G9a and ZNF644 Physically Associate to Suppress Progenitor Gene Expression during Neurogenesis

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http://dx.doi.org/10.1016/j.stemcr.2016.06.012

SUMMARY

Proliferating progenitor cells undergo changes in competency to give rise to post-mitotic progeny of specialized function. These cell-fate transitions typically involve dynamic regulation of gene expression by histone methyltransferase (HMT) complexes. However, the composition, roles, and regulation of these assemblies in regulating cell-fate decisions in vivo are poorly understood. Using unbiased affinity purification and mass spectrometry, we identified the uncharacterized C2H2-like zinc finger protein ZNF644 as a G9a/GLP-interacting protein and co-regulator of histone methylation. In zebrafish, functional characterization of ZNF644 orthologs, znf644a and znf644b, revealed complementary roles in regulating G9a/H3K9me2-mediated gene silencing during neurogenesis. The non-overlapping requirements for znf644a and znf644b during retinal differentiation demarcate critical aspects of retinal differentiation programs regulated by differential G9a/ZNF644 associations, such as transitioning proliferating progenitor cells toward differentiation. Collectively, our data point to ZNF644 as a critical co-regulator of G9a/H3K9me2-mediated gene silencing during neuronal differentiation.

INTRODUCTION

Alterations in histone methylation instruct developmental gene-expression programs that enable proliferating progenitor cells to exit the cell cycle and differentiate. These changes are mediated by conserved, multiprotein macromolecules that “write” and “read” histone methylation marks, such as Set1/COMPASS-like complexes (Shilatifard, 2012), Polycomb repressor complexes (Margueron and Reinberg, 2011), and assemblies containing the histone methyltransferases (HMTs) G9a and GLP (Shinkai and Tachibana, 2011). Developmental regulation of specific genomic loci involves complex physical interactions involving tissue-specific transcription factors (TFs), non-coding RNA, and other co-factors (Guttman et al., 2011). Histone methylation in pluripotency-related gene regulation has been characterized extensively (Watanabe et al., 2013), yet the composition of relevant HMT complexes and, specifically, the identity of physically associated co-regulators that modulate activity during cellular differentiation are incompletely defined.

G9a/EHMT2 and GLP/EHMT1 are responsible for dimethylated H3K9 (H3K9me2) in transcriptionally repressed euchromatin (Tachibana et al., 2005) and are essential for cell differentiation during embryogenesis (Shinkai and Tachibana, 2011). In embryonic stem cells (ESCs), G9a/GLP facilitate the long-term silencing of pluripotency-associated genes (Tachibana et al., 2008). In hematopoietic stem and progenitor cells (HSPCs), inhibition of G9a/GLP delays lineage commitment and prevents the formation of large H3K9me2 chromatin territories (Chen et al., 2012). In neural contexts, loss of G9a or GLP in the mouse forebrain reactivates neural progenitor gene expression, leading to cognitive and adaptive behavioral defects (Schaefer et al., 2009). Disruption of the GLP/EHMT1 gene in humans is associated with congenital intellectual disability (Kleefstra et al., 2005), and heterozygous GLP/EHMT1 knockout mice exhibit behavioral and neurodevelopmental abnormalities (Balemans et al., 2010, 2014). Although G9a/GLP-associated proteins have been reported (Bian et al., 2015; Maier et al., 2015; Ueda et al., 2006), their precise contributions to G9a/GLP-mediated neural differentiation are largely unknown.

Retinal progenitor cells (RPCs) are proliferative multipotent cells that produce differentiated cells in an evolutionary conserved birth order (Bassett and Wallace, 2012; Cepko et al., 1996). RPC proliferation and differentiation must be closely integrated and coordinated with eye growth for proper morphology and structure (Green et al., 2003; Wong et al., 2015), with impaired proliferative capacity resulting in microphthalmia, degeneration, and visual impairment (Levine and Green, 2004). RPC differentiation requires G9a/H3K9me2-mediated suppression of genes that sustain a proliferative multipotent state (Katoh...
Additional evidence suggested that the functions of functional cooperativity and physical binding to G9a, neurons, respectively, both of which were dependent on viability and ensuring the proper differentiation of retinal involved in chromatin and transcriptional regulation (Mak et al., 2012), including Vsx2/Chx10, a TF uniformly expressed in fully multipotent RPCs (Burmeister et al., 1996; Duffy et al., 2005; Vitorino et al., 2009), and Ccnd1/Cyclin D1, the predominant D-cyclin in the developing retina (Barton and Levine, 2008). Co-regulators that facilitate G9a/H3K9me2-mediated silencing during neurogenesis are unknown.

High-grade myopia involves progressive axial elongation of the eye that predisposes to degeneration and blindness. Genetic factors linked to high-grade myopia (Hawthorne and Young, 2013) include mutations in the C2H2-like zinc finger (ZF) protein ZNF644, which segregates with autosomal dominant inheritance (Shi et al., 2011). These findings suggest a role for ZNF644 in maintaining proper eye morphology and/or growth, yet its function in neural contexts is currently uncharacterized.

To better understand the molecular basis of histone methylation, we applied a lentiviral-based affinity purification and mass spectrometry (AP-MS) approach to isolate protein complexes that “write” and/or “read” histone methylation. We found that ZNF644 physically interacts with G9a and GLP and serves as a co-regulator of H3K9me2. By characterizing zebrafish ZNF644 orthologs znf644a and znf644b, we pinpointed roles for these genes in regulating G9a/H3K9me2-mediated silencing in neural progenitor cells. Loss-of-function experiments demonstrated roles for znf644a and znf644b in maintaining cell viability and ensuring the proper differentiation of retinal neurons, respectively, both of which were dependent on functional cooperativity and physical binding to G9a. Additional evidence suggested that the functions of g9a and znf644a are recapitulated in midbrain progenitor cells, suggesting a common gene-silencing complex mediating differentiation in distinct neuronal progenitor populations. Collectively, our findings highlight G9a-ZNF644 as a critical regulator of gene silencing and cell-fate transitions during neural differentiation.

RESULTS

ZNF644 Is a Co-regulator of Histone Methylation

We previously described a systematic AP-MS approach, based on stable lentivirus-based expression of epitope-tagged proteins, to characterize stable protein complexes involved in chromatin and transcriptional regulation (Mak et al., 2010; Ni et al., 2011). As part of a larger study (Marcon et al., 2014), we targeted human HMT protein complexes linked to histone methylation during development. These included confirmed and putative homologs of Polycomb, Set1/COMPASS, and G9a/GLP-containing protein complexes (Table S1). We created a scored physical interaction map encompassing components of histone methylation-related protein complexes with emphasis on regulators of developmental gene expression. Our dataset comprised 33 replicate AP-MS analyses of 25 histone methylation-related and eight transcription-related proteins.

In addition to accurately recapitulating established transcriptional assemblies, such as Mediator (Conaway et al., 2005), RNA polymerase II (RNAPII) (Ni et al., 2011), and cleavage and polyadenylation-specific termination factor (Kaufmann et al., 2004) (Figure 1 and Table S1), we identified dozens of previously unknown associations among a filtered network of 573 interactions. Pairwise hierarchical clustering of the highly correlated interaction profiles of the different bait proteins revealed components residing within the same complex(es) (Figure 1B). Most (312 of 428, 72.9%) of the annotated interactions detected were corroborated in public protein-protein interaction repositories such as BioGRID (Chatr-Aryamontri et al., 2015) and iRefWeb (Turner et al., 2010), suggesting a high-confidence interaction network.

In addition to its known dimerization partner GLP, affinity-tagged G9a reproducibly co-purified the C2H2-like ZF proteins, WIZ and ZNF644 (Figure 1C). WIZ was previously reported to interact with G9a/GLP in ESCs via association of the SET domains of G9a/GLP with an atypical C2H2-like ZF motif in WIZ (Ueda et al., 2006). ZNF644 contains a highly similar atypical C2H2-like ZF motif, and GFP-tagged full-length ZNF644 as well as an isoform comprising the atypical ZF region (ZF8) was sufficient to co-purify G9a/GLP and WIZ (Figure 1D). In addition, mutations in the putative zinc ion-chelating residues of the atypical ZF motif (C1263A or H1283A) abolished complex formation (Figure 1D). The shortened ZF8 region of ZNF644, expressed as a recombinant glutathione S-transferase (GST)-fusion protein, efficiently pulled down recombinant His-tagged G9a SET domain homodimer, but not the C1263A or H1283A mutants (Figure 1E). Small hairpin RNA (shRNA)-mediated knockdown of either G9a or ZNF644 significantly and specifically reduced H3K9me2 (Figures 1F and S1A). Collectively, these data demonstrate that ZNF644 binds to G9a/GLP and co-regulates H3K9me2.
Figure 1. ZNF644 Is a G9a/GLP-Interacting Protein and Co-regulator of H3K9me2

(A) Physical interaction map of histone methylation-related complexes. Nodes represent proteins identified by AP-MS, black edges represent high-confidence interactions, and known interactions are in green.

(B) Hierarchical clustering of bait interaction profile similarity. Colored boxes indicate proteins known to reside in the same complex.

(C) (Top) Protein domain architecture of WIZ and ZNF644. (Bottom) Sequence alignment of atypical C2H2-like ZF motifs. Putative zinc ion-chelating residues are circled.

(D) Summary of AP-MS analyses showing average peptide spectral counts of GFP-tagged forms of full-length WT ZNF644, the C1263A or H1283A point mutants, or the ZF8-containing C-terminal region only.

(E) Western blot analysis of pulldown assays using GST-tagged WT, C1263A, or H1283A versions of the ZF8 domain of ZNF644 with the His-tagged G9a SET domain.

(F) Western blot assays monitoring G9a, H3K9me2, H3K9me3, histone H3, and GAPDH levels in HEK293 cell lines expressing shRNAs targeting a non-silencing control, G9a, or ZNF644.
G9a/GLP. Indeed, AP-MS assays based on GFP-tagged zebrafish Znf644a or Znf644b proteins in HEK293 cells confirmed the ability to interact with G9a/GLP and WIZ (data not shown). Throughout embryogenesis, znf644a and znf644b mRNAs were expressed in most tissues, with slightly higher levels in the neuroepithelium of the retina and midbrain at 24 hr post fertilization (hpf) that was reduced by 48 hpf (Figures 2B and S1B–S1D). Since the 24-hpf retina is largely composed of proliferating RPCs on the verge of differentiation (Agathocleous and Harris, 2009; Li et al., 2000), we analyzed the developing retina to investigate potential functional cooperativity with g9a.

Figure 2. Zebrafish znf644 Paralogs Are Specifically Expressed In and Regulate the Development of Retinal and Midbrain Progenitor Cells
(A) Conservation of C2H2-like ZF motifs between human and zebrafish ZNF644 genes.
(B) Representative lateral views, retinal cross-sections, and midbrain cross-sections of whole-mount in situ hybridization (WISH) assays monitoring znf644a or znf644b expression at 24 hpf (n > 15 in each group). Arrows denote the inner part of the central retinal epithelium.
(C) Lateral views of g9a (20/32), znf644a (10/18), or znf644b (14/21) morphant embryos compared with WT (21/21). Dashed circles denote 2D area of the WT retina. Arrowheads denote the dorsal hindbrain region.
(D) Quantitation (mean ± SD) of retinal 2D cross-sectional area (μm²) of znf644a, znf644b, or g9a morphants relative to WT embryos. Error bars represent SD. *p < 0.005, Student’s t test.
(E) RT-PCR assays monitoring unspliced (unsp) and spliced (spl) znf644a or znf644b transcripts at 24 hpf.
The Forming Retina and Midbrain Are Disrupted in znf644a or znf644b Morphants

Loss-of-function studies using g9a-MO were consistent with previous reports (Rai et al., 2010) in that these morphants showed severe morphological abnormalities in a dose-dependent manner (Figures S2A and S2B). Although overall far less severe, splice-block MOs targeted against znf644a (znf644a-MO) or znf644b (znf644b-MO) showed significant dose-dependent reductions in eye size (microphthalmia) and disrupted midbrain morphology (Figure 2C). Mismatch control MOs showed no discernible phenotypes (data not shown), supporting targeting specificity. MO injection doses eliciting a ~50% reduction in retinal cross-sectional area (Figure 2D) corresponded to a 57% reduction in znf644a, nearly complete splicing disruption of znf644b, and ~35% drop in g9a (Figure 2E and data not shown). Co-injection of cognate mRNA fully rescued the znf644a morphant defect and, to a lesser but still appreciable degree, znf644b morphants (Figures S2C and S2D), further confirming targeting specificity. These data highlight the developing retina and midbrain as neural structures dependent on coordinate g9a/znf644a/znf644b function.

Znf644a and znf644b Regulate Gene Silencing in the Retina via H3K9me2

To assess the extent to which znf644a and znf644b affect H3K9me2 in vivo, we isolated cells from the head regions (heterogeneous mixture of retinal, brain, and other cell types) of znf644a and znf644b morphant embryos at 48 hpf and mapped the positioning of H3K9me2 marks by chromatin immunoprecipitation (ChIP) sequencing (GEO: GSE63225). H3K9me2 peaks near transcriptional start sites (TSSs), which were prominent in control embryos, were largely depleted in znf644a and znf644b morphants (Figures 3A and 3B; Table S1), consistent with roles in the global regulation of H3K9me2.

Although some affected H3K9me2 peaks localized near genes implicated in neural development (Table S1), sample heterogeneity likely precluded sampling of specific loci relevant to neural progenitor cell subpopulations. We therefore performed targeted investigations of the vsx2 and ccnd1 genes, two key regulators of multipotency and proliferation reported to be suppressed by G9a/H3K9me2 in mouse RPCs (Katoh et al., 2012). ChIP-PCR assays of zebrafish vsx2 and ccnd1 revealed reductions in H3K9me2 in the proximal promoters of vsx2 and ccnd1 in znf644a and znf644b morphant embryos (Figure 3C). In addition, these defects in H3K9me2 coincided with aberrantly mislocalized vsx2 and ccnd1 expression throughout the central retina at 48 hpf (Figure 3D), with similar changes observed in g9a morphant retinas (Figures S3D and S3E). Collectively, these data demonstrate shared roles for znf644a and znf644b in H3K9me2-mediated gene silencing in RPCs.

znf644a and znf644b Suppress Neural Progenitor Gene Expression and Proliferation

To distinguish whether mislocalized expression of vsx2 and ccnd1 simply represented failed gene silencing during retinal differentiation or a maintained proliferative progenitor identity, we examined additional multipotency (rx3, otx2) and proneural (ath5) markers (Stenkamp, 2007). Strikingly, znf644a morphants exhibited elevated and persistent expression of rx3, ath5, and otx2 (Figure 3D), consistent with continued expression of multiple genes that typify the RPC expression program. In contrast, znf644b morphants showed minimal changes in rx3 and ath5 expression, but elevated otx2 expression (Figure 3D), suggesting the deregulated expression of only a subset of RPC-specific genes. These changes in gene expression were rescued by co-injection of cognate mRNAs (Figures S3A and S3B). Together, these data suggest at least partially non-overlapping requirements for znf644a and znf644b in suppressing the RPC gene-expression program.

We assessed the proliferative state of retinal cells in the morphants by quantitating 5'-bromo-2-deoxyuridine (BrdU)+ (S phase), phospho-histone H3 (pH3)+ (mitotic), and proliferating cell nuclear antigen (PCNA)+ cells (cycling cells). While the central retina of wild-type (WT) embryos contained few BrdU+ cells (0.23 ± 0.04 cells/μm²) and low PCNA expression at 48 hpf, there was an order-of-magnitude increase in the number of central BrdU+ cells (1.86 ± 0.29 cells/μm²) and markedly increased PCNA levels in znf644a morphants (Figures 4A and 4B). Likewise, znf644b morphants showed a significant (~3.5-fold) increase in BrdU+ cells (0.81 ± 0.28 cells/μm²) and elevated PCNA (Figures 4A and 4B); however, the number of pH3+ cells was largely unchanged in the znf644a or znf644b morphant central retina, although a decreasing trend in znf644a morphants was noted (Figures 4C and 4D). In contrast, a substantial number of pH3+ cells persisted in the central retina at 72 hpf in znf644a morphants, but not znf644b morphants, compared with controls, where none were observed (Figures 4C–4E). Likewise, g9a morphant retinas exhibited similar increases in the number of proliferating cells that persisted in the central retina into late development (Figures S4A and S4B). Collectively, these results establish roles for znf644a and znf644b in suppressing retinal cell proliferation.

CRISPR/Cas9 gene editing provided additional confirmation of znf644b morpholino target specificity (Figure S5A). Compared with phenotypically normal F1 embryos that were heterozygous for targeted mutations, embryos with two mutant (e.g., frameshift) alleles showed significantly reduced brain and retinal growth and elevated expression of ccnd1 (Figure S5B), coinciding with the presence of pH3+ cycling cells in the central retina at 48 hpf.
Thus, results from an independent genetic loss-of-function approach are consistent with the phenotype observed in the znf644b morphants. Consistent with the prolonged proliferation and maintained RPC gene-expression program, the central retinas of znf644a morphants lacked various types of differentiated A B C D

Figure 3. znf644a and znf644b Are Required for H3K9me2-Mediated Gene Silencing in the Forming Retina
(A) Frequency and distance from TSS at which H3K9me2 peaks were identified in WT, znf644a, and znf644b morphants.
(B) Venn diagram illustrating the number and overlap of H3K9me2 peaks at 48 hpf in znf644a or znf644b morphants or WT embryos.
(C) ChIP-PCR assays monitoring H3K9me2 levels near the TSSs of vsx2 or ccnd1 genes at 48 hpf.
(D) WISH assays monitoring expression of progenitor or proneural genes in WT, znf644a, or znf644b morphant retinas. Arrows highlight expression domains.

(Figure S5C). Thus, results from an independent genetic loss-of-function approach are consistent with the phenotype observed in the znf644b morphants.
Figure 4. znf644a and znf644b Morphant Retinas Are Composed of Distinct Populations of Retinal Cells with Differing Characteristics

(A) Immunostaining of retinal cross-sections monitoring BrdU incorporation and PCNA+ cells at 48 hpf in WT, znf644a, and znf644b morphants.

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neurons, as late as 48–72 hpf, including protein kinase C (PKC)\(^*\) bipolar neurons, GABA\(^*\) amacrine cells, and Zpr1\(^*\) cone photoreceptor cells (Figure 4F; data not shown). A similar lack of differentiated retinal neurons was likewise observed in g9a morphants (Figure S4C). In contrast, the persistent expression of vsx2 and ccnd1 notwithstanding, znf644b morphant retinas, as a population, formed differentiated neurons by 48–72 hpf (Figure 4F and data not shown). These data imply distinct regulatory requirements for znf644a and znf644b vis-à-vis cellular differentiation in the retina.

**znf644a Morphant Retinas Preserve Characteristics of Proliferative Progenitors**

To further characterize the znf644a morphant retina, we assessed the temporal emergence of RGCs and amacrine cells, which are among the first-born differentiated retinal cells (Agathocleous and Harris, 2009; Li et al., 2000). Isl2b\(^*\) RGCs and Pax6\(^*\) amacrine cells, which are normally amply visible by 48 hpf, were not detected in znf644a morphant retinas until 72 hpf (Figure 5A), consistent with delayed onset of differentiation. Rather, the znf644a morphant retinas were also largely composed of Vsx2\(^*\) cells at 48 hpf, suggestive of a maintained RPC identity (Figure 5A). We also detected H3K9me2\(^*\) nuclei among differentiated cells of the ganglion and inner nuclear layer at 72 hpf (Figure 5A), reflecting global increases in H3K9me2 during differentiation. However, znf644a morphant retinas lacked H3K9me2\(^*\) nuclei even at 72 hpf (Figure 5A), suggesting that H3K9me2\(^*\) nuclei emerge downstream of cell-cycle exit within differentiated retinal neuron populations.

To confirm an RPC-specific role, we transplanted znf644a morphant donor cells (membrane-GFP\(^*\)) into WT host embryos, thereby generating chimeric retinas. We observed a significant (21.6\%) increase in pH3\(^*\) cells in donor cells (76 cells in total; n = 2) compared with an absence of pH3\(^*\) cells in embryos with control donor cells (94 cells in total; n = 2) (Figures S4D and S4E), indicating an RPC-autonomous role of znf644a.

**Impaired Viability of Differentiated Neurons in znf644b Morphant Retinas**

Misexpression of proliferation genes during differentiation, as observed in znf644b morphants, is known to cause apoptosis (Bonda et al., 2009; Busser et al., 1998; Kuan et al., 2004; Zhu et al., 2004). Indeed, we observed increased cleaved Caspase3 in differentiated regions of znf644b morphant retinas at 72 hpf (28.7 ± 10.3) relative to WT retinas (2.0 ± 0.0; Figures 5B and 5C). High levels of apoptotic cells were seen in g9a morphant retinas not only at 72 hpf (67.0 ± 10.8), but also at earlier time points (Figures 5B and 5C). Cleaved Caspase3 was rescued in znf644b and g9a morphants by co-injecting a morpholino targeting p53 (p53-MO), suggesting intrinsic apoptosis pathway activation (Figures 5B and 5C). We also found evidence of concurrent expression of Pax6, a marker for differentiated amacrine and ganglion cells, and PCNA in znf644b morphant retinal cells at 56 hpf, coinciding with the onset of excessive cell death (Figure S5E). Overall, these data suggest znf644b function is required to maintain the long-term viability of differentiated retinal cells, at least in part through H3K9me2-dependent repression of progenitor and/or cell-cycle genes.

**Extensive Functional Cooperativity between g9a, znf644a, and znf644b**

We used a genetic cooperativity approach wherein subphenotypic doses of MOs that individually do not yield observable developmental defects were co-injected to reveal potentially additive or synergistic effects. Co-injected subphenotypic g9a-MO and znf644b-MO gave rise to a g9a morphant-like phenotype with a substantial number of Caspase3\(^*\) apoptotic cells emerging early in the central retina (Figure 6A). Similar cooperativity occurred in mosaic retinas derived from in vivo cell transplantation assays, where a higher incidence (14.6\%) of apoptosis was observed among retinal cells derived from co-injected subphenotypic g9a-MO plus znf644b-MO GFP\(^*\) donor cells (123 cells in total; n = 4) (Figures S4F–S4H), compared with only 0.56\% subphenotypic g9a-MO GFP\(^*\) donor cells (142 cells in total; n = 2) or 3.58\% subphenotypic znf644b-MO GFP\(^*\) donor cells (139 cells in total; n = 3). These data suggest a cell-autonomous role for g9a and znf644b in maintaining retinal cell viability.

The number of PCNA\(^*\) cells localized to the peripheral ciliary marginal zones (CMZ) in control retinas was 64.4 ± 10.2 cells per section (n = 5), with none seen in

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(B) Quantitation of BrdU\(^*\) cells/\(\mu m^2\) in central retina of WT, znf644a, or znf644b morphants at 48 hpf (n = 3 for each group). Error bars represent SD. *p < 0.05, Student’s t test.

(C–E) Immunostaining of pH3 expression in retinal cross-sections at 48 and 72 hpf in WT, and znf644a morphants (C). Quantitation of pH3\(^*\) cells/\(\mu m^2\) in the central retina of WT, znf644a, or znf644b morphants at (D) 48 hpf (n = 5 for each group) and (E) 72 hpf (n = 2 for each group). Quantitation represented as mean ± SD.

(F) Immunostaining monitoring marker protein expression in retinal cross-sections. At 72 hpf, amacrine cells express GABA, bipolar neurons express PKC, and cone photoreceptors express Zpr1.

White arrows highlight populations of proliferative or differentiated cells.
Figure 5. *znf644a* and *znf644b* Morphant Retinas Exhibit Distinct Cellular Defects

(A) Immunostaining of retinal cross-sections of Vsx2 (red), Pax6 (red), Isl2b (green), or H3K9me2 (red) in WT or *znf644a* morphant embryos at indicated time points. Vsx2 expression at 48 hpf marks proliferating RPCs in the CMZ (gray arrows) and in a subpopulation of bipolar neurons in the central retina (yellow arrows). Pax6 expression marks amacrine cells and a subset of RGCs, and Isl2b expression marks a subset of RGCs. Differentiated retinal neurons with H3K9me2+ nuclei are highlighted.

(B and C) Immunostaining monitoring cleaved Caspase3 (red) in retinal cross-sections from WT, *g9a*, or *znf644b* morphant embryos at 48 or 72 hpf (B). Yellow arrows highlight Caspase3+ apoptotic cells. (C) Quantitation of Caspase3+ cells in WT, *znf644a* morphant, or *znf644b* morphant retinas with or without co-injection of p53-MO at 72 hpf. Quantitation represented as mean ± SD (n=3 for each group). *p < 0.05, **p < 0.005, ***p < 0.0005, Student’s t test.
### Individual subphenotypic MO doses

|                      | Combined subphenotypic MO doses |
|----------------------|---------------------------------|
| g9a-MO (2ng)         | g9a-MO (2ng) + znf644b-MO (10ng)|
| znf644a-MO (15ng)   | znf644a-MO (15ng) + znf644b-MO (10ng) |

### Combined subphenotypic MO doses

|                      | Combined subphenotypic MO doses |
|----------------------|---------------------------------|
| g9a-MO (2ng) + znf644a-MO (15ng) | g9a-MO (2ng) + znf644b-MO (10ng) |

### Individual subphenotypic MO doses

|                      | Combined subphenotypic MO doses |
|----------------------|---------------------------------|
| g9a-MO (2ng)         | g9a-MO (2ng) + znf644b-MO (10ng)|
| znf644a-MO (10ng)   | znf644a-MO (10ng) + znf644b-MO (10ng) |

### Combined subphenotypic MO doses

|                      | Combined subphenotypic MO doses |
|----------------------|---------------------------------|
| g9a-MO (2ng) + znf644a-MO (15ng) | g9a-MO (2ng) + znf644b-MO (10ng) |

### Individual subphenotypic MO doses

|                      | Combined subphenotypic MO doses |
|----------------------|---------------------------------|
| g9a-MO (2ng)         | g9a-MO (2ng) + znf644b-MO (10ng)|
| znf644a-MO (15ng)   | znf644a-MO (15ng) + znf644b-MO (10ng) |

### Combined subphenotypic MO doses

|                      | Combined subphenotypic MO doses |
|----------------------|---------------------------------|
| g9a-MO (2ng) + znf644a-MO (15ng) | g9a-MO (2ng) + znf644b-MO (10ng) |

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the central retina in control or single MO-injected embryos at subphenotypic doses. However, embryos co-injected with subphenotypic doses of znf644b-MO and g9a-MO showed modestly increased PCNA+ cell numbers (74.0 ± 19.7 cells/section; n = 2), but only at the retinal periphery, while no change was observed in the central retina, which correlated with a modest effect in vsx2 expression (Figure 6B). In contrast, co-injection of znf644a-MO and g9a-MO led to a synergistic increase in PCNA+ cells (129.0 ± 22.6 cells/section; n = 3) throughout the central retina at 48 hpf (Figure 6B), comparable with full znf644a-MO morphant retinas (Figure 4B), yet without any observable effects on cell viability (Figure 6A) and with minimal changes in vsx2 expression (Figure 6B). In addition, compared with controls, where pH3+ cells (17.0 ± 1.0; n = 3) and BrdU+ cells (15.3 ± 6.4; n = 3) were confined primarily to the marginal zone, co-injected subphenotypic doses of g9a-MO and znf644b-MO resulted in marked increased levels in pH3+ (50.6 ± 1.5; n = 3) and BrdU+ cells (31.0 ± 12.5; n = 3) and mislocalized ccd1 expression across the entire central retina and margin (Figure S6B). Co-injection of g9a-MO and znf644a-MO resulted in similar overall changes (Figure S6B). These data suggest that extensive functional cooperativity between g9a and znf644b or znf644a maintains the viability of differentiated retinal cells or suppresses a prolonged proliferative state in the retina, respectively.

The misexpression of vsx2 and ccd1 and prolonged retinal cell proliferation in znf644a and znf644b morphant retinas at 48 hpf (Figures 3A, 4A, and 4B) led us to investigate the role of functional cooperatively among these paralogs in suppressing RPC identity. As seen in znf644a and znf644b morphants (Figure 3A), co-injection of subphenotypic doses of znf644a-MO plus znf644b-MO resulted in mislocalized vsx2 and ccd1 expression throughout the central retina at 48 hpf (Figures 6B and S6A), consistent with functional cooperativity. In addition, increased numbers of PCNA+ (129.3 ± 25.2 cells/section; n = 6), pH3+ (33.0 ± 6.2; n = 3), and BrdU+ (42.0 ± 9.8; n = 3) cells were evident throughout the central retina (Figures 6B and S6B), suggesting that znf644a and znf644b work together to suppress RPC gene expression and retinal cell proliferation.

Human ZNF644 Rescues znf644a and znf644b Morphant Phenotypes

To assess the conservation of human and zebrafish ZNF644 orthologs, we performed functional complementation assays. Full-length native version of human ZNF644 rescued the retinal defects observed in both the znf644a and znf644b morphant retinas; specifically, mislocalized expression of vsx2 and elevated apoptosis in znf644b morphants (Figures 6D), and the mislocalized expression of vsx2, prolonged cellular proliferation, and lack of differentiated neurons in znf644a morphants (Figures 6C and S4F). In contrast, the C1263A and H1283A mutants failed to rescue as efficiently (Figures 6C, 6D, S4C, and S4D; data not shown). These data establish functional conservation of human ZNF644 with both zebrafish paralogs and further demonstrate the critical importance of physical association with G9a to ZNF644 function.

Retinal Defects in znf644a and znf644b Morphants Are Recapitulated in the Midbrain

The cellular and molecular defects of g9a and znf644 morphant retinas were recapitulated in the midbrain, which, unlike other neural regions (Figures S3A and S3C), showed more conspicuous morphological defects. Specifically, znf644a and znf644b morphant midbrains exhibited upregulated and mislocalized expression of ccd1 at 48 hpf.
(Figure 7A). Strikingly, combined subphenotypic doses of g9a-MO, znf644a-MO, or znf644b-MO again gave rise to localized ccnd1 expression in midbrain cells at 48 hpf (Figure S6A), suggesting cooperative roles in gene suppression in the midbrain. As in the retina, g9a and znf644b morphants exhibited increased apoptosis in the midbrain, which was again rescued by co-injection of p53-MO (Figure 7B). Moreover, znf644a morphants exhibited prolonged PCNA expression and elevated numbers of pH3+ cells throughout the midbrain (Figure 7C and data not shown). Collectively, these data provide evidence that the cooperative functions of g9a, znf644a, and znf644b in the retina are recapitulated in other populations of neural progenitor cells.

**DISCUSSION**

**ZNF644 Is a Co-regulator of G9a/H3K9me2-Mediated Gene Silencing during Neurogenesis**

Our high-confidence interaction dataset recapitulated annotated complexes and many other recently reported protein-protein interactions (Figures 1A and 1B; Table S1). Among the physical interactions identified, we functionally characterized the association of the C2H2-like ZF protein ZNF644, linked to high-grade myopia, with the HMTs G9a and GLP.

As with another documented interactor, WIZ, ZNF644 contains an atypical C2H2-like ZF motif, which we determined is necessary and sufficient for physical association with the G9a and GLP via direct binding to the SET domain (Figures 1D and 1E). These findings contrast with a recent report that a poorly conserved N-terminal region of ZNF644 is responsible for associating with G9a/GLP (Bian et al., 2015) and imply that the conserved atypical C2H2-like ZF motifs of ZNF644 and WIZ constitute a common G9a/GLP recognition module. Point mutations in this atypical C2H2-like ZF motif that abolished interaction with G9a/GLP also failed to rescue znf644a and znf644b morphant phenotypes (Figures 6C and 6D). Furthermore, using shRNA-based knockdown assays in human cell culture, we determined that ZNF644 specifically co-regulates H3K9me2 levels but not other histone methylation marks, such as H3K9me3 (Figure 1F), suggesting a specialized function for ZNF644 related to G9a/H3K9me2-mediated gene silencing. Collectively, these data establish ZNF644 as a functionally relevant G9a/GLP-interacting protein and a specific co-regulator of H3K9me2 formation.

**Multifaceted Regulation of Differentiation by Functional and Physical Interactions**

Shortly after 24 hpf, most zebrafish RPCs exit the cell cycle to generate post-mitotic differentiated progeny. These changes in RPC identity occur with concomitant suppression of multipotency-related and cell-cycle-related genes (Dixit et al., 2014), although little is known regarding the physical interactions or tissue-specific co-factors governing this transition. The regulatory properties of ZNF644 vis-a-vis G9a/H3K9me2 and expression patterns of the zebrafish paralogs in neural progenitor cell populations at 24 hpf (but reduced at later stages of retinal development) imply that they are co-regulators of H3K9me2-mediated gene silencing. We showed that disruption of znf644a or znf644b function leads to downstream defects in retinal differentiation that particularly manifest at 48–72 hpf.

Despite distinct multipotency-related gene-expression programs (e.g., vsx2 expression in RPCs, but not in midbrain progenitors), similar histone methylation-related complexes may dictate gene silencing and cellular differentiation in distinct populations of neuronal progenitors. While loss of ZNF644 function in cell culture and in vivo leads to deficits in H3K9me2, additional investigations are needed to determine the precise molecular mechanism whereby ZNF644 regulates G9a/GLP activity. Although ZNF644 has multiple C2H2-like ZF motifs and evidence for a role in recruiting G9a/GLP to chromatin has been proposed (Bian et al., 2015), we have failed to establish whether ZNF644 is truly capable of sequence-specific nucleic acid binding (our unpublished data). Neither human nor zebrafish ZNF644 contain the modular three-ZF array typically found in the DNA-binding variety of C2H2 ZF-containing proteins (Lam et al., 2011). One possibility, however, is that ZNF644 may indirectly impart target gene specificity to G9a/GLP by serving as a molecular scaffold bridging HMTs to DNA/RNA-binding repressor molecules and/or other co-repressors to facilitate the stepwise, stable suppression of multipotency- and cell-cycle-related loci. Such a possibility is supported by findings in ESCs that WIZ serves as a physical scaffold bridging G9a/GLP to the CtBP co-repressor complex (Ueda et al., 2006). An alternative possibility, not necessarily mutually exclusive, is that ZNF644 directly stimulates the catalytic activity of G9a/GLP via direct physical interaction to the SET domain. Regardless, given the complex molecular mechanisms involved in G9a/GLP targeting in other contexts (Cedar and Bergman, 2009), additional investigations are needed to define the precise functions of ZNF644 and WIZ in G9a/GLP-mediated silencing during neurogenesis.

Roles for G9a/GLP in regulating cell differentiation in other contexts, such as in HSPCs (Chen et al., 2012) and ESCs (Tachibana et al., 2008), have been reported. In neural progenitors, disruption of G9a/ZNF644 results in the extended maintenance of a proliferative progenitor identity, resulting in delayed formation of differentiated
retinal neurons. We observed H3K9me2+ nuclei within populations of differentiated retinal neurons, which may represent expanded regions of H3K9me2-marked chromatin reported elsewhere (Wen et al., 2009). The emergence of H3K9me2+ nuclei was preceded temporally by suppression of retinal progenitor identity and by expression of lineage commitment, suggesting that formation of expanded H3K9me2-marked chromatin may be a late-phase process occurring within differentiated neuronal cells. Given the reduced retinal expression at 48 or 72 hpf (Figure S1D), when H3K9me2+ nuclei are detectable, znf644a and znf644b are unlikely to directly regulate

Figure 7. The Retinal Defects of g9a, znf644a, and znf644b Morphants Are Recapitulated in the Midbrain
(A) WISH assays monitoring ccnd1 expression in WT, znf644a, or znf644b morphant midbrain cross-sections at 48 hpf. Arrows indicate mislocalized ccnd1 expression.
(B) Immunostaining of cleaved Caspase3 in g9a or znf644b morphant midbrain cross-sections at 72 hpf and rescue by p53-MO (n = 3 in each group). Arrows denote the position of Caspase3+ cells.
(C) Immunostaining of PCNA in WT or znf644a morphant midbrain cross-sections (n = 3 in each group). Arrows denote the position of PCNA+ cells.
formation of expanded regions of H3K9me2-marked chromatin in differentiated cell populations, but rather may regulate initial “seeding” of H3K9me2 marks at isolated genomic regions in retinal cells, at around 24 hpf, such as the promoter regions of vsx2 and ccdn1, as we demonstrated.

**Possible Insights into ZNF644 Function and High-Grade Myopia**

It is generally thought that longitudinal stretching (or lack thereof) can alter axial elongation, resulting in myopia or hyperopia (Wallman and Winawer, 2004). However, some reports implicate developmental-related genes and signaling pathways in post-natal retinal growth (Hysi et al., 2010; Sundin et al., 2005). In primates, disruption of the CMZ, which harbors a proliferating RPC niche, can cause high-grade myopia (Tkatchenko et al., 2006). It is not obvious how SNPs in the coding sequence and 3’ UTR might affect ZNF644 function. Further investigations are needed to determine the precise role of ZNF644 (and impact of mutations associated with high-grade myopia) in controlling G9a/H3K9me2 during retinal growth and differentiation in adult stem cell populations.

**EXPERIMENTAL PROCEDURES**

**Affinity Purification and Mass Spectrometry**

Sequence-verified open reading frames from the Human ORFeome library (Open Biosystems) and UltimateORF collection (Invitrogen) from cDNA were cloned into pDONR221 or pDONR223. Replicate two-step enrichment AP-MS experiments were performed from HEK293 cell lysates essentially as previously described (Mak et al., 2010). GFP-tagged AP-MS assays used Flp-In T-Rex HEK293 cells (a kind gift from Anne-Claude Gingras) cultured in media containing 250 \( \mu \)g/ml doxycycline for 24 hr, followed by affinity purification using 1 \( \mu \)g of rabbit anti-GFP ABfinity recombinant antibody (Invitrogen, G10362). See Supplemental Experimental Procedures for additional information.

**Zebrafish Husbandry**

Danio rerio (AB strain) were raised and bred at 28°C in accordance with regulations on animal experimentation established by the Canadian Council on Animal Care. Care and procedures were approved by the University of Toronto Animal Care Committee. The Tg(Isl2b:mGFP) strain was a kind gift from Dr. Chi-Bin Chien (University of Utah).

**Morpholino, mRNA Injections, and Statistics**

Embryos were injected at the 1- to 2-cell stage with antisense morpholino oligonucleotides (Gene Tools) and/or in vitro transcribed, capped full-length mRNA. Morpholino sequences, targeted splice junctions, and injection doses are listed in Supplemental Experimental Procedures. RT-PCR assays were performed from cDNA derived from RNA derived from de-yolked embryos (tails removed) using TRIZol (Invitrogen). In all cases, “n” refers to the number of individual embryos assayed in the experiments.

**Whole-Mount In Situ Hybridization**

Whole-mount in situ hybridization (WISH) was essentially carried out as described in Supplemental Experimental Procedures. RNA antisense probes were obtained from the following sources: znf644a and znf644b, cloned from 24-hpf embryos; lef1 and otx2, kind gifts from Ashley Bruce; ath5, a kind gift from Jeremy Kay; rx3, a kind gift from Pamela Raymond, vsx2, Open Biosystems; and ccdn1 (cb161), ZIRC.

**Chromatin Immunoprecipitation**

ChIP assays were performed using cells derived from ~100 zebrafish heads at 48 hpf essentially as previously described (Lindeman et al., 2009). In brief, cells were crosslinked using 1% formaldehyde and quenched with 0.125 M glycine. Sonication was performed by Bioruptor (Diagenode) using seven cycles. ChIP was performed using the EZ-ChIP kit (Millipore), H3K9me2 (AbCam, ab1220) and isotype control (AbCam, ab170191) antibodies were used at 2 \( \mu \)g per sample. See Supplemental Experimental Procedures for further information.

**Immunostaining and BrdU Incorporation Assays**

Samples were fixed in 4% paraformaldehyde and processed as described (Wong et al., 2010). Sections were stained with Hoechst 33,342 and mounted with 80% glycerol. Antibodies used are listed in Supplemental Experimental Procedures. BrdU (5 mM) was injected into the tectal brain ventricle of 48-hpf embryos and fixed with 4% paraformaldehyde, followed by cryosection and immunostaining. Slides were treated with 20 U/mL BrdU, detected using an anti-BrdU antibody (Cedarlane, MCA2060), and imaged by confocal microscopy imaging.

**ACCESSION NUMBERS**

The accession number for znf644a reported in this paper is RefSeq: NM_001278254.1. The accession number for znf644b reported in this paper is RefSeq: XM_003198781.4.

**SUPPLEMENTAL INFORMATION**

Supplemental Information includes Supplemental Experimental Procedures, six figures, and four tables and can be found with this article online at http://dx.doi.org/10.1016/j.stemcr.2016.06.012.

**AUTHOR CONTRIBUTIONS**

J.O. and L.W. designed and conducted most of the experiments, analyzed the data, and wrote the manuscript. S.D., A.M., and H.G. conducted experiments and analyzed data. Y.L. and Z.Z. analyzed data. J.F.G. provided reagents, technical support, and scientific advice. V.T and A.E. designed the study, analyzed the data, wrote the manuscript, and funded the research.
ACKNOWLEDGMENTS

We thank Guoqing Zhong, Xinghua Guo, and Oxana Pogouste for assistance with affinity purifications as well as Sadhna Phanse for assistance with mass spectrometry data analysis. This work was supported in part by grants from the Ontario Ministry of Innovation (ORF-GL2) and the Canadian Institutes of Health Research (TEAM grant 8294) to A.E.; a grant from the Canadian Institutes of Health Research (MOP-119415) to V.T; an Ontario Graduate Scholarship to J.O; and Vision Science Research Program Scholarships to L.W. and A.M.

Received: November 16, 2015
Revised: June 28, 2016
Accepted: June 29, 2016
Published: August 18, 2016

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