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The H3K4me3/2 histone demethylase RBR-2 controls axon guidance by repressing the actin-remodeling gene wsp-1

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ABSTRACT

The dynamic regulation of histone modifications is important for modulating transcriptional programs during development. Aberrant H3K4 methylation is associated with neurological disorders, but how the levels and the recognition of this modification affect specific neuronal processes is unclear. Here, we show that RBR-2, the sole homolog of the KDM5 family of H3K4me3/2 demethylases in Caenorhabditis elegans, ensures correct axon guidance by controlling the expression of the actin regulator wsp-1. Loss of rbr-2 results in increased levels of H3K4me3 at the transcriptional start site of wsp-1, with concomitant higher wsp-1 expression responsible for defective axon guidance. In agreement, overexpression of WSP-1 mimics rbr-2 loss, and its depletion restores normal axon guidance in rbr-2 mutants. NURF-1, an H3K4me3-binding protein and member of the chromatin-remodeling complex NURF, is required for promoting aberrant wsp-1 transcription in rbr-2 mutants and its ablation restores wild-type expression of wsp-1 and axon guidance. Thus, our results establish a precise role for epigenetic regulation in neuronal development by demonstrating a functional link between RBR-2 activity, H3K4me3 levels, the NURF complex and the expression of WSP-1.

KEY WORDS: Epigenetics, H3K4 methylation, Histone demethylase, Neuronal development, Axon guidance, C. elegans

INTRODUCTION

Chromatin organization regulates gene expression by changing DNA accessibility and is thought to have a key role in developmental processes (Eisenberg and Shilatifard, 2010; Greer and Shi, 2012). Post-translational modifications on the N-terminal tails of histone proteins are one of the elements that can influence chromatin structure. To date, several chromatin factors with the ability to catalyze the addition (‘writers’) or removal (‘erasers’) of these modifications have been identified (Zhang and Pradhan, 2014). Modified histones can modulate the strength of DNA/histone interactions as well as recruit ATP-dependent chromatin remodelers and other factors with the ability to decode the modifications (‘readers’) (Baker et al., 2008). Methylation of lysine 4 on histone 3 (H3K4) is one of the most studied modifications, with its trimethylated form (H3K4me3) enriched at transcriptional start sites (TSSs) of actively transcribed genes (Barski et al., 2007). In mammals, members of the lysine methyltransferase 2 family (KMT2) catalyze di- and tri-methylation of H3K4 (Eisenberg and Shilatifard, 2010), whereas its demethylation is executed by the four members of the lysine demethylase 5 family (KDM5) (Benevolenskaya, 2007; Pedersen and Helin, 2010; Kooistra and Helin, 2012). The biological consequences of changes in H3K4 methylation are mediated, at least in part, by molecules that recognize methylated forms of H3K4, including the plant homeodomain (PHD) zinc-finger proteins (Baker et al., 2008).

The relevance of chromatin organization in many aspects of cell biology is testified by the identification of several chromatin factors that are mutated in cancer (Helin and Dhanak, 2013; Suzuki et al., 2013). Recently, next-generation sequencing approaches have uncovered mutations in genes that encode multiple chromatin regulators in neurodevelopmental and psychiatric disorders, including intellectual disability syndromes, schizophrenia and autism spectrum disorders (for reviews, see Berdasco and Esteller, 2013; Ronan et al., 2013). Interestingly, among the mutated genes, members of the KMT2 (KMT2A/C/D) and KDM5 (KDM5A/B/C) families were identified (Shen et al., 2014; Vallianatos and Iwase, 2015), strongly suggesting that the regulation of H3K4 methylation is important to achieve correct neuronal development and functionality.

Although the catalytic properties of the KDM5 proteins are well established, relatively little is known about their biological functions. Knockout mice have been generated for some of the KDM5 family members and their characterization supports a role for this class of demethylases in neuronal development (Klose et al., 2007; Schmitz et al., 2011; Albert et al., 2013). However, the complexity of the murine nervous system and the early lethality associated with loss of some KDM5 members (Cox et al., 2010; Catchpole et al., 2011; Albert et al., 2013) have challenged the identification of their specific neuronal functions and relevant downstream targets. Thus, the study of the KDM5 family in a biologically relevant but smaller and more tractable model nervous system could provide information regarding the roles of these proteins in different aspects of neuronal development. The KDM5 family is evolutionarily conserved and is represented uniquely by the RBR-2 protein in C. elegans. Previous analyses have shown that RBR-2 is a regulator of vulva formation (Christensen et al., 2007) and aging (Greer et al., 2010, 2011), but no data are available regarding its potential function in neuronal development.

The C. elegans nervous system is the most complex tissue of the animal both in terms of numbers and diversity of cells. It consists of 302 cells, for which the lineage, morphology, migration patterns and axonal routing have been described (Sulston et al., 1983; White et al., 1986). A large number of evolutionarily conserved secreted molecules, such as Slits, Netrins, Semaphorins and Ephrins, as well as extracellular matrix components, are known to orchestrate neuronal cell migration and axon guidance (reviewed by Hatten, 2002; Hobert and Bülow, 2003; Killeen and Sybenga, 2008).
Axon guidance defects were also identified in another rbr-2 mutant allele (tm1231) (Fig. S1B), indicating that rbr-2 is required for correct axon guidance.

**RBR-2 acts during embryogenesis in the nervous system to ensure correct axon guidance**

To study the functional role of RBR-2 in axon guidance, we focused on the PVQ neurons, which fully develop during embryogenesis and are easily visualized by using the oysl14 transgene. As several molecules have been shown to play a role in maintaining rather than establishing correct PVQ axon guidance (Aurelio et al., 2002; Bülow et al., 2004; Bénard et al., 2006, 2009, 2012; Woo et al., 2008; Pocock et al., 2008), we first investigated when RBR-2 is required for the correct guidance of the PVQs. As shown in Fig. 2A, the aberrant axon guidance occurs in freshly hatched L1 mutant larvae to the same extent as observed at the adult stage, suggesting that RBR-2 is essential for establishing correct axon guidance of the PVQs during embryonic development, but it does not play any role in its maintenance. Second, we investigated in which cells/tissues RBR-2 is required to control PVQ axon guidance. The generation of transgenic lines in which RBR-2::GFP expression is driven by the rbr-2 promoter showed that RBR-2 is ubiquitously expressed in embryos and larvae (Fig. 2B). Importantly, when expressed in the rbr-2(tm3141) mutant, this transgene completely rescued the PVQ defects, further confirming that RBR-2 has a crucial role in axon guidance in these cells (Fig. 2C). To uncover the specificity of the role of rbr-2 in neuronal development, we re-expressed RBR-2 in the rbr-2 mutant background using well-characterized tissue-specific promoters, and assayed the transgenic lines for the PVQ phenotypic rescue. These experiments showed that RBR-2 is required specifically in the nervous system (F25B3.3 promoter), but its presence in hypodermal cells (dpy-7 promoter) and in muscles (myo-3 promoter) is not essential (Fig. 2C). The neuronal-specific promoter used in this assay (F25B3.3 promoter) is active, in the embryos, in four types of neurons, namely PVPs, PVQs, AVKs and embryonic motoneurons (eMN)s (Boulin et al., 2006). As the re-expression of RBR-2 in PVQs (sra-6 promoter) only partially rescues the PVQ defects observed in rbr-2 mutants (Fig. 2D), we suggest that RBR-2 acts, to some extent, cell-autonomously, but its expression might be also required in other neuronal cells to ensure correct PVQ axon guidance.

Multiple redundant pathways are known to act in parallel to drive axon guidance and genetic interaction assays have been used to identify their specific components. To test the functional relationship between rbr-2 and these pathways, we created double mutants carrying rbr-2(tm3141) together with mutations of genes acting in axon guidance pathways and measured the penetrance of PVQ defects. As shown in Table S1, we found that rbr-2 genetically interacts with all the genes tested, suggesting that rbr-2 acts in concert with the main pathways involved in axon guidance.

**The catalytic activity of RBR-2 is required for correct axon guidance and controls the expression of actin-regulating genes**

RBR-2 is the only known H3K4me3/2 demethylase in *C. elegans* (Christensen et al., 2007) and, accordingly, the global level of H3K4me3 is increased in rbr-2(tm3141) (Fig. 3A). To investigate whether the enzymatic activity of RBR-2 is required for axon guidance, we performed rescue experiments expressing a catalytically inactive form of the protein in the tm3141 genetic background (Fig. 3B; Fig. S2). In contrast to wild-type RBR-2, the mutant protein (called RBR-2DD for Demethylase Dead) was unable to
restore H3K4me3 levels (Fig. 3A) or correct axon guidance of PVQs (Fig. 3B). These results indicate that the catalytic activity of RBR-2 in the removal of the H3K4me3 mark is strictly required for the process of axon guidance.

The genetic interaction assays, showing that \( rbr-2 \) acts in concert with the main pathways implicated in axon guidance, suggest that RBR-2 might interfere with a common downstream effector and/or process. The complex signaling network initiated by guidance cues...
ultimately regulates, at growth cones, the dynamics of filopodia and lamellipodia, actin-based structures required for axon growth, branching and guidance (Kaili and Dent, 2005; Dent et al., 2011; Chia et al., 2014; Gomez and Letourneau, 2014). The role of WASP, WAVE and Ena/VASP in regulating axon growth and guidance by controlling filopodia formation (Drees and Gertler, 2008; Bear and Gertler, 2009; Tahirovic et al., 2010) is evolutionarily conserved and their C. elegans homologs (WSP-1, WVE-1 and UNC-34) are described as the main downstream targets of the signaling pathways regulating axon migration (Fig. 3C) (Shakir et al., 2008; Norris et al., 2009; Mohamed et al., 2012). Therefore, we hypothesized that the aberrant accumulation of H3K4me3 in the rbr-2 mutant could affect the transcription of actin-regulating genes, leading to the axon guidance defects observed. This hypothesis prompted us to test the role of RBR-2 in regulating the transcription of the three main actin regulators, wsp-1, wve-1 and unc-34. First, we measured the expression levels of these genes and found that they were all upregulated in rbr-2 mutant embryos (Fig. 3D). Second, we investigated whether the overexpression of wsp-1, wve-1 and unc-34 could result from higher H3K4me3 levels at their TSSs. Indeed, as shown in Fig. 3E, the H3K4me3 levels were increased in the regions surrounding the TSS of all three genes in the rbr-2 mutant compared with wild-type animals and intergenic regions. Finally, we tested the direct involvement of RBR-2 in H3K4me3 regulation by performing chromatin immunoprecipitation (ChIP)-qPCR with a GFP antibody on chromatin isolated from RBR-2::GFP-expressing worms. RBR-2::GFP was significantly enriched at the TSS of wsp-1, wve-1 and unc-34, in comparison with intergenic regions (Fig. 3F). Thus, the catalytic activity of RBR-2 is required for correct axon guidance and for the inhibition of major actin-regulating genes through a mechanism controlling the levels of H3K4me3 at their TSS.

### RBR-2 controls axon guidance by regulating WSP-1 expression

To test whether wsp-1, wve-1 and unc-34 are key downstream targets of RBR-2 in the regulation of PVQ axon guidance, we used genetic approaches. We postulated that if the PVQ defects in the rbr-2 mutant depended on the overexpression of these genes, their removal should ameliorate the phenotype. When wsp-1 was ablated in the tm3141 background, we observed a full suppression of the rbr-2 phenotype (Fig. 4A). By contrast, loss of wve-1 or unc-34, the other two major regulators of actin polymerization, did not improve the PVQ phenotype associated with rbr-2 deletion (Fig. 4A), demonstrating that WASP-1 is a key component for how RBR-2 regulates axon guidance in these cells.

To investigate further the role of wsp-1, we tested whether its overexpression in wild-type animals could reproduce the defects of the rbr-2 mutant. Indeed, overexpression of WSP-1 in wild-type animals induced the same PVQ defects observed in rbr-2(tm3141) (Fig. 4B) and re-expression of WSP-1 in rbr-2;wsp-1 double mutants also resulted in a significant increase of PVQ defects (Fig. 4C). Importantly, the effect of the increased levels of WSP-1 on PVQ axons was observed only when WSP-1 was overexpressed in neuronal tissue (F25B3.3 promoter), where we reported the action of RBR-2 in axon guidance (Fig. 4B; Fig. 2C). Collectively, these data show that the increased levels of WSP-1 expression are responsible for the axonal defects in rbr-2 mutants. In mammals, WASP has been shown to promote F-actin formation through its VCA domain, which interacts directly with the Arp2/3 complex and globular actin and stimulates actin nucleation and branching from existing filaments (Blachon et al., 2000; Pantaloni et al., 2000; Amann and Pollard, 2001; Beltzner and Pollard, 2008). Overexpression of the VCA domain leads to excess cellular protrusions in cultured mammalian cells (Yamaguchi et al., 2000) and induces axon guidance defects in C. elegans (Shakir et al., 2008). To determine whether the Arp2/3 activation domain was responsible for the rbr-2 phenotype, we overexpressed the VCA domain in the nervous system using the F25B3.3 promoter. Interestingly, ectopic expression of the sole VCA domain in wild-type animals mimicked the PVQ axonal cross-overs observed in rbr-2 mutants (Fig. 4D), strongly suggesting that such defects might arise from abnormal actin dynamics due to increased WSP-1 levels and activity. Moreover, overexpression of the VCA domain in PVQ neurons (sra-6 promoter) was sufficient to induce the phenotype, further corroborating our hypothesis that RBR-2 could act cell-autonomously (Fig. 4D).

The activity of the mammalian WASP is regulated by a plethora of proteins, of which many are evolutionarily conserved and present in C. elegans. Among others, WIP (WIPF1), CDC42 and NCK (NCK1) are known to regulate WASP activity by influencing its conformation and subcellular localization (Rivero-Lezcano et al., 1995; Symons et al., 1996; Abdul-Manan et al., 1999; Martinez-Quiles et al., 2001; Rohatgi et al., 2001; Anton and Jones, 2006; Sawa and Takenawa, 2006). We hypothesized that the removal of their C. elegans homologs, WIP-1, CDC-42 and NCK-1, could revert the axonal defects in rbr-2 worms. Strikingly, loss of wip-1, cdc-42 or nck-1 resulted in full recovery of the normal PVQ guidance in rbr-2 mutants (Fig. 4E), further supporting the key role of WSP-1 in regulating this process downstream of RBR-2.

### Role of H3K4me3 readers in axon patterning

Epigenetic regulation of transcription involves proteins that can interpret or ‘read’ the post-translational modifications of histones (Baker et al., 2008). High levels of H3K4me3 often correlate with transcriptional activation and, in agreement with this, H3K4me3 is involved in the recruitment of ATP-dependent chromatin remodeling factors (Wysocka et al., 2006) and proteins of the basal transcription machinery (Vermeulen et al., 2007). Several
reports have illustrated that H3K4me3 can be specifically recognized by proteins with a PHD zinc-finger domain (Musselman and Kutateladze, 2011), including the recombination activating gene RAG2, the large subunit of the nucleosome remodeling factor complex (NURF) BPTF, the inhibitor of growth ING2, the TAF3 subunit of TFIID (TBP) and the KDM7 family members PHF2/8 and KIAA1718 (KDM7A) (Badenhorst et al., 2002; Li et al., 2006; Peña et al., 2006; Shi et al., 2006; Wysocka et al., 2006; Matthews et al., 2007; Vermeulen et al., 2007; Palacios et al., 2008; Kleine-Kohlbrecher et al., 2010; Fortschegger and Shiekhattar, 2011). These PHD-containing proteins, except RAG2, are all conserved in C. elegans and viable mutant alleles are available for most of them (Table S2). We hypothesized that in the rbr-2 mutant the increased levels of H3K4me3 could be interpreted by one or more H3K4me3 readers, leading to inappropriate transcriptional activation of wsp-1. If this were the case, genetic ablation of the specific reader(s) in the rbr-2 background should rescue the PVQ axon guidance phenotype. We found that the removal of nurf-1 (homolog of BPTF), but not of ing-3, jmjd-1/1.2 and lsy-13, significantly ameliorated the rbr-2 phenotype (Fig. 5A; Table S3), suggesting that NURF-1 is responsible for sensing and interpreting the increased levels of H3K4me3 in the rbr-2 mutant. Accordingly, loss of isw-1, homolog of the catalytic component of the NURF complex ISWI, also fully suppressed the rbr-2 axonal defects (Fig. 5B). Importantly, the level of wsp-1 mRNA was significantly reduced and resumed wild-type expression in rbr-2;nurf-1 and rbr-2;isw-1 double mutants in comparison with rbr-2 alone (Fig. 5C). These results strongly suggest that NURF-1 and ISW-1 are required for the inappropriate activation of wsp-1 transcription that occurs in the absence of rbr-2.

Fig. 2. RBR-2 is required during embryogenesis in the nervous system to ensure correct axon guidance. (A) Quantification of PVQ axonal cross-over defects in rbr-2(tm3141) mutants at L1 and adult stages. n>100, **P<0.01, ***P<0.001, n.s., not significant (one-way ANOVA followed by Tukey’s multiple-comparison test). (B) Expression of the Prbr-2::rbr-2::GFP translational reporter at embryonic and larval stages. Ventral nerve cord (VNC), muscle (m) and intestinal cells (i) are indicated. Anterior to the left. Scale bars: 10 μm. (C,D) Tissue- (C) and cell-specific (D) rescue analyses. Promoters used for transgenic rescue are: PF25B3.3, nervous system; Pmyo-3, body-wall muscles; Pdpy-7, hypodermis; Psra-6, PVQs. Statistical significance was calculated in relation to non-transgenic controls (values not shown) for each transgenic line. n>100, *P<0.05, **P<0.01, ***P<0.001, ****P<0.0001, n.s., not significant (one-way ANOVA followed by Tukey’s multiple-comparison test). At least two independent lines for each transgene were analyzed. Error bars represent standard error of proportion.
DISCUSSION
In this study, we show that the unique member of the KDM5 class of H3K4me3/2 demethylase RBR-2 is required for correct axon guidance in *C. elegans* through a mechanism involving the transcriptional regulation of *wsp-1*. Loss of RBR-2 leads to increased levels of H3K4me3 at the TSS of *wsp-1* and...

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Fig. 3. See next page for legend.
FIG. 3. RBR-2 modulates the expression of proteins regulating actin dynamics. (A) Top: Alignment of a part of the Jumonji C domain of human RBP2 with wild-type (WT) RBR-2 and with the catalytically inactive RBR-2DD (DD, Demethylase Dead). Asterisks mark two conserved amino acids in the iron-binding domain (HXD/EXH) of the JmC domain, modified in the RBR-2DD. Bottom: Quantified levels (relative to WT) of H3K4me3 in wild-type, rbr-2(tm3141) and rbr-2(tm3141) animals carrying a translational GFP fusion of catalytically inactive RBR-2 (rbr-2DD) or wild-type RBR-2 (rbr-2WT). H3 is used as loading control. (B) Quantification of PVQ axonal cross-over defects in rbr-2(tm3141) expressing a translational construct encoding for wild-type (rbr-2WT) and catalytically inactive (rbr-2DD) RBR-2. Data are representative of three independent lines for each transgene and statistical significance was calculated in relation to non-transgenic controls for each transgenic line. n=100, ***P<0.001, n.s., not significant (one-way ANOVA followed by Tukey’s multiple-comparison test). Error bars represent standard error of proportion. (C) WSP-1/WASP, VHE-1/WAVE and UNC-34/Ena/VASP integrate multiple signaling pathways to regulate actin dynamics. (D) mRNA levels of wsp-1, wve-1 and unc-34 in rbr-2(tm3141) embryos as measured by qPCR, using pmp-3 as internal control. (E) ChIP-qPCR assay for H3K4me3 on wsp-1, wve-1 and unc-34 TSS and intergenic regions in wild-type and rbr-2(tm3141) embryos. ChIPs with antibody against IgG served as negative controls. ChIP enrichments are normalized to input. (F) ChIP-qPCR assay for GFP on wsp-1, wve-1 and unc-34 TSS and intergenic regions in wild-type embryos carrying a translational GFP fusion of RBR-2 (WT-RBR-2::GFP). ChIPs with antibody against IgG served as negative controls. ChIP enrichments are normalized to input. In D-F, data are the average of three biological independent experiments and are expressed as means±e.m.; *P<0.05, **P<0.01 (Student’s t-test).

concomitant upregulation of its transcription, resulting in axonal defects. Furthermore, we show that the NURF complex is required for wsp-1 upregulation in the rbr-2 mutant and thus for translating the increased H3K4me3 signal into aberrant axon guidance.

Our analysis indicates that RBR-2 is required throughout development, as it functions in neurons that mature during embryonic (PVPs, PVQs) and postembryonic (HSNs) development. Projections of other neurons are unaffected in the rbr-2(tm3141) mutant. Thus, the function of rbr-2 appears to be restricted to a subset of neurons and its loss has a limited effect on global neuronal architecture. Genetic interaction assays suggest that RBR-2 acts in the most common pathways known to guide axon migration. The simplest explanation for this result is that the action of RBR-2 is required for regulating downstream targets of these pathways and cytoskeletal proteins are therefore potential candidates.

Using genetic and molecular approaches we identified WSP-1, an activator of the actin-nucleating complex Arp2/3, as a key protein required for the misguidance of PVQ axons in rbr-2 mutant worms. Briefly, we have demonstrated that: (1) RBR-2 is enriched at the wsp-1 promoter; (2) loss of rbr-2 results in increased levels of H3K4me3 at the wsp-1 TSS, in association with its transcriptional upregulation; (3) loss of wsp-1 in rbr-2 mutants fully restores normal PVQ axon guidance; (4) the overexpression of WSP-1 or its VCA domain lead to axon guidance defects similar to those observed in rbr-2(tm3141); and (5) inactivation of genes that encode proteins required for WSP-1 stability, localization and activation suppresses the rbr-2 mutant PVQ phenotype. The effect of loss of rbr-2 on the expression of wsp-1 is rather modest, as expected considering the cell-specific action of rbr-2 and the fact that we use full embryos to analyze the levels of wsp-1. Based on the well-established role of WSP-1 mammalian homologs, WASP/N-WASP, in positively regulating filopodia formation (Symons et al., 1996; Miki et al., 1998; Castellano et al., 1999; Kessels et al., 2011), we suggest that the upregulation of WSP-1 in C. elegans might lead to misregulated filopodia dynamics at the growth cone of the PVQ axons and to an aberrant response to different extracellular signals.

This hypothesis is corroborated by tissue-specific rescue experiments, pointing to a cell-autonomous role of RBR-2 and WSP-1 in the PVQs.

The identification of actin-remodeling genes as targets of RBR-2 is interesting in the context of neurological disorders. The H3K4me3 demethylase KDM5C is mutated in X-linked mental retardation (Iwase et al., 2007; Abidi et al., 2008; Adegbola et al., 2008; Gonçalves et al., 2014). Further, mutations in KDM5B have been reported in individuals with non-syndromic intellectual disability and autism spectrum disorders (Athanaskis et al., 2014; De Rubeis et al., 2014; Lissiof et al., 2014) and mutations in KDM5A have been linked to an autosomal recessive form of intellectual disability (Najmabadi et al., 2011). Similarly, several genes implicated in cytoskeleton dynamics are mutated in intellectual disability and autism spectrum disorders (Nadif Kasri and Van Aelst, 2008; Ba et al., 2013; Hu et al., 2014; Srivastava and Schwartz, 2014). Thus, our results help to establish a functional link between chromatin regulators and the actin-remodeling genes required for axon/dendrite formation and synaptic plasticity, which are processes compromised in neurodevelopmental and psychiatric disorders. However, considering the complexity underpinning intellectual disability disorders, it is possible that distinct mechanisms might act in cell-specific manners. Indeed, although ectopic expression of RBR-2 is able to rescue the axon migration defects of another pair of neurons (HSNs) observed in the rbr-2 mutant, loss of wsp-1 is not (Fig. S3), suggesting that RBR-2 might control axon guidance through different molecules/mechanisms. Thus, future analyses will be needed to molecularly dissect RBR-2 functions in other neuronal contexts.

Rescue analysis indicates that the H3K4me3 demethylase activity of RBR-2 is required for achieving correct axon guidance, providing a strong link between misregulated H3K4 methylation levels and the axonal phenotype. Several proteins, including those containing the evolutionarily conserved PHD domain, are known to bind specifically to H3K4me3, thereby translating the methylation status into a cellular process (reviewed by Sims and Reinberg, 2006; Vermeulen and Timmers, 2010). We showed that inactivation of nuref-1, coding for the large subunit of the NURF nucleosome-remodeling complex, reverts the rbr-2 phenotype. Consistently, the genetic ablation of isw-1, homolog of the ISWI catalytic component of the NURF complex, results in a similar outcome. In mammals, the NURF complex is recruited to chromatin through the interaction of its PHD domain with H3K4me3 where it catalyzes ATP-dependent nucleosome sliding at core promoters, facilitating transcription both in vitro and in vivo (Mizuguchi et al., 1997; Barak et al., 2003). Interestingly, the NURF complex is known to regulate some aspects of neuronal development (Lazzaro and Picketts, 2001; Barak et al., 2003), including neurite outgrowth (Barak et al., 2003). Thus, based on our results, we propose a model (Fig. 5D) in which increased levels of H3K4me3 at promoter regions occurring in the absence of RBR-2 could result in recruitment of the NURF complex to RBR-2 target genes, e.g. wsp-1, leading to changes in chromatin structure that favor transcription. In turn, increased levels of wsp-1 result in aberrant actin dynamics and axon guidance. The lack of a specific antibody against NURF-1 has prevented us from directly demonstrating increased levels of NURF-1 binding to RBR-2-regulated promoters in the rbr-2 mutant. However, the fact that inactivation of nuref-1 in rbr-2 mutant worms is accompanied by a reduction of wsp-1 expression to wild-type levels and by re-establishment of normal axon guidance strongly suggest that NURF-1 contributes to the overexpression of wsp-1 by binding to H3K4me3 at the TSS of wsp-1.
Fig. 4. See next page for legend.
In conclusion, our study establishes a precise role for epigenetic regulation in neuronal development by connecting the catalytic activity of RBR-2 to H3K4me3 levels and the NURF complex in controlling WSP-1 transcription. Future work will be aimed at identifying the molecules required for the cell-specific action of RBR-2 and the neuronal functions of other H3K4 regulators mutated in intellectual disability syndromes, with the hope of providing mechanistic insights that may be relevant for developing therapeutic treatments.

MATERIALS AND METHODS

Genetics and strains

C. elegans strains were cultured using standard growth conditions at 20°C on Escherichia coli OP50 (Brenner, 1974). Strains used were as follows: wild type Bristol: N2; rbr-2(tm3141) IV; rbr-2(tm1231) III; CX5334: oys14 [[(Para-6::GFP)+lin-15(+)] V; lin-13(n765) X; VH648: hda26(Podr-2::CFP; P ras-6::DsRed2) III; MUL1085: bws26([Pflp-1::GFP]+ [P rbf4::rol-6] (sa1006)]; EG1285: oxs121[(Punc-47::GFP)+lin-15(+)] X; lin-13(n765) X; NW1160: evk8282([Punc-129::GFP]+ [Prap-20(+)V]; ady-1(e2082) IV]; HTI1593: unc-119(ed3) III; NG324: wsp-1(gm324) IV; VC2706: wve-1 (ok3080) Vht72(rol-497); let-47(q82); gfs-68 (III); CB656: unc-34 (e566) V; CV2053: wsp-1(gm342)III H72(rol-497); let-47(q82); gfs-68 (UIII); VW198: rdc-42(gm386); mls14; dpy-10(c128) V; nck-1(ok694) VI; MT13649: nuf-1(jn2459) II; 1g-3(m2530) II; RB1433: T0611.0.4dly-13(ok1475) IV; jmd-1(jm3980) IV; jmd-1(jm3713) II; MT15795: isw-1(m2394) III; CX3198: sax-3(ky123) X; CX5000: slt-1 (eh5) X; OH1439: [vak-1(dx31) II]; oys14; CZ4111: vak-2(μ1) IV; NW1549: evf-2(ev658); evf-3(ev696) X; OH1487: hse-5(m472) III; AH205: sdn-1(zh10) X; NW1700: [pl-2(ev775) II; him-5(f1940) V]. The strains rphp6Exb(Pphk-1::GFP; P ras-6::RFP) and unc-5(e53) IV were generous gifts from Roger Pocock (Department of Anatomy and Developmental Biology, Monash University, VIC, Australia). Double mutant animals with specific genetic backgrounds were generated by standard crossing procedure. For a complete list of the transgenic strains generated for this study, see Table S4.

rbr-2 alleles

The allele rbr-2(tm3141) carries a deletion of 365 bp and an insertion of 8 bp, leading to a frameshift and a premature stop codon. The predicted protein, if translated, contains only the N-terminal portion of RBR-2, until the first PHD domain (Fig. 1A). The C. elegans mutant strain rbr-2(tm1231) carries an in-frame-deletion of 648 bp and the putative mutant transcript can give rise to a protein of 1261 amino acids lacking the first PHD domain (Fig. S1B). The alleles m3141 and m1231 were identified by the National BioResource Project (NBRP), Japan, and were backcrossed at least three times with wild-type animals before phenotypic analyses.

Generation of transgenic constructs

For the rbr-2::GFP construct, a 6081-bp fragment of rbr-2 (ZK593.4 in WormBase) containing the entire coding region and a 2540-bp promoter region were PCR-amplified from N2 genomic DNA. The resulting fragments were inserted in the multiple cloning sites of the pDONR pCR8 and the pDONR P4-P1R vectors, respectively (Gateway Cloning System, Life Technologies). The rbr-2 genomic sequence was amplified with the primers rbr-2_Fw (ATTCGTGACTGCTGCAAGA) and rbr-2_Rv (AATCATGGAAACTCAGGA) and the primers prom_rbr-2_Fw (GGGACACCTTTGTGAATAAAGTTTGAATc) and prom_rbr-2_Rv (GGGACTGCTTTTGGTACAAACTGTGTcc) were used to amplify the rbr-2 promoter region. Plasmids were constructed using MultiSite Gateway Three-Fragment Vector Construction Kit (Life Technologies). Tissue-specific promoters were cloned into the pDONR P4-P1R vector. The pDONR P2-PR3 vector containing the GFP sequence followed by unc-54 3‘UTR was a generous gift from Erik Jorgensen (Department of Biology, University of Utah, Salt Lake City, UT, USA). Final constructs were cloned into the pDEST R4-R3 destination vector.

To generate an integrated line, the vectors pDONR P4-P1R [Prbr-2], pDONR pCR8 [rbr-2] and pDONR P2-PR3 [GFP; unc-54 3‘UTR] were recombined with the destination vector pCG150 (Addgene), which includes the unc-119 rescue fragment into the vector backbone pDEST R4-R3.

For the rbr-2DD::GFP construct, the rbr-2::GFP construct was mutated using the QuickChange Site-Directed Mutagenesis Kit (Stratagene). Specifically, the DNA sequence was mutated so that the histidine at position 514 (H514) and the glutamic acid at position 516 (E516) were changed to alanine. Primers rbr-2_H514E516A_FWD (GTTCGCTGGCGCACCGGATGACTTTGGA) and rbr-2_H514E516A_Rv2 (GTCACAGTATCGGCAGACAGAAG) were used for the site-directed mutagenesis to generate the rbr-2 catalytic dead mutant construct.

The vectors pZ4 [PF25B3.3::wsp-1a cDNA] and pZ3 [Psy-3::wsp-1a cDNA] were generous gifts from Terry Kubesium (Department of Biology, York University, Toronto, ON, Canada).

The cDNA encoding the VCA domain of WSP-1 [9108-9579 of the wsp-1 gene (C07G1.4a in WormBase)] was PCR-amplified from pZ4 with the primers VCA_GW-fw (aaacatatgTCAGGAGCCGGAGGACCT) and VCA_GW-rv (ATCTGACCATTCATTTTTGTCAT). Of note, the distribution of PVQ defects between wild type and some lines. Of note, the distribution of PVQ defects between wild type and some lines. Of note, the distribution of PVQ defects between wild type and some lines. Of note, the distribution of PVQ defects between wild type and some lines.
Imager M2) and MicroManager software (version 1.4.11). All pictures were exported in preparation for printing using Photoshop (Adobe).

Dil staining of amphid and phasmid neurons
Young adult hermaphrodites cultured at 25°C were transferred from a plate into an eppendorf tube with 1 ml M9, spinned down at 1500 g and washed twice. Worms were resuspended in 1 ml M9 and 5 μl were added from a stock dye solution containing 2 mg ml⁻¹ Dil (Molecular Probes, catalog # D-282) in dimethyl formamide. Eppendorf tubes were incubated wrapped in foil on a slow shaker for 3 h. Worms were spinndown at 1500 g and washed twice with M9 before being analyzed by fluorescence microscopy using the Texas Red filter, as described above.

Chromatin immunoprecipitation (ChIP)
Gravid hermaphrodites cultured at 25°C were treated with hypochlorite solution and embryos were flash-frozen in liquid nitrogen and stored at −80°C before chromatin immunoprecipitation. ChIP was performed with a protocol modified from that of Kolasinska-Zwierz et al. (2009). Chromatin was disrupted by sonication using a Diagenode Bioruptor sonicator UCD-300 to obtain fragments of 200-500 bp in size. Suitable amounts of chromatin were incubated with the following antibodies overnight: polyclonal anti-H3K4me3 (Abcam, ab8580; 1:1000), polyclonal anti-IgG (Sigma, I8140; 1:1000), polyclonal anti-GFP (Abcam, ab290; 1:500). Immunoprecipitated complexes were recovered on magnetic Protein G Dynabeads (Invitrogen) and, after extensive washes, DNA was isolated by reverse cross-linking and purification using the QIAquick PCR Purification Kit (Qiagen). For ChIP-qPCR, immunoprecipitated DNA and input were quantified by real-time qPCR as described below. The measures were normalized to the input. All reactions were performed in duplicate, in three independent experiments.

Real-time quantitative PCR (RT-qPCR)
Late embryos were obtained by hypochlorite treatment of gravid hermaphrodites followed by 6-8 h of suspension of the embryos in M9 at room temperature. Embryos were flash-frozen in liquid nitrogen and stored at −80°C before use.

Fig. 5. Loss of the NURF complex rescues the PVQ axon guidance defects associated with rbr-2(tm3141) and restores correct level of wsp-1 mRNA.
(A) Quantification of PVQ axonal cross-over defects in rbr-2(tm3141) and double mutants with PHD-containing proteins nurf-1(n4295), ing-3(tm2530), lsy-13(ok1475), jmd-1.1(tm3980) and jmd-1.2(tm3713). n>100, **P<0.01, n.s., not significant (one-way ANOVA followed by Tukey’s multiple-comparison test). Error bars represent standard error of proportion. (B) Quantification of PVQ axonal cross-over defects in rbr-2(tm3141), isw-1(n3294) and double mutant. n>100, **P<0.01 (one-way ANOVA followed by Tukey’s multiple-comparison test). Error bars represent standard error of proportion. (C) mRNA levels of wsp-1 in rbr-2(tm3141), rbr-2(tm3141);nurfl(n4295) and rbr-2(tm3141);isw-1(n3294) embryos as measured by qPCR, using pmp-3 as internal control. Data are the average of three biological independent experiments and are expressed as means±s.e.m. *P<0.05 (Student’s t-test). (D) Model of the mechanism of action of RBR-2 as a negative regulator of wsp-1 transcription. In wild type, RBR-2 regulates the level of H3K4me3 at wsp-1 TSS and secures correct axon guidance. Loss of rbr-2 leads to increased levels of H3K4me3 and of NURF complex recruitment at wsp-1 TSS and consequently to overexpression of WSP-1, resulting in aberrant actin remodeling and axon guidance.
–80°C before RNA extraction. Total RNA was isolated using TRIzol reagent (Life Technologies) and the RNeasy Mini Kit (Qiagen). cDNA was synthesized using oligo(dT)16 primers and reagents from the TaqMan Reverse Transcription Kit (Applied Biosystems). qPCR was performed using Maxima SYBR Green/ROX qPCR Master Mix 2× (Thermo Scientific). Reverse Transcription Kit (Applied Biosystems).

qPCR was performed to be unusually stable (Zhang et al., 2012), with similar results. All reactions were performed in duplicate, in three independent experiments.

**Western blot analysis**

Total protein extracts were prepared from L4 hermaphrodites grown on OP50 at 25°C. Protein concentration was estimated using the modified micro-Lowry assay and equal amounts of protein were loaded. The following antibodies were used: polyclonal anti-H3K4me3 (Abcam, ab8580, lot GR152455-1; 1:5000); polyclonal anti-H3 (Abcam, ab1791, lot GR9204-1; 1:30,000); peroxidase-labeled anti-rabbit secondary antibody (Vector; 1:10,000). Western blots were quantified using the ImageJ program (National Institutes of Health).

**Statistical analyses**

All phenotypes were scored as percentages of defective animals and results are shown with error bars representing the standard error of proportion. Statistical analyses were performed in GraphPad Prism 6 using Fisher’s exact test, for pairwise comparisons, or one-way ANOVA followed by Bonferroni post-hoc test. Differences with a P-value <0.05 were considered significant.

**Competing interests**

The authors declare no competing or financial interests.

**Author contributions**

L.M. carried out all the experimental work with the contribution of Y.C.L., J.V. and A.R. A.E.S. and L.M. designed the experiments, analyzed the data and wrote the manuscript.

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**Supplementary information**

Supplementary information available online at http://dev.biologists.org/lookup/suppl; doi:10.1242/dev.132985/DC1

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