Taken together, these results reveal that, in the C5–T1 DRGs, TrkC neurons segregate into five distinct subpopulations according to their dependence on various combinations of RE-mediated Runx3 expression: (1) R1-dependent (A1, B1, and C1), (2) R3-dependent (B2 and C2), (3) R1/R3-dependent [A2], (4) R1/R2 and R3-dependent [A3 and B3], and (5) R1-dependent with R3 repressed by R2 (C3). The differences in abundance of these subpopulations along consecutive brachial DRGs are likely to reflect subtle changes in the proprioceptive properties of their respective target muscles.

Importantly, these neuronal subpopulations differ functionally, as demonstrated by the behavioral tests. Specifically, the precision of foot placing in the beam test singled out the R1-dependent subpopulations (A1, B1, C1, and C3), as the loss of these subpopulations increased foot slippage [Figs. 4C, 5A–C, AR1 in A4,B4,C4]]. Conversely, the home cage locomotion scores of mutants lacking these R1-dependent populations were similar to those of wild-type mice, suggesting that they are less important for gross motor function. However, upon double deletion of R1 and R2 [ΔR1,2,P2GFP], which resulted in loss of the main R1-dependent subpopulations A1, B1, and C1 as well as the A3 and B3 subpopulations, we observed a reduction in home cage hourly activity and exacerbation of foot slippage in comparison with ΔR1. Thus, the further removal of the A3 and B3 subpopulations, retaining only the R3-dependent subpopulations (B2 and C2), impaired both mouse locomotion and coordination. Moreover, deletion of these R3-dependent subpopulations [Fig. 5B, C, ΔR3 and AR3,2 in B4,C4]] was not associated with any behavioral defect, supporting the conclusion that R3-dependent populations are not essential for gross locomotion or precise foot placing. Thus, an intricate combinatorial interplay among the three REs drives Runx3 expression in distinct DRG neuronal subpopulations that play specific functional roles.

R1 represses R2-driven Runx3 expression in TrkA neurons

In addition to the high expression levels in wild-type TrkC neurons, we also detected lower Runx3 levels in non-TrkC neurons of E11.5 and E12.5 embryos [Fig. 2A,F]. Interestingly, this early Runx3 expression was confined to ~40%–50% of TrkA neurons [Fig. 6A,C]. Because the percentage of Runx3-expressing TrkA neurons in mutant embryos lacking the three REs was similar to that in wild-type/P2GFP mice, we deduced that early Runx3 expression in TrkA neurons was driven solely by R2. Importantly, P2 activity in TrkA neurons disappeared at a later embryonic stage, as demonstrated by the lack of Runx3 in either CRΔR1,2,3/P2GFP or wild-type E14.5 TrkA neurons [Fig. 6B,C]. Of note, upon deletion of R1 alone (CRΔR1/P2GFP), Runx3 was readily detected in most E14.5 TrkA neurons [Fig. 6B,C]. Thus, R1 simultaneously promotes Runx3 expression in TrkC neurons and represses it in TrkA neurons. This conclusion was supported by a similar result obtained using BAC transgenics [Fig. 6D,E]. Importantly, in contrast to its inability to drive Runx3 expression in TrkC neurons, R2 alone (CRΔR1,3/P2GFP) conferred Runx3 expression in TrkA neurons [Fig. 6B,C]. Moreover, in BAC transgenics, reporter expression in TrkA neurons was driven only by constructs bearing R2 and lacking R1 but not in mutants lacking R2. Collectively, the complementary results of in vivo CRΔRE/P2GFP mutants and BAC transgenics demonstrate that R2 positively regulates Runx3 expression in TrkA neurons. R3 can cooperate with R2, as a higher percentage of TrkA neurons expressed Runx3 when both R2 and R3 were present (CRΔR1,3/P2GFP), as compared with R2 alone [Fig. 6C]. This occurrence was particularly pronounced in E14.5 mutant embryos. In the context of the spatiotemporal regulation, the data demonstrate that Runx3 expression specificity in TrkC neurons is determined by not only R1-enhancive activity but also R1-mediated Runx3 repression in TrkA neurons [Fig. 6F].

Runx3 is not expressed in wild-type TrkB neurons [Supplemental Fig. S8A,C]. However, deletion of R2 from BAC-C [BAC-C-delR2] revealed an early stage [E11.5] BAC-mediated GFP expression in a small number of TrkB neurons that, at E12.5–E14.5, comprised nearly 80% of these neurons [Supplemental Fig. S8B,D]. This observation implies that, in the context of BAC transgenics, R2 mediates Runx3 repression in TrkB neurons and that R2 activity may be essential for TrkB neuron survival. However, analysis of E12.5 CRΔR2/P2GFP or homozygous AR2/ΔR2 embryos revealed only a small population of Runx3-expressing TrkB neurons [Supplemental Fig. S8E]. This small population was absent at E17.5, with a minute decline in the number of TrkB neurons in AR2/ΔR2 mice compared with control [data not shown]. Thus, excision of R2 from BAC-C led to reporter-GFP expression in ~80% of TrkB neurons, while its deletion in vivo [AR2/AR2] produced only a small number of TrkB/Runx3 neurons, indicating that R2 alone could not repress Runx3 expression in these cells in-vivo. Together, these results suggest that Runx3 protein levels in TrkB neurons may be regulated at the
translation/degradation level and that the absence of Runx3 in wild-type TrkB neurons may be controlled by mechanisms yet to be elucidated. Alternatively, BAC-C could lack an additional silencing element that is present in the genome, leading to the difference between the two model systems.

**BRNF TF-binding sites regulate R1 and R3 activity**

We next sought to identify upstream TFs that regulate Runx3 RE activity. The amount of TrkC neurons that can be isolated from DRGs is insufficient for TF-specific ChIP-seq (chromatin immunoprecipitation [ChIP] combined with high-throughput sequencing) analyses. Therefore, we conducted RNA sequencing (RNA-seq) using RNA isolated from purified TrkC/GFP neurons of wild-type/P2<sup>GFP</sup> and ΔR1,2,3/P2<sup>GFP</sup> mice. As seen in Supplemental Table S6, several TFs were highly expressed by these neurons, most of which were not affected by the loss of Runx3. Among these TFs, the POU homeodomain Brn3a was implicated previously in Runx3 regulation (Dykes et al. 2010). A conserved Brn3a-binding site located 94 kb upstream of Runx3 (marked as BrnT in Fig. 7A) was shown previously to bind Brn3a (Dykes et al. 2010). Interestingly, this −94-kb genomic region colocalized with R3, implying that Brn3a is one of the TFs regulating R3 activity. Sequence alignment analysis of the R3 genomic region for conserved TF-binding sites identified both BrnT and an additional Brn3a-binding site (marked BRNF-R3). To directly demonstrate the in vivo participation of Brn3a in the regulation of Runx3 in TrkC neurons, we mutated either the BrnT site or the two BRNF-R3-binding motifs in BAC-C. Both the BrnT and BRNF-R3 mutations caused a similar effect, mirroring the deletion of an entire R3 [Fig. 7B,C]. These results support the notion that Brn3a is an important upstream regulator of R3-dependent Runx3 expression in TrkC neurons.
Figure 7. BRFN-binding sites are essential for R1 and R3 activity. (A) Evolutionarily conserved BRFN-binding sites within R3 core region. Red boxes mark the BRFN-binding sites identified by DiAllign TF software in five or six species using core similarity 0.75 and matrix similarity optimized. The underlined binding site, marked as BrnT, corresponds to the site identified by Dykes et al. (2010). Mutations introduced into BRFN-R3-binding sites and BrnT-binding sites, labeled as mutR3 and mutR3T, respectively, are shown below the red boxes. (B) Expression of GFP in TrkC but not TrkA neurons in E14.5 BAC transgenic embryos. (C) The percentage of TrkC neurons expressing GFP at E12.5 (white) or E14.5 (gray) in transgenic DRGs expressing intact BAC-C (Ct) and either deleted or mutated BAC-C. ∗ P < 0.001 compared with Ct. (D) An evolutionarily conserved BRFN-binding site within the R1 [BRFN-R1] core region is marked by dark green, while a conserved SORY-binding site is marked by light green. Both sites were identified by DiAllign TF software in five or six species using core similarity 1 and matrix similarity 0.9. (E) Expression of GFP by BAC-C-delR1 (left) and BAC-C-mutR1 (right) in TrkC and TrkA neurons of E14.5 transgenic embryos. (F) The percentage of GFP-expressing TrkC neurons in BAC transgenics (white, left panel) and the percentage of Runx3-expressing TrkC neurons in endogenously mutated embryos (gray, right panel) at E14.5. The mean for DRGs C6–T1 is presented. ∗∗ P < 0.0001 compared with Ct. (G) As detailed in F for TrkA neurons. Bar, 50 μm.
Because Runx3 expression is abolished in Brn3a-deficient mice (Eng et al. 2007), we hypothesized that, besides its impact on R3 activity, Brn3a also regulates the activity of R1. Inspection of the core genomic region of R1 for conserved TF-binding sites revealed a BRNF-R1-binding site (Fig. 7D). Mutating this binding site in either BAC-C or the mouse genome caused partial impairment of R1-enhancing activity [Fig. 7E]. Specifically, mutations in the BRNF-R1-binding site of BAC-C reduced GFP expression in TrkC neurons as compared with intact BAC-C. However, the effect was significantly smaller than that upon complete R1 deletion [Fig. 7F, left panel]. A similar tendency was observed in CRISPR/Cas9-mediated mutation [Fig. 7F, right panel]. Interestingly, the BRNF-R1 mutation also affected the repression activity of R1 in TrkA neurons. As noted previously, Runx3 expression in TrkA neurons is mediated by R2 and inhibited by R1. Consistently, E14.5 transgenics bearing intact BAC-C did not express GFP in TrkA cells. BAC-C-mutR1 conferred Runx3 expression in 50% of TrkA neurons [Fig. 7G, left panel], whereas CRISPR-mutR1 drove Runx3 in 30% of TrkA neurons [Fig. 7G, right panel]. In both cases, the mutation had a lesser effect relative to complete removal of R1.

Together, these data indicate that BRNF-binding sites participate in the enhancive and repressive activities of R1. However, the notably milder phenotype of BRNF-R1 mutants as compared with mutants lacking the entire R1, raises the possibility that additional TFs are involved in regulating R1 function in TrkC neurons. Indeed, RNA-seq analysis revealed that besides Brn3a [Pou4f1], three other TFs—Isl1, Sox11, and NeuroD1—were highly expressed in purified TrkC neurons [Supplemental Table S6]. Sox11, which was shown to regulate sensory neuron development and survival [Lin et al. 2011], is an interesting candidate because R1 contains a conserved SORY-binding site [Fig. 7D]. The R1 Isl1-binding site is probably of lesser importance because the loss of Isl1 hardly affects Runx3 expression [Huang et al. 1999; Quina et al. 2009; Dykes et al. 2011], while NeuroD1 emerged as a Runx3-responsive gene [Supplemental Table S6], placing it downstream from Runx3. Of note, comparison of RNA-seq data from wild-type/P2ΔGFP and P2ΔGFP/GFP neurons [GSE81140] [Supplemental Table S7] revealed that a group of genes previously identified as targets of Brn3a [Dykes et al. 2011] was Runx3-responsive. This overlap may indicate that those presumable Brn3a target genes are, in effect, regulated by Runx3 [Supplemental Fig. S9].

Discussion

REs located 100–125 kb upstream of P2 regulate TrkC neuron expression of Runx3

In the present study, we delineated the Runx3 gene transcriptional unit. It spans ~170 kb of genomic DNA, encompassing the gene itself and the REs that control its spatiotemporal expression specificity in different tissues. Ten CNEs were identified in this region, three of which—R1, R2, and R3—are essential for specific Runx3 expression in TrkC neurons. These REs most likely mediate Runx3 expression exclusively in DRGs and TGs, as none of the other putative REs that regulate Runx3 in other tissues overlap with these TrkC-specific REs [Ghisletti et al. 2010; Nakayamada et al. 2011; Vahedi et al. 2012; Hnisz et al. 2013; McClellan et al. 2013; Ohba et al. 2015; Zhou et al. 2015; Gunnell et al. 2016].

As shown previously [Levanon et al. 2003; Sullivan et al. 2008], Runx3 functions as an important component of the ancient stretch reflex neuronal circuit. This role corresponds with the finding that the three REs collaborate with the more evolutionarily conserved promoter P2 [Levanon and Groner 2004], i.e., the three REs skip the more upstream, less conserved P1 in favor of P2. Furthermore, homozygous P2ΔGFP/GFP mice bearing an intact P1 developed severe limb ataxia due to loss of Runx3 in early developing TrkC neurons. Thus, in TrkC neurons, unlike in other cell types, P1 does not compensate for the absence of P2 [Levanon et al. 2014]. Of note, two additional P2 features—the upstream distant location of REs and the presence of an Inr motif within the promoter [Bangsow et al. 2001]—are typically found in developmentally regulated genes [Lorberbaum and Barolo 2015].

Combinatorial RE cross-talk regulates Runx3 in different TrkC neuronal subpopulations

Proprioceptive neurons consist of a highly diverse population. Accordingly, interneurons and motor neuron pools receive specialized input from selected groups of sensory neurons [Eccles et al. 1957; Mears and Frank 1997; de Nooij et al. 2013], yet little is known about proprioceptor specification. de Nooij et al. [2013] have shown that proprioceptive neurons destined to innervate distinct muscle targets differ markedly in their Etv1 TF-dependent survival and differentiation. Using single-cell RNA-seq, two proprioceptive subtypes were detected in adult mouse DRGs [Usoskin et al. 2015]. It was also reported that mesenchymal signals expressed in restricted dorsoventral and proximodistal domains of the developing limb are essential for endowing muscle-type identity to two distinct proprioceptive subtypes, marked by the expression of either Cdh13, Sema5, or Crtac1 [Poliak et al. 2015].

Our analysis revealed that TrkC neurons also diverge in the pattern of Runx3 expression regulation. Differential usage of Runx3 REs defines five TrkC subpopulations in brachial DRGs. Of these, R1- and R3-dependent subpopulations dominate in C6–T1 ganglia. The R1-dependent population [Fig. 5, A1,B1,C1], which is characterized by intense Runx3 expression commencing at early embryonic stages, is the major subgroup of TrkC neurons in these ganglia. Neither R3 nor R2 could rescue these TrkC neurons upon R1 deletion. The importance of R1 is further emphasized by the observation that, in its absence (∆R1), the precision of foot placing deteriorates. Foot-placing precision depends on proprioceptive feedback from muscle spindles [Akay et al. 2014], raising the possibility that the R1-dependent subpopulation is involved in signaling from muscle spindles. The R3-dependent population [Fig. 5, B2, C2] is smaller and develops at later embryonic stages. In this population, Runx3 expression...
was hardly detected at E12.5 and increased only later on. Deletion of R3 led to loss of the B2 and C2 subgroups but did not have a notable effect on either home cage locomotion or the beam test. However, mice with deletion of R1 and R2 (ΔR1,2)—i.e., the R3-alone mice—performed poorly in the beam test and exhibited reduced home cage locomotion. Nevertheless, these mice were not ataxic. Moreover, analysis of ΔR1,2 spinal cords revealed fewer afferents, most of which did not reach the ventral zone. These observations indicate that the R3-dependent subgroup is not sufficient to support normal locomotion and precision of foot placing. In addition, R2, which is not active on its own in TrkC neurons from E12.5 onward, does affect R3 activity. However, we cannot rule out the possibility that the behavioral defects are due to changes in higher brain functions that were not assayed in this study and are secondary to the loss of spinal connectivity.

R1 and R3 also differ in their dependence on the POU domain TF. Mutations in BRFN-R3-binding sites led to a phenotype similar to upon deletion of the entire R3. In contrast, a mutation in the BRFN-R1-binding site had a significantly lesser effect. In both REs, this binding site is probably occupied by Brn3a, which is the major POU domain TF in E11.5 TrkC neurons. RNA-seq data analysis demonstrated a particularly high expression level of both Brn3a and Sox11. The finding that R1 has a conserved SORY-binding site raises the possibility that Brn3a and Sox11 TFs cooperate in regulating R1 activity in TrkC neurons. Thus, it was interesting to note that POU- and Sox-binding sites are also situated adjacent in genomic regions regulating subtype specification of dorsal spinal cord neurons (Borromeo et al. 2014). The fact that the three REs are active already at E11.5 implies that a common TF, most likely Brn3a, regulates Runx3 expression at this developmental stage. Interestingly, after E11.5, as additional TFs come into play, the relative contribution of various TFs to RE function diverges, resulting in the observed differential activity of the three REs.

The stringent cell-specific Runx3 expression observed in TrkC neurons is attained by not only the REs’ enhancing capacity but also their inhibitory activity in non-TrkC neurons. The establishment of cell-type specificity through repressing expression in the irrelevant cells is a widely accepted paradigm [Lanier et al. 2009]. In the developing DRGs, R1 functions as a repressor in TrkA neurons, abolishing the R2-mediated Runx3 expression in these neurons. Additionally, R2 exerts Runx3 repression in TrkB neurons, facilitating the gain of their identity. REs that function in an opposing manner to specify cell identity were also reported for other genes, including Sonic hedgehog (Shh) [Lettice et al. 2012].

The spatiotemporal specificity of developmental TFs such as Runx3 is commonly regulated by multiple REs [Lagha et al. 2012]. Several modes of action have been described for gene expression regulation by multiple REs. The elements act either synergistically to increase robustness or antagonistically to enhance specificity [Andrey et al. 2013; Levine et al. 2014]. Alternatively, REs may act sequentially or as shadow REs that balance genetic or environmental perturbations [Schwarzer and Spitz 2014] or else respond dynamically to various stimuli [Joo et al. 2016]. At the onset of Runx3 expression, R1, R2, and R3 act synergistically but segregate later to establish the spatiotemporal expression patterns in distinct TrkC subgroups. Thus, each RE has a specific role, and none acts as a shadow enhancer. Overall, this Runx3 RE ensemble functions as an integral unit, underscoring the Heinz et al. [2015] paradigm of superenhancers controlling expression of lineage-specific genes.

**Transitory Runx3 expression at E11.5 fails to rescue TrkC neurons**

As noted earlier, low amounts of Runx3 were detected in E11.5 DRGs of ΔR1,3 mice. However, the exclusively R2-driven Runx3 expression completely disappeared at E12.5, as did the prospective TrkC neurons. Accordingly, these mice encountered a severe neonatal ataxia that recapitulated the Runx3Δ−/− phenotype (Levanon et al. 2002). This finding indicates that transient Runx3 expression commencing at E11.5 does not rescue proprioceptor development. In contrast, ΔR1,2 mice expressing only low Runx3 levels at E12.5 showed a later gain of Runx3 expression and exhibited no ataxia, although they did display locomotion defects in home cage and beam tests. Given that normal locomotion requires proper function of extensor and flexor muscles, which form connections with distinct TrkC subtypes [Poliak et al. 2016], the fact that ΔR1,2 mice are not ataxic supports the notion of functional heterogeneity within this R3-only TrkC neuron subgroup. Moreover, given that R3 is less conserved than R1 [Supplemental Table S2], this subgroup may be associated with a phylogenetically more recent function of TrkC neurons.

A somewhat similar situation occurs in the Isl1 gene, where the less evolutionarily conserved CREST2 enhancer, which coappears with limb development, is required for a new subtype of motor neurons [Kim et al. 2015].

**Materials and methods**

**Mouse strains**

The experiments were conducted in strict accordance with the recommendations of the US National Institutes of Health Guide for the Care and Use of Laboratory Animals. The protocols were approved by the Weizmann Institute of Science Committee on the Ethics of Animal Experiments. P1^{APP} and P2^{APP} mice were described previously [Levanon et al. 2011]. Further information regarding mouse strains is in the Supplemental Material.

**CNE identification**

For CNE identification, we performed sequence alignments using genomic sequences spanning the mouse Runx3 locus [chromosome 13: 134,953,991–135,208,237; mm10] according to the University of California at Santa Cruz [UCSC] genome browser [http://genome.ucsc.edu] and the corresponding genomic regions from humans [human February 2009 assembly], rats [rat November 2004 assembly], chickens [chicken November 2011 assembly], frogs [Xenopus tropicalis November 2009 assembly], and fugu [fugu October 2011 assembly]. The mouse genomic
sequence served as the basis for sequence alignments using M-LAGAN [http://genome.lbl.gov/vista/index.shtml] (Brudno 2007).

**BAC reporter constructs**

Six BACs were obtained from Children’s Hospital Oakland Research Institute [http://bacpac.chori.org] [Supplemental Table S1]. Each BAC was modified using Quick & Easy conditional knockout kit –FRT (GeneBridges) by insertion of LacZ or EGFP in-frame into exon 3 of the murine Runx3 at genomic position chromosome 4: 134,711,240, mm9 assembly, UCSC genome browser [http://genome.ucsc.edu].

To reduce variability, transient transgenic embryos or established BAC construct lines (at least two) were selected based on E14.5 whole-mount reporter intensity and precision in recapitulating the endogenous Runx3 expression in the skeleton, whiskers, eyelids, and nose (Levanon et al. 2001, 2011).

**CRISPR/Cas9-mediated in vivo deletions**

The single guided RNA (sgRNA) sequences used in this study are in Supplemental Table S4. Cas9 and sgRNA plasmids encompass the T7 promoter. sgRNA and Cas9 RNA were purified using MEGAclear kit (Life Technologies, AM 1908). sgRNAs were microinjected (2.5 µg each of 5 sgRNAs) into fertilized mouse eggs. At least two distinct mouse lines were analyzed for each deleted or mutated RE. Further information regarding CRISPR/Cas9 deletions is included in the Supplemental Material.

**LacZ, GFP, and immunofluorescence analyses**

X-Gal staining, GFP expression, and immunofluorescence analyses were performed on brachial DRG sections as described previously [Levanon et al. 2001, 2011]. Further information is included in the Supplemental Material.

**Determination of the percentage of GFP- or Runx3-retaining TrkC neurons**

The percentage of either BAC-derived GFP-expressing neurons or Runx3 expression in CRISPR/Cas9-derived mutants was determined by monitoring TrkC, GFP, and Runx3 immunofluorescence. At least two different lines of either transgenic BAC or CRISPR/Cas9 mutants were analyzed. For each line, sagittal sections of at least four brachial-level DRGs from three to seven embryos were monitored. Percentage was calculated by dividing the number of Runx3/TrkC/GFP neurons by the total number of TrkC/GFP neurons (in CRISPR/Cas9 mutants) or by TrkC/Runx3 neurons (in BAC transgenics).

**Cell number determination**

Neuronal cell counting was performed on serial transverse sections of C5–T1 brachial ganglia from two to four embryos for each genotype and from four wild-type/P2GFP control embryos at E15.5 (4 µm) and E17.5 (10 µm). Given that ImagePro analysis determined the average diameter of a TrkC neuron as ~19 µm, every fourth section of right and left ganglia was analyzed using the ImageJ software. To obtain the number of positive neurons per ganglion, the average number of positive cells per section was multiplied by the number of sections and divided by 4.75 [19 µm diameter, 4-µm section]. Further information regarding determination of Runx3 intensity is included in the Supplemental Material. Kolmogorov-Smirnov statistical test was used for analyses of immunohistochemical data. Results obtained from lines of the same genotype were compiled based on their insignificant deviation. Data are presented throughout as mean and SEM.

**Behavioral studies**

Home cage locomotion was assessed on males using the InfraMot system [TSE Systems, http://www.tse-systems.com/products/behavior/homecage/phenomaster/activity/inframot.html]. The beam-walking test was conducted using a 6-mm-wide beam as detailed in the Supplemental Material. The following indices were examined: time to cross over (in seconds), number of steps, and falls ratio (percentage = number of falls/number of steps).

**Analysis of TF-binding sites**

Sequences of R1 and R3 were analyzed for TF-binding sites using DiAlign TF; multiple alignment and TF-binding sites [Morgenstem et al. 1998] are available at the Genomatix Web site [http://www.genomatix.de]. Further information regarding gene expression analysis is included in the Supplemental Material. RNA-seq data were deposited into the Gene Expression Omnibus repository under the accession number GSE81140.

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