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FoxO6 affects Plxna4-mediated neuronal migration during mouse cortical development

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The forkhead transcription factor FoxO6 is prominently expressed during development of the murine neocortex. However, its function in cortical development is as yet unknown. We now demonstrate that cortical development is altered in FoxO6^{+/-} and FoxO6^{-/-} mice, showing migrating neurons halted in the intermediate zone. Using a FoxO6-directed siRNA approach, we substantiate the requirement of FoxO6 for a correct radial migration in the developing neocortex. Subsequent genome-wide transcriptome analysis reveals altered expression of genes involved in cell adhesion, axon guidance, and gliogenesis upon silencing of FoxO6. We then show that FoxO6 binds to DAF-16-binding elements in the *Plexin A4 (Plxna4)* promoter region and affects *Plxna4* expression. Finally, ectopic *Plxna4* expression restores radial migration in FoxO6^{+/-} and siRNA-mediated knockdown models. In conclusion, the presented data provide insights into the molecular mechanisms whereby transcriptional programs drive cortical development.

FoxO6 | cortex | development | radial migration | Plxna4

The FoxO (forkhead box O) family of transcription factors is known to be involved in a wide array of biological processes, including apoptosis, cellular survival, differentiation, and stress resistance. FoxOs are part of the conserved PI3K-AKT-FoxO signaling pathway, which has been shown to convey extracellular signals to intracellular transcriptional programs. Insulin and insulin-like growth factors (IGFs; e.g., Igf1) regulate FoxO transcriptional activity mainly via posttranslational modifications, resulting in the nuclear exclusion of the protein, thereby rendering it inactive (1–5). In humans, de novo mutations in genes involved in the PI3K/AKT pathway result in a wide array of cortical malformations, including cortical dysplasia, megalencephaly-polymicrogyria-polydactyly-hydrocephalus (MPPH), and megalencephaly-capillary malformation-polymicrogyria syndrome (MCAP) (6, 7). Studying the downstream effectors of this pathway (e.g., FoxO proteins) provides in-depth insights into the molecular mechanisms underlying these cortical malformations.

Recent studies have shown that FoxOs are required for the coordinated regulation of postnatal neuronal stem cell (NSC) homeostasis, demonstrating a prominent role for FoxO3 in NSC proliferation and renewal (8–10). In a combined *FoxO1,3,4*-deficient mouse model, an initial increase in brain size was observed followed by a decline in the adult NSC pool. Furthermore, it was shown, both in vivo and in vitro, that the single-*FoxO3*-deficient animal results in a similar decline in NSC number (10). FoxO6 was shown to be required for the regulation, in the hippocampus, of a set of genes involved in synaptic function (11) and is essential in Pak1-mediated cellular polarity in the cerebellum (12), demonstrating the importance of this forkhead member in the mammalian brain.

Apart from the functions of FoxO6 in the postnatal hippocampus and cerebellum, FoxO6 is likely to play a role during the embryonic development of the cortex, judging from its specific temporal and spatial expression pattern (13). From embryonic day (E)12.5 during murine embryonic development, *FoxO6* expression is found mainly in the hippocampus, cortex, and striatum (14). This

pattern is conserved at later stages until birth, when expression of *FoxO6* is mainly found in the hippocampus (14). Importantly, in early developmental stages, expression is observed in proliferating areas in the cortex whereas at later stages *FoxO6* is also prominently expressed in the postmitotic cortical plate, suggesting different functions for FoxO6 during development.

During cortical development, neuroepithelial cells, located in the cortical ventricular zone of the neural tube, will elongate at mouse E11 and assume a radial glial morphology. These radial glial cells divide asymmetrically to generate new radial glial cells and intermediate progenitor cells, destined to become postmitotic neurons (15, 16). Postmitotic cells undergo outward migration to form the upper layers of the cortical plate and eventually generate the six-layered adult neocortex. Although several factors, such as guidance molecules and cell-adhesion molecules (10, 17, 18), have been implicated in coordinating migration, molecular programming underlying these processes remains poorly understood.

In this study, we show that FoxO6 affects radial migration events during cortical development. Transcriptome analysis revealed that *Plexin A4 (Plxna4)* is regulated by FoxO6 in different mouse models. Moreover, we show that FoxO6 occupies DAF-16-binding sites present in the *Plxna4* promoter and is able to regulate its transcription. Finally, in *FoxO6* knockdown models, radial migration is restored by ectopic overexpression of *Plxna4*. These

Significance

The molecular basis of radial migration of cortical neurons is a well-studied process showing prominent roles for axon guidance, cell adhesion, cell polarity, and cytoskeleton remodeling. Remarkably, knowledge about transcriptional control of such processes is scarce. In this study, we show that the forkhead transcription factor FoxO6 influences *Plexin A4 (Plxna4)* expression, a key component of the Semaphorin signaling pathway, known for its role in axonal guidance and cortical migration. FoxO6 knockdown animals show a hampered migration of embryonic day 14.5-born neurons, which can be rescued by recombinant *Plxna4* expression constructs. Altogether, our data provide insights into the molecular mechanisms whereby transcriptional programs influence cortical development.

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The authors declare no conflict of interest.

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Data deposition: The microarray data reported in this paper have been deposited in the Gene Expression Omnibus (GEO) database, www.ncbi.nlm.nih.gov/geo (accession no. GSE71954).

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postmitotic regions of the developing cortex were affected by the loss of *FoxO6* at E18, we performed immunohistochemistry for B cell leukemia/lymphoma 11B (Ctip2) and special AT-rich sequence binding protein 2 (satb2) to identify the subcortical and callosal projecting pyramidal neurons (Fig. 1 *B* and *C*). The amount of subcortical projecting neurons, located in the deep layers V and VI (marked by Ctip2), was not significantly altered in *FoxO6*^{-/-} cortices at E18 (Fig. 1 *D* and *F*; *n* = 4 per condition). We next divided the Satb2-positive cortical region into two bins covering the upper-layer Satb2-positive neurons and the deep-layer, still migrating, Satb2-positive neurons, revealing both populations to be not significantly changed (Fig. 1 *E* and *G*; *n* = 4 per condition). Supporting the results described above, the gross morphology of the developing layers was unaffected in *FoxO6*^{-/-} cortices as shown by an unchanged thickness of the VZ/SVZ, intermediate zone (IZ), and CP at E18 (Fig. S1*E*; *n* = 3 per condition).

***FoxO6*^{-/-} Mice Display Hampered Radial Migration During Cortical Development, Resulting in a Decreased Amount of Cut-Like Homeobox 1-Positive Cells.** Because global cortical markers did not reveal significant changes in developing *FoxO6*^{-/-} cortices, we set out to

detect more subtle deviations in the developing cortex caused by *FoxO6* ablation. Therefore, we introduced a *GFP* expression plasmid in E14.5 *FoxO6*^{-/-} and *FoxO6*^{+/+} cortices by in utero electroporation, thereby tracing neurons born at the peak of neurogenesis. At E18, GFP-positive transfected cells in control cortices (Fig. 2*A*) migrated outward, reaching the cortical plate. Interestingly, in *FoxO6*^{-/-} cortices (Fig. 2*A*), the proportion of GFP-positive cells reaching the cortical plate was decreased significantly. Quantification revealed that a majority of the GFP-positive cells (54.40%; SD ±0.53) in wild-type control cortices reached the cortical plate, whereas only 37.2% (SD ±6.1; *P* < 0.05) of these cells in *FoxO6*^{-/-} cortices reached the cortical plate (Fig. 2*B*; *n* = 4 per condition). In line with these findings, 42.83% (SD ±3.10) of GFP-positive cells in *FoxO6*^{-/-} cortices are located in the IZ compared with 34.50% (SD ±0.35; *P* < 0.05) in wild-type littermate controls (Fig. 2*B*). Furthermore, we observed a significantly increased ratio of GFP-labeled cells in the VZ of *FoxO6*^{-/-} cortices (19.4%; SD ±3.8) compared with *FoxO6*^{+/+} controls (11%; SD ±0.35; *P* = 0.01) (Fig. 2*B*), indicating the necessity of proper FoxO6 function for the migration of these

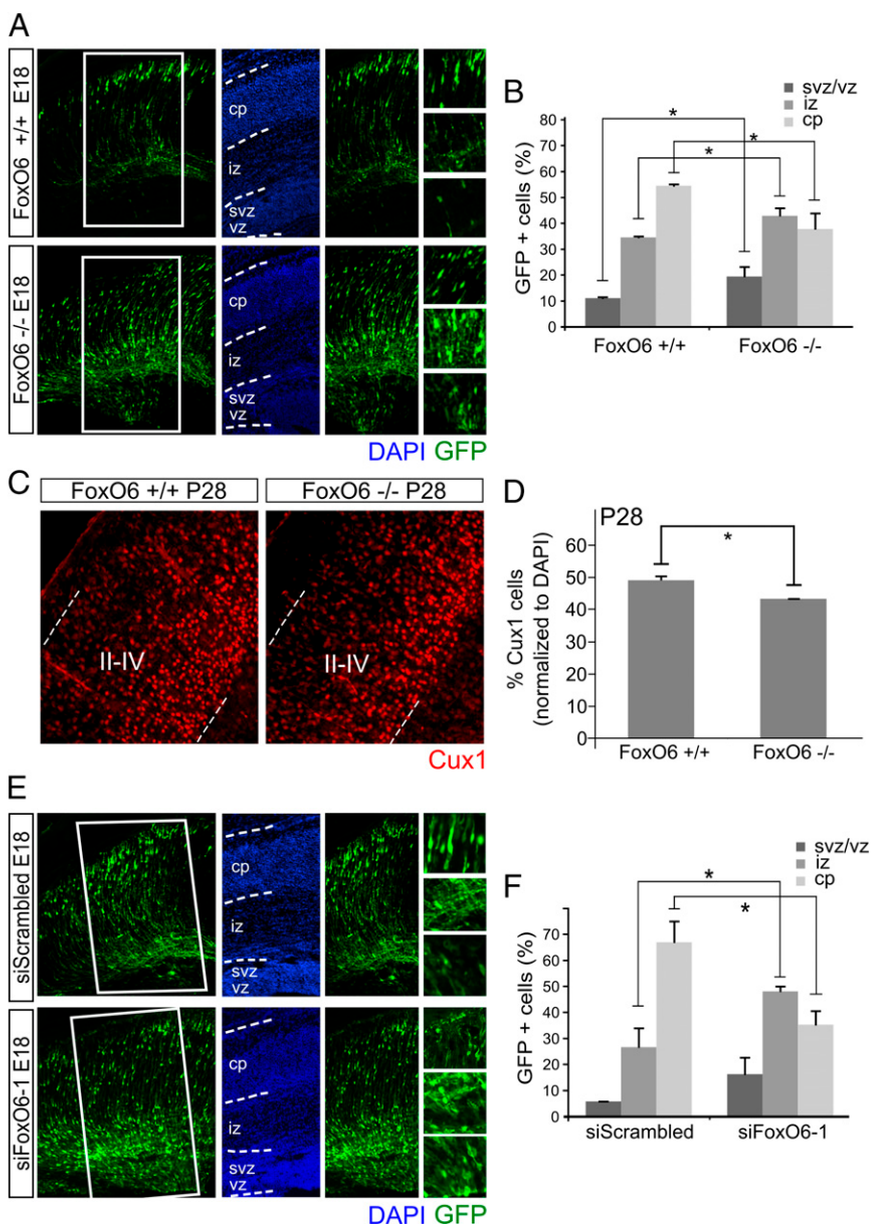


Fig. 2. *FoxO6*^{-/-} mice display hampered radial migration during cortical development, which is exemplified by the altered presence of Cux1-positive callosal projection neurons. (A) Tracing newborn neurons by in utero electroporation (IUE) at E14.5 with a *GFP* expression vector revealed an altered distribution of cortical cells in *FoxO6*^{-/-} cortices compared with *FoxO6*^{+/+} cortices at E18. Magnifications are shown for the cortical plate, intermediate zone, and ventricular/subventricular zone. (B) Quantification of triplicate experiments as shown in A. Ratios between the ventricular zone, intermediate zone, and cortical plate are shown. (C) Immunostaining for Cux1 in *FoxO6*^{-/-} cortices compared with *FoxO6*^{+/+} cortices. (D) Quantification of the number of Cux1-positive cells for a fixed region covering layers II–IV in *FoxO6*^{+/+} and *FoxO6*^{-/-} cortices. Data were normalized against the number of DAPI cells in that region. (E) Cortices in utero electroporated with siFoxO6-1 at E14.5 and cotransfected with a *GFP* expression vector display altered neuronal distribution at E18 compared with siScrambled (control) electroporated cortices. Magnifications are shown for the cortical plate, intermediate zone, and ventricular/subventricular zone. (F) Quantification of triplicate experiments as shown in E. Ratios between the ventricular zone, intermediate zone, and cortical plate are shown. The error bars show the SD. Two-tailed Student's *t* test; **P* < 0.05 (Bonferroni-corrected significance level in B and F).

developing neurons. Although FoxOs have been implicated to regulate cell polarity in the cerebellum (12), magnifications of the different cortical bins revealed that GFP-labeled cells, ablated for *FoxO6*, display a normal bipolar morphology (Fig. 2A, Right), which makes polarity changes unlikely to be related to the observed phenotype. These data, together with the observation that global cortical markers are unaffected in *FoxO6*^{-/-} cortices, suggest that only a relatively small proportion of cells born at E14.5 is affected by the loss of *FoxO6*. In line with this, we analyzed the amount of cut-like homeobox 1 (*Cux1*)-positive cells in postnatal day (P)28 *FoxO6*^{-/-} and *FoxO6*^{+/+} cortices (Fig. 2C and D). As shown, the amount of *Cux1*-positive cells in the upper layers of the *FoxO6*^{-/-} cortex is decreased significantly (12%; $P < 0.05$; $n = 3$ per condition), indicating that ablation of *FoxO6* influences a proportion of cortical neurons born at E14.5.

siRNA-Mediated Down-Regulation of *FoxO6* Expression in the Embryonic Cortex Phenocopies the Radial Migration Defect Observed in *FoxO6*^{-/-} Mice. To substantiate the cell-autonomous role of *FoxO6* in cortical migration, we followed an acute knockdown strategy in which *FoxO6* expression in the embryonic cortex was down-regulated via in utero electroporation of siRNAs specifically directed against *FoxO6*. Out of two different siRNAs directed against *FoxO6*, we used the most efficient one in these experiments (siFoxO6-1; Fig. S2). To monitor radial migration, we coelectroporated a GFP expression vector and analyzed these cortices at E18. Mimicking *FoxO6*^{-/-} animals, a hampered radial migration was observed at E18 when cortices were electroporated with siFoxO6-1 at E14.5 (Fig. 2E and F). At E18, 3.5 d post-electroporation, cortices electroporated with scrambled control siRNA showed 27% (SD ± 2.5) of transfected cells located in the intermediate zone compared with 48% (SD ± 1.1 ; $P < 0.001$) in cortices down-regulated for *FoxO6*. Furthermore, only 35.2% (SD ± 1.4) of the transfected cells in cortices electroporated with siFoxO6-1 reached the cortical plate, compared with 66.9% (SD ± 1.2 ; $P < 0.001$) in controls (Fig. 2E and F). Furthermore, we observed an increased ratio of GFP-labeled cells in the ventricular and subventricular zones in cortices down-regulated for *FoxO6* (16.2% SD ± 6.6) compared with control cortices (5.7%; SD ± 2.2 ; $P < 0.05$). Taken together, the phenotype as observed in *FoxO6*^{-/-} electroporated cortices resembles the *FoxO6* siRNA-mediated knockdown phenotype. To elaborate upon the specificity of the siRNA, we introduced the siRNA targeting *FoxO6* in *FoxO6*^{-/-} cortices. As expected, in these cortices, the distribution of the GFP-labeled cells was unaltered (Fig. S3). In addition, to exclude the possibility that part of the observed siRNA-mediated phenotype is due to off-target effects, we performed similar in utero electroporation experiments but now using a different siRNA directed against *FoxO6* (siFoxO6-2) targeting a different part of the *FoxO6* mRNA. Importantly, cortices electroporated with siFoxO6-2 showed an identical phenotype as cortices electroporated with siFoxO6-1 (Fig. S4), indicating that the phenotype observed is due to the down-regulation of *FoxO6*. Taken together, siRNA-mediated knockdown of *FoxO6* in the developing cortex starting at E14.5 results in hampered radial migration similar to the effects observed in the full *FoxO6* knockout.

***FoxO6* Knockdown Results in Altered Expression of Genes Involved in Cell Adhesion, Axon Guidance, and Gliogenesis.** To gain insight into the acute molecular mechanisms governing *FoxO6*-mediated cortical radial migration, genome-wide transcriptome analysis was performed. To this end, we specifically silenced *FoxO6* expression in the cortex at E14.5 by in utero electroporation of siFoxO6-1 together with a GFP expression plasmid. GFP-positive cells were FACsorted at E16.5, which is at the peak of radial migration and neurogenesis, followed by RNA isolation (Fig. 3A). We used two different siRNAs targeting *FoxO6* (siFoxO6-1 and siFoxO6-2) and independently compared the results with a control siRNA (siScrambled, common reference) to minimize off-target effects and to select the most significant genes regulated by *FoxO6* activity (Table S1). For each condition at least

three cortices were pooled (randomized over litters) to minimize the possibility of finding initial differences due to the electroporation procedure itself or due to between-nest variation. In total, 12 cortices were used, whereby for each biological replicate 3 cortices were pooled, generating a total 4 replicates. Electroporated cortices from different litters were pooled to minimize effects due to electroporation and litter differences. To investigate whether the different siRNAs and the separate in utero electroporations provided similar data, we performed a correlation analysis on the false discovery rate (FDR)-corrected values (FDR cutoff 0.1). The correlation coefficient (R) was at least 0.85 (siFoxO6-2, $R = 0.85$, $P < 0.001$; siFoxO6-1, $R = 0.89$, $P < 0.001$), indicating that the gene expression profile as a result of the *FoxO6* knockdown by the two different siRNAs is highly comparable (Fig. 3B). Subsequently, we performed a Gene Ontology analysis (19) on the separate datasets, showing that similar processes are highly enriched (Fig. 3C). The results from siFoxO6-2 ($n = 3$) and siFoxO6-1 ($n = 3$) and control siRNA (common reference) ($n = 9$) were used in a three-way comparison, thereby generating two independent datasets that contained ± 200 and ± 300 *FoxO6*-regulated genes ($P < 0.05$ cutoff) (FDR) (Fig. 3D). Overlap of these two restricted datasets ($P < 0.05$, for separate list selection) yielded a list of 24 regulated genes (Fig. 3D). We confirmed the transcription modulation of several genes in independently electroporated and FACsorted cortices using quantitative (q)PCR [Fig. S5; LIM homeobox transcription factor 1 alpha (*Lmx1a*) was used as an unchanged reference], thereby confirming the validity of the transcriptome analysis.

***FoxO6* Affects Radial Cortical Migration in a *Plxna4*-Dependent Manner.** One of the identified *FoxO6*-regulated genes, *Plxna4* (Fig. 3D), is known to be required for proper radial migration of newborn neurons in the developing cortex. *Plxna4* plays a part in the Semaphorin signaling cascade, which is disrupted upon *Plxna4* ablation, resulting in an impaired radial migration of cortical neurons (20). Importantly, the observed defect found in *FoxO6*-deficient cortices, with neurons halted in the deeper layers of the developing cortex, phenocopies the defect observed when rat embryonic cortices are electroporated with siRNA specifically directed against *Plxna4* (20). To further explore the possibility that the correct radial migration of neurons in the developing cortex is influenced by a *FoxO6*-mediated regulation of *Plxna4*, we first analyzed the spatial and temporal expression profile of *Plxna4* by performing in situ hybridization of E16.5 and E18 cortices (Fig. 4A). At E16.5, from rostral to caudal, *Plxna4* is most prominently expressed in the superficial layers of the cortical plate and intermediate zone with an apparent gradient from lateral to medial regions. At E18, expression is not restricted to the lateral regions of the developing cortex but is now prominently expressed in the entire outer layer of the cortical plate (Fig. 4A), resembling, in part, the expression domain of *FoxO6* (Fig. 4B), as shown in Fig. 1. To substantiate the possible functional interaction of *Plxna4*, *FoxO6*, and cortical migration, we designed an in utero electroporation experiment in which we overexpress *Plxna4* in *FoxO6*-deficient animals.

As demonstrated earlier, in *FoxO6*^{-/-} cortices, radial migration is altered (Fig. 2A and B), showing GFP-labeled cells primarily in the IZ of the developing cortex. To investigate whether *Plxna4* is related to the disrupted radial migration in *FoxO6*^{-/-} animals, we expressed *Plxna4* in these cortices at E14.5 and analyzed the distribution of GFP-labeled cells at E18. Our results show that in *FoxO6*^{-/-} cortices, ectopic expression of *Plxna4* was insufficient to restore normal radial migration (Fig. 4C and D). To investigate whether expression of *Plxna4* is altered in full *FoxO6* mutant cortices similar to neurons silenced for *FoxO6* (Fig. 3D), we introduced a GFP expression plasmid at E14.5 in cortices and FACsorted the resulting GFP-labeled cells from *FoxO6*^{-/-}, *FoxO6*^{+/-}, and *FoxO6*^{+/+} cortices at E16.5, followed by qPCR for *Plxna4*. We did not observe a significant difference in the expression of *Plxna4* in *FoxO6*^{+/+} and *FoxO6*^{-/-} cortices. However, we did detect a significant reduction of *Plxna4* in *FoxO6*^{+/-} cortices (Fig. S6; $P < 0.05$). This indicates that full *FoxO6* knockout

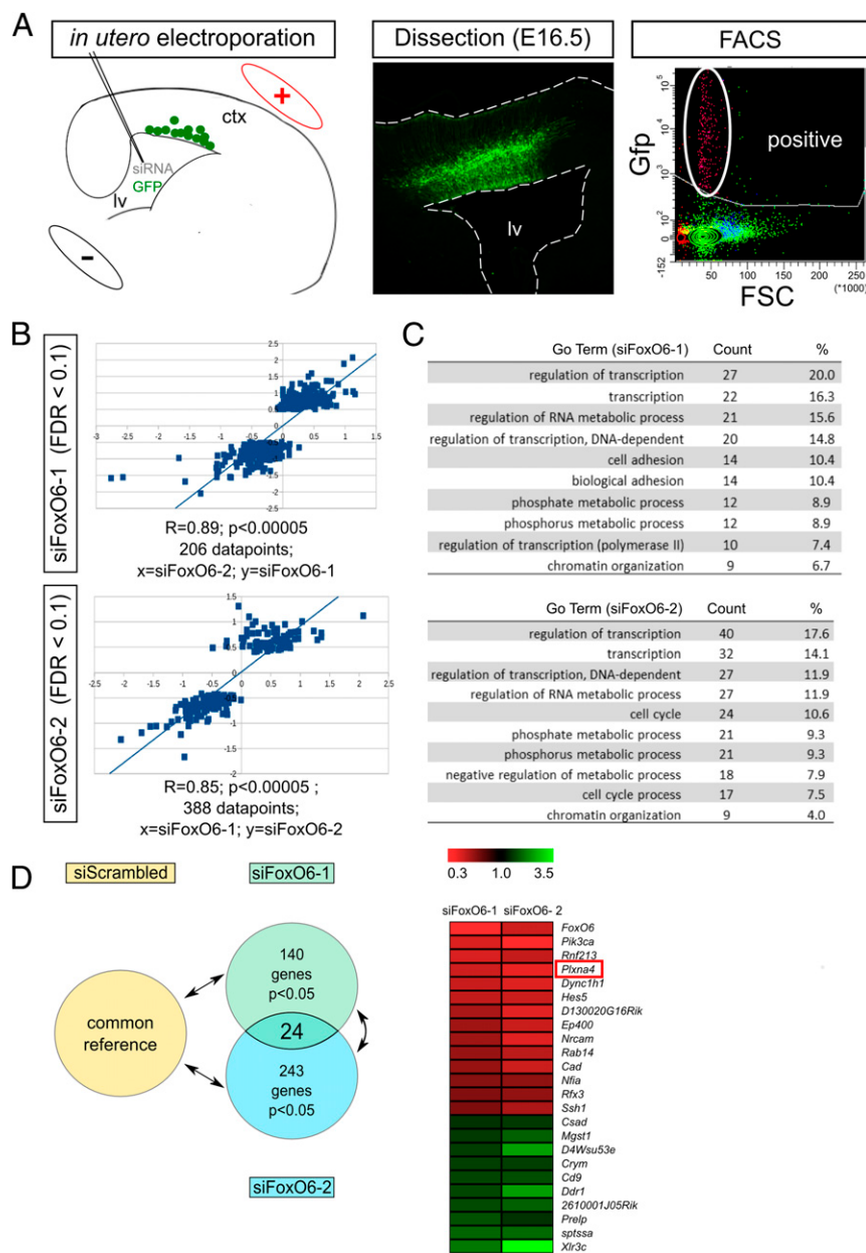


Fig. 3. Genome-wide transcriptional profiling of *FoxO6* knockdown neurons. (A) A graphical representation of the methodology used. *FoxO6* is down-regulated by in utero electroporation at E14.5 using two different siRNAs targeting *FoxO6* expression. Brains are dissected and dissociated at E16.5. Cortices are coelectroporated with GFP to FACS sort the transfected cells followed by RNA isolation and microarray analysis. Three electroporated cortices were pooled for each *N* per condition to minimize the effects due to electroporation differences and between-litter differences. (B) Regulated genes by *FoxO6* (FDR cutoff 0.1), identified as a result of the two different siRNAs, display a strong correlation indicating that in both datasets similar genes are represented, which are changed in a similar direction; siFoxO6-2, $R = 0.85$, $P < 0.01$; siFoxO6-1, $R = 0.89$, $P < 0.01$. (C) GO term analysis reveals that similar processes are regulated by *FoxO6* knockdown through the use of siFoxO6-2 and siFoxO6-1. (D, Left) Graphical representation of the microarray analysis setup and resulting amount of regulated genes (243, siFoxO6-2; 140, siFoxO6-1) and the exact overlap (24 genes) in the $P < 0.05$ FDR-restricted datasets. (D, Right) Heatmap of the 24 genes that are present in such an overlap, including *FoxO6*. ctx, cortex; FACS, fluorescence-activated cell sorting; FSC, forward scatter; GFP, green fluorescent protein; lv, lateral ventricle.

animals, which lack *FoxO6* from the beginning, have compensated for this loss of functionality, suggesting that the migration defect is not caused by limiting levels of *Plxna4* in *FoxO6*^{-/-} animals. However, in developing cortices that do contain *FoxO6*, but in limiting levels, a reduction of *Plxna4* is observed, which would suggest that in such animals *Plxna4* overexpression might influence the radial migration events. To investigate this further, we performed a rescue experiment in *FoxO6*^{+/-} cortices. To this end, we coelectroporated E14.5 cortices with expression plasmids for *GFP* and *Plxna4* or an empty vector (EV) control (Fig. 4 E and F). Importantly, overexpression of *Plxna4* in *FoxO6*^{+/-} cortices significantly increased the amount of GFP-positive cells in the CP ($P < 0.05$).

To further substantiate the role of *FoxO6* in modulating *Plxna4* expression, we performed similar rescue experiments in cortices silenced for *FoxO6* expression via siRNA-mediated knockdown. To this end, we coelectroporated E14.5 cortices with siFoxO6-1 and an expression vector for *Plxna4* or an EV control. We hypothesized that *Plxna4* is able to, at least partly, rescue radial migration in cells deficient for *FoxO6*, resulting in restored radial migration of

transfected cells (Fig. 5A). The overexpression of *Plxna4* was confirmed by immunohistochemistry (Fig. S7). Confirming earlier results, cortices electroporated with siFoxO6-1 and EV control ($n = 4$) showed 18.8% (SD ± 1.2) of transfected cells located in the ventricular zone compared with 14.0% (SD ± 0.36 ; $P < 0.001$) in cortices coelectroporated with siFoxO6-1 and the *Plxna4* expression vector (Fig. 5 B and C; $n = 4$). Furthermore, only 26.6% (SD ± 2.4) of the transfected cells in cortices electroporated with *Plxna4* were located in the IZ compared with 43.5% (SD ± 1.7 ; $P < 0.001$) in cortices coelectroporated with siFoxO6-1 and the EV control plasmid. Finally, 59.4% (SD ± 4.4) of the GFP-labeled cells reached the cortical plate when coelectroporated with *Plxna4* compared with 37.7% (SD ± 0.8 ; $P < 0.01$) in EV controls (Fig. 5 B and C). In conclusion, our data suggest that cortices electroporated with siFoxO6-1 and a *Plxna4* expression vector display restored migration of developing neurons toward the cortical plate compared with *FoxO6* knockdown controls. To test the possibility that overexpression of *Plxna4* results in improved migration of wild-type neurons by itself, wild-type E14.5 cortices were transfected with *Plxna4* or EV control plasmids. Analysis of

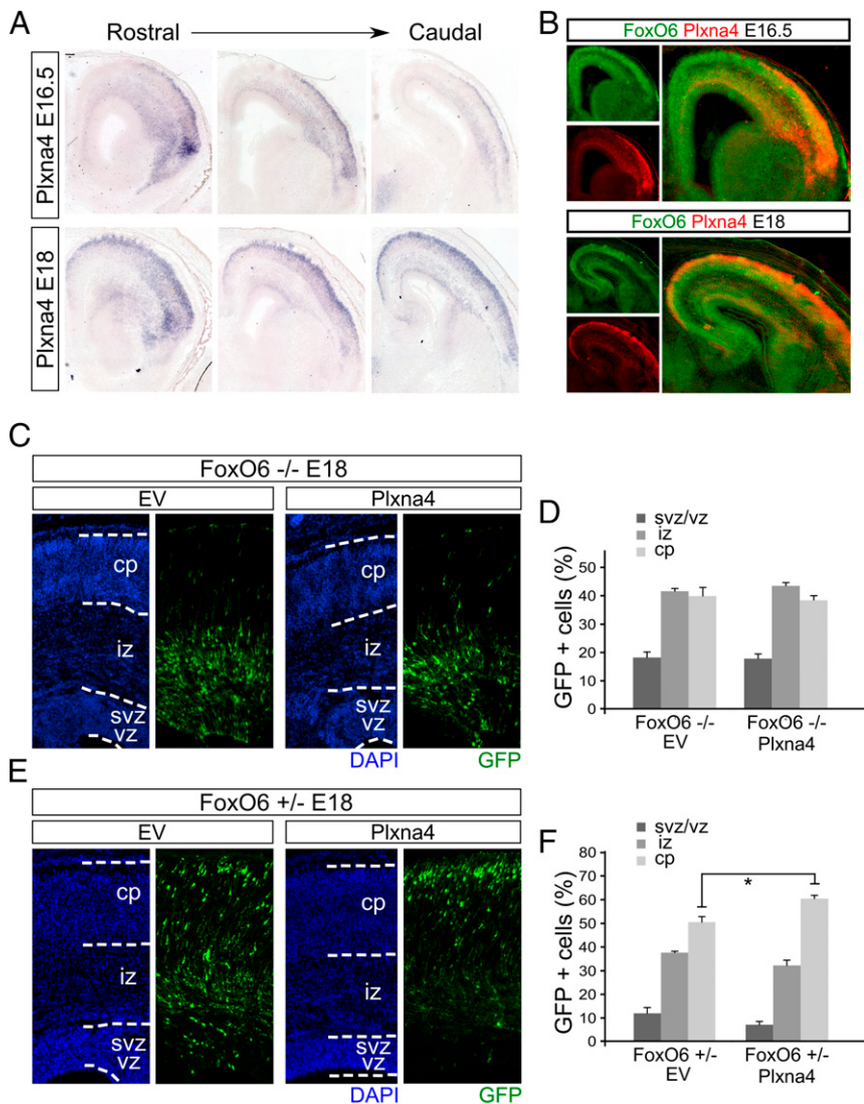


Fig. 4. *Plxna4* is expressed in the developing cortex and restores radial migration defects in *FoxO6*^{+/-} animals. (A) In situ hybridization for *Plxna4* at E16.5 and E18. At E16.5, from rostral to caudal, *Plxna4* is most prominently expressed in the superficial layers of the cortical plate and intermediate zone with an apparent gradient from lateral to medial regions. At E18, expression is not restricted to the lateral regions of the developing cortex but is now prominently expressed in the entire outer layer of the cortical plate (20). (B) Pseudooverlay of adjacent sections analyzed for *FoxO6* and *Plxna4* expression indicate regional overlap. (C) GFP-traced neurons in *FoxO6*^{-/-} cortices of E18 embryos in utero electroporated with a GFP expression vector and either empty vector control or *Plxna4* expression vector. Cortices were counterstained with DAPI. (D) The distribution of the transfected cells was quantified for the indicated layers for at least three brains per condition using multiple sections per brain. (E) GFP-traced neurons in *FoxO6*^{+/-} cortices of E18 embryos in utero electroporated with a GFP expression vector and either empty vector control or *Plxna4* expression vector. Cortices were counterstained with DAPI. (F) The distribution of the transfected cells was quantified for the indicated layers for at least three brains per condition using multiple sections per brain. The error bars show the SD. Two-tailed Student's *t* test; **P* < 0.05.

these cortices at E16.5 and E18 did not show an enhanced migration phenotype (Fig. S8), indicating that the induction of *Plxna4* expression above normal levels does not alter radial migration in wild-type neurons. To compare all used genetic models, we chose to calculate the difference in percentage of the GFP cells in the CP and IZ because these two parameters are mainly changed upon *FoxO6* modulation (Fig. 5 D and E). This representation clearly shows that the siRNA-mediated knockdown of *FoxO6* results in a negative value, indicating that more cells reside in the IZ compared with the CP. A similar pattern is observed in *FoxO6*^{-/-} animals. However, in *FoxO6*^{+/-} cortices, the amount of cells is higher in the CP compared with the IZ. This indicates that a mild phenotype is present in *FoxO6*^{+/-} animals. Introduction of *Plxna4* increases the migration to the CP in the knockdown and *FoxO6*^{+/-} cortices in concordance with the lower *Plxna4* levels in both the knockdown and *FoxO6*^{+/-} cortices. Taken together, our data indicate that radial migration events, in the developing murine cortex, are influenced by *FoxO6*.

FoxO6 Interacts with Specific Daf-16–Binding Elements Within the *Plxna4* Promoter, Regulating *Plxna4* Gene Expression. The results as described above indicate that the cortical migration defect, as observed in *FoxO6* knockdown models, is mediated by a decrease of *Plxna4* expression. To investigate whether *Plxna4* transcription is directly regulated by *FoxO6* via interaction with four in silico identified Daf-16–binding elements (DBEs) (21) located within

10 kb up- and downstream of the mouse *Plxna4* promoter (Fig. 6A), we performed chromatin immunoprecipitation (ChIP) analysis.

To this end, we performed two independent in vivo *FoxO6* ChIP-qPCR experiments on E16.5 cortices. First, we compared the *FoxO6* IP and the control IgG IP in *FoxO6*^{+/-} cortices. Using this approach, we measured the highest enrichment of *FoxO6* at DBE4 (Fig. 6B; *n* = 2). We then performed a *FoxO6* IP on E16.5 *FoxO6*^{+/-} and *FoxO6*^{-/-} cortices. Within this experimental design, *FoxO6*^{-/-} cortices served as the negative control. Here we observed an enrichment of *FoxO6* on each DBE, showing the highest enrichment on the locus covering DBE4 (Fig. 6C; *n* = 3). We then asked whether binding of *FoxO6* to DBEs located in the *Plxna4* promoter is able to regulate *Plxna4* transcription. To this end, we used part of the *Plxna4* promoter with and without confirmed *FoxO6*-binding elements and cloned this 5' of a luciferase reporter (pGL3). In this approach, luminescence (measured in relative light units; RLUs) is a readout for the activity of the cloned promoter regions. In these experiments, a positive control (pGL3-6xDBE) (5) showed a large increase in RLUs (*FoxO6*-GFP/GFP), confirming our approach (Fig. 6D). The pGL3-EV showed a 6.5-fold increase in luminescence (*FoxO6*-GFP/GFP) (Fig. 6D), suggesting that *FoxO6* may regulate transcription in general, as has previously been suggested (22). Importantly, promoter elements containing the confirmed in vivo *FoxO6*-binding element (pGL3-DBE4) show a significant increase in RLUs (*P* < 0.01) (*FoxO6*-GFP/GFP) compared with the control (pGL3-EV)

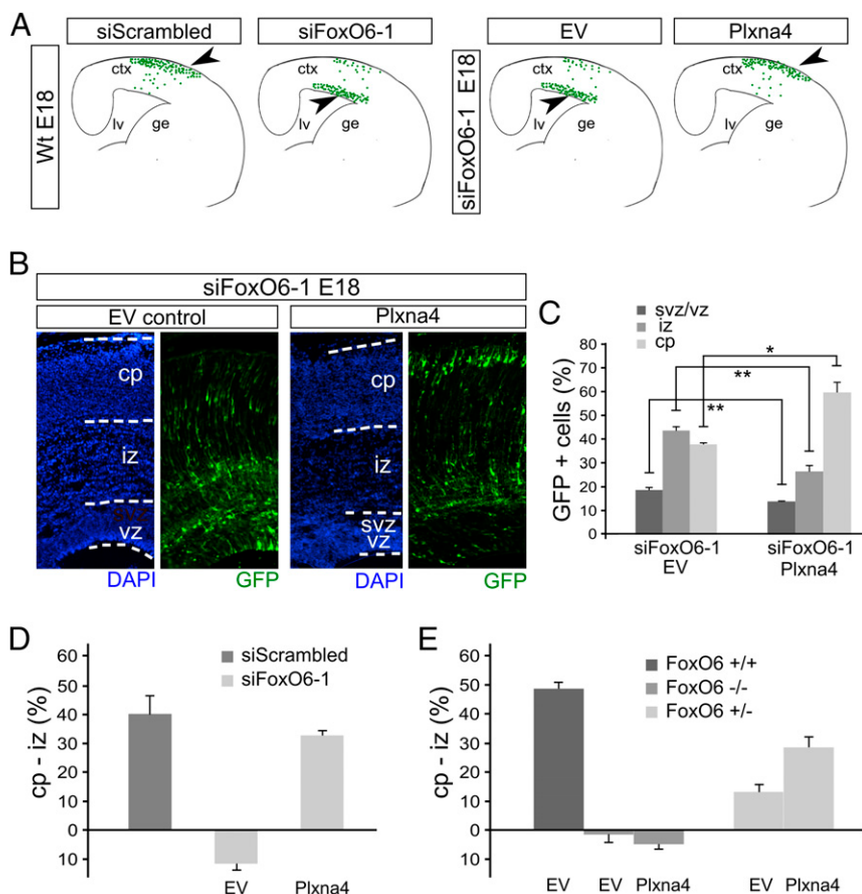


Fig. 5. *Plxna4* restores radial migration in *FoxO6* knockdown-induced radial migration defects. (A) Schematic overview of cortical migration phenotypes in *FoxO6*^{+/+} cortices transfected with the indicated siRNAs showing hampered radial migration in cortices transfected with siFoxO6-1. Schematic representation of the hypothesis that *Plxna4* rescues altered radial migration in cells down-regulated for *FoxO6* (Right). Arrowheads point to the hypothesized positions of the neurons in the different treatment situations. (B) GFP-traced neurons in cortices of E18 embryos in utero electroporated with siFoxO6-1, a GFP expression vector, and either empty vector control (20) or *Plxna4* expression vector. Cortices were counterstained with DAPI. (C) The distribution of the transfected cells was quantified for the indicated layers for at least three brains per condition using multiple sections per brain. The error bars show the SD. Two-tailed Student's *t* test; **P* < 0.05 and ***P* < 0.001. (D) Meta-analysis of the distribution of GFP-labeled cells in cortices treated with siScrambled, siFoxO6-1, and either empty vector control or *Plxna4* expression vector. The analysis represents the CP minus IZ value (%). (E) Similar representation as in D for *FoxO6*^{+/+}, *FoxO6*^{-/-}, and *FoxO6*^{+/-}, electroporated with either EV or *Plxna4*. Introducing *Plxna4* restores in part the percentage of GFP-labeled cells in the cortical plate in *FoxO6*^{+/-} cortices. The error bars show the SD. Wt, wild type.

(Fig. 6D). Together, these data indicate that *FoxO6* affects *Plxna4* transcription through binding to the identified DBE located within the *Plxna4* promoter.

It has been described that transcriptional activity of FoxO transcription factors is negatively regulated via insulin and insulin-like growth factors (3–5, 13). This would imply that treatment with insulin, or IGFs in general, would negatively regulate the RLUs activated via the *Plxna4* DBE4 enhancer element, thereby resembling the *FoxO6* DNA-binding mutant used in the ChIP experiment described above. To investigate this possibility, we repeated the reporter experiments as described above under conditions with and without serum-mediated repression of *FoxO6* transcriptional activity. To this end, cells were grown in serum-free media and treated with 10% heat-inactivated FCS (repression of FoxO activity) or kept under serum-free conditions (3, 4, 13). Interestingly, luciferase levels significantly dropped to ~27% (Fig. 6D) in 3T3 cells grown under serum-containing conditions, compared with luciferase levels in cells grown under serum-free conditions. These results indicate that serum-mediated signaling can modulate DBE4-driven *Plxna4* transcription.

Discussion

In this study, we established evidence that *FoxO6*, a forkhead transcription factor predominantly expressed during cortical development, affects radial migration. We show that in siRNA-mediated knockdown and in *FoxO6*^{+/-} animals the level of *FoxO6* expression is linked to the expression level of *Plxna4*, a critical component in cortical migration (20). However, in full *FoxO6* knockouts, we were unable to rescue the observed migration phenotype, which was confirmed by the absence of a down-regulation of *Plxna4*. Our data suggest that, although the migration phenotype in *FoxO6*^{-/-} animals resembles the migration phenotype in neurons conditionally silenced for *FoxO6*, the underlying molecular mechanisms differ. The cause could lie in the fact that

FoxO6 expression in the KO is ablated in neural progenitors before E14.5, the time point of the in utero electroporation. This could affect the molecular signature of early progenitors expressing *FoxO6* and as a consequence lead to cortical abnormalities as observed.

Our data demonstrate that in *FoxO6*^{-/-} cortices a subpopulation of E14.5 neurons fail to localize to the correct position in the developing cortex. Although we did not observe a defect in global cortical markers, we did detect a significant difference in the number of Cux1-positive cells in the upper layers, which supports the notion that a small subgroup is affected by ablation of *FoxO6*.

The genome-wide transcriptome analysis performed in this study has identified 24 genes that were deregulated upon silencing of *FoxO6*. Next to *Plxna4*, the down-regulation of *Pik3* catalytic subunit alpha (*Pik3ca*) was of particular interest. In humans, de novo mutations in core components of the PI3K/AKT pathway, such as *Pik3ca*, result in a wide array of cortical malformations, including cortical dysplasia, MPPH, and MCAP (6, 7). These mutations generally lead to a gain of function and activation of the PI3K/AKT pathway, which, in turn, results in the phosphorylation of FoxOs, thereby rendering them inactive (23). Moreover, a subset of patients with mutations in PIK3CA suffers from autism or display autistic features, which may imply a role for *FoxO6* in the development of autism (6).

Plxna4, the coreceptor for *Plxna*/NPI receptors, has been known to be part of the Sema3/6 signaling pathway. The hampered radial migration as observed upon *FoxO6* silencing resembles to a great extent the disrupted migration upon the in utero silencing of *Plxna2* and *Plxna4* (20, 24, 25). This is in line with our hypothesis that *FoxO6* may influence radial migration through *Plxna4* regulation in the developing cortex.

Studies investigating the role of FoxO transcription factors in mitotic regions of the adult mammalian cortex have illustrated the general importance of FoxO factors in maintaining the population of

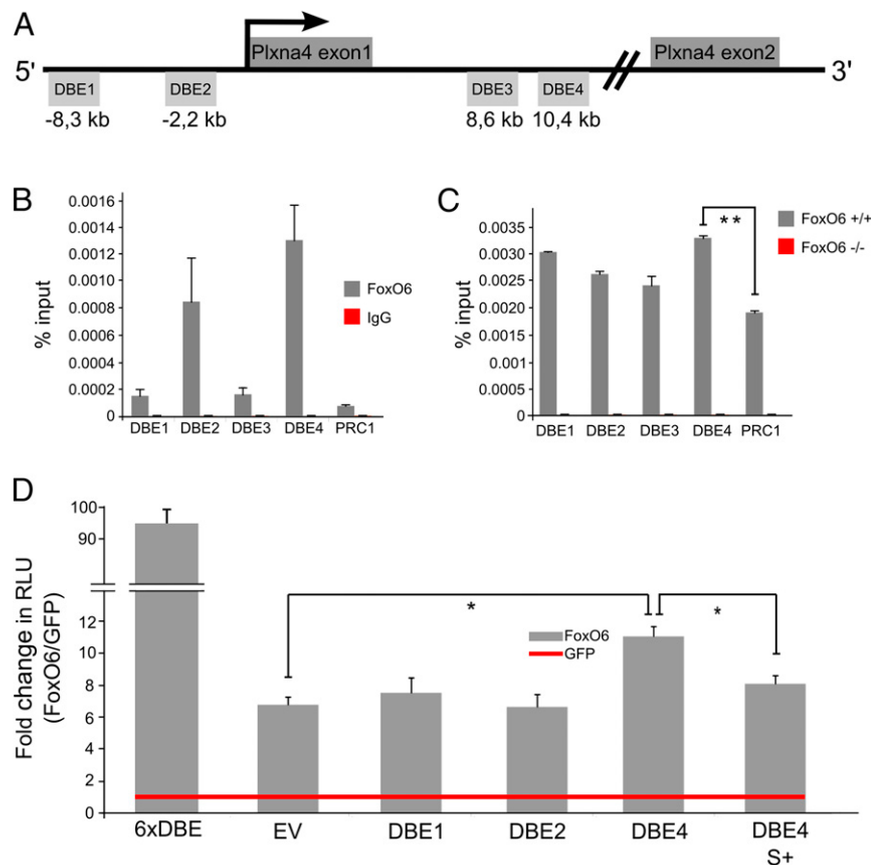


Fig. 6. FoxO6 binds and is able to regulate the *Plxna4* promoter via an identified FoxO6 DNA-binding element. (A) Schematic representation of four putative FoxO DBEs in a 10-kb region surrounding the mouse *Plxna4* transcription start site. (B) In vivo FoxO6 ChIP-qPCR in E16.5 cortices showing enrichment of DBE2 and DBE4 ($n = 2$). (C) FoxO6 ChIP-qPCR showing highest enrichment ($n = 3$) at DBE4 of the *Plxna4* promoter. (D) Dual-luciferase assays in 3T3 cells with the different DBEs showing increased activity for DBE4 ($n = 3$, $P < 0.05$), which is diminished by the addition of serum ($n = 3$). Values are normalized to Renilla. DBE, DAF-16-binding element; S+, serum added. The error bars show the SD. Two-tailed Student's *t* test; * $P < 0.05$ and ** $P < 0.001$.

NSCs (8, 10). Studies in *FoxO3*-deficient mice have shown that adult NSC homeostasis is regulated by FoxO3, mainly via the regulation of genes involved in quiescence (10). In a combined *FoxO1,3,4*-deficient mouse model, a progressive decline in adult NSC numbers was observed, also showing deregulation of genes linked to human brain size and the control of cellular proliferation and differentiation (8). Therefore, it is plausible that in the absence of FoxO6 proliferation and NSCs, homeostasis is altered. The numbers of NSCs and intermediate progenitor cells (IPCs) were unchanged in *FoxO6*^{+/+} and *FoxO6*^{-/-} cortices (Fig. S1), suggesting that this effect is subtle and not likely to contribute to the hampered radial migration.

It is known that FoxO proteins are regulated by the PI3K/AKT signaling pathway. In the presence of insulin or IGFs, FoxO proteins are phosphorylated at specific amino acids (3), resulting in decreased DNA binding and shuttling to the cytoplasm (3–5, 13). It is noteworthy that these upstream repressors of FoxO activity are normally present in embryonic cerebrospinal fluid (CSF) (reviewed in ref. 26). Hypothetically, this hints toward a mechanism in which extrinsic factors, present in the embryonic CSF, regulate the transcriptional activity of FoxO proteins, thereby regulating corticogenesis via downstream target genes such as *Plxna4*.

In addition to the down-regulation of *Plxna4*, our results revealed an altered expression of a diverse set of genes via which FoxO6 potentially executes its functions during corticogenesis. Remarkably, several of these additional identified genes are known to be involved in the process of neuronal migration, including *Dync1h1*, a dynein motor protein, and *Nrcam*, a neuronal cell-adhesion protein associated with psychiatric disorders such as autism (27–29). The expression of *Nfia*, a family member of *Nfix*, which has been shown to be involved in regulating the rostral migratory stream of SVZ-derived neuroblasts (30), was, like *Plxna4*, significantly decreased upon siRNA-mediated knock-down of *FoxO6*.

The molecular basis of radial migration of cortical neurons is a well-studied process showing prominent roles for axon guidance (20,

31, 32), cell adhesion (33, 34), cell polarity (35, 36), and cytoskeleton remodeling (28, 37, 38). Remarkably, knowledge about transcriptional control of genes involved in radial migration is scarce. Transcription factors known to be important in radial migration include *Brn1* and *Brn2*, regulating Cdk5-mediated cortical lamination (39); *Aff3*, controlling Mdg2-mediated cortical migration (40); *RP58*, a zinc finger transcription factor, which negatively regulates Rho family GTPase 2 (*Rnd2*) (41); *COUP-TFI*, which also regulates *Rnd2* during mid to late corticogenesis (42); and *FoxP1*, found to regulate morphogenesis of cortical cells, thereby also critical for neuronal migration (43). In this perspective, the study as presented here is important, as it identifies a player in the transcriptional control of neuronal migration during cortical development.

Materials and Methods

Animals. The C57BL/6J mouse line was used for in utero electroporations silencing *FoxO6* expression and rescuing *Plxna4* expression. Adult *FoxO6* mutant animals (-/-) were backcrossed to the C57BL/6J line and used in heterozygous breeding generating *FoxO6* wild-type (+/+) and mutant progeny (-/-). Embryos were collected at E14.5, E16.5, and E18.5 (E0.5 is defined as day of copulatory plug). Genotyping was performed by PCR analysis using the following primers: forward *FoxO6*-1, 5'-ccagttgtccagctctc-3'; reverse *FoxO6*-1, 5'-cagagccaggtacacgag-3'; and reverse *FoxO6*-2, 5'-ctaaagcgcagtgctccagac-3'. The *FoxO6* wild-type allele produced a band of 257 bp, and the *FoxO6*^{-/-} allele produced a band of 174 bp in a PCR. All procedures were according to and fully approved by the Dutch Ethical Committees for Animal Experimentation (University Medical Center Utrecht and University of Amsterdam).

In Situ Hybridization. In short, coronal brain sections (16- μ m) were collected using a Leica CM3050 S cryostat. Sections were fixed in 4% (wt/wt) paraformaldehyde (PFA) dissolved in PBS and blocked by acetylation. Next, sections were prehybridized for 2 h in prehybridization solution [50% (vol/vol) deionized formamide, 5 \times SSC, 5 \times Denhardt's solution, 250 μ g/mL baker's yeast tRNA, 500 μ g/mL sonicated salmon sperm DNA]. Hybridization was performed overnight at 68 $^{\circ}$ C with 400 ng probe per mL hybridization mix.

The probes used were as follows: *FoxO6*, base pairs 1514–2430 of mouse coding sequence (CDS) and 3' UTR; and *Plxna4*, base pairs 1976–3232 of mouse CDS. The slides were washed in 2× SSC at 68 °C followed by 0.2× SSC at room temperature (RT). Slides were blocked in 10% (wt/wt) heat-inactivated (HI) FCS and incubated overnight at 4 °C with anti-DIG alkaline phosphatase, diluted 1:5,000 in TBS containing 1% HI FCS. Next, the slides were treated with nitro-blue tetrazolium/5-bromo-4-chloro-3'-indolylphosphate solution. Incubation took 4 to 16 h, depending on the probe. The color reaction was stopped in 10 mM Tris-HCl, 5 mM EDTA, and the slides were dehydrated with ethanol and embedded using Entellan.

Immunohistochemistry. E16.5 and E18.0 heads were fixed in 4% (wt/wt) PFA in PBS for 4 and 8 h, respectively, followed by incubation in 30% (wt/wt) sucrose in PBS for at least 48 h. Coronal brain sections (16- μ m) were collected using a Leica CM3050 S cryostat. Antigen retrieval was performed for several antibodies. In short, sections were incubated in 10 mM sodium citrate (pH 6.0) for 15 min at 90 °C. Sections were blocked with 4% (wt/wt) HI FCS in THZT (50 mM Tris, 500 mM NaCl, 0.5% Triton X-100, pH 7.6). Primary antibody was incubated overnight at 4 °C. Primary antibodies and dilutions were as follows: rabbit anti-GFP (1:1,000; ab290; Abcam), chicken anti-GFP (1:1,000; ab13970; Abcam), rabbit anti-Sox2 (1:1,000; ab97959; Abcam), rabbit anti-Tbr2 (1:1,000; ab23345; Abcam), rabbit anti-Ctip2 (1:1,000; ab28488; Abcam), mouse anti-Satb2 (1:1,000; ab51502; Abcam), rabbit anti-Cux1 (1:500; sc-13024; Santa Cruz Biotechnology), and Armenian hamster anti-Plxna4 (kindly provided by F. Suto, National Center of Neurology and Psychiatry, Tokyo). Secondary antibodies were incubated at room temperature for 2 h (Molecular Probes; goat anti-rabbit IgG Alexa Fluor 488; goat anti-rabbit IgG Alexa Fluor 594; goat anti-mouse IgG Alexa Fluor 594; goat anti-Armenian hamster IgG Alexa Fluor 594). Sections were counterstained with DAPI and embedded in FluorSave. Fluorescent images were taken with a Leica microscope (DFC310 FX).

Transfection and Western Blot. To test the efficiency of two siRNAs targeting *FoxO6*, we transfected HEK293 cells, using Lipofectamine 2000, with a plasmid expressing a translational fusion of *FoxO6* and GFP, a GFP expression plasmid, and either siScrambled, siFoxO6-1, or siFoxO6-2. At one day posttransfection, the cells were lysed directly in sample buffer [62.5 mM Tris-HCl, 2% (wt/wt) SDS, 10% (vol/vol) glycerol, 50 mM DTT, 0.01% bromophenol blue] and sonicated for 5 min for optimal lysis. The lysates were centrifuged at 13,300 × g for 20 min. The supernatant was collected and denatured at 95 °C. Proteins were separated by 10% (wt/wt) SDS/PAGE and blotted onto a 0.2- μ m nitrocellulose membrane at 100 V for 2 h at 4 °C. The blot was blocked for 1 h at room temperature in Tris-buffered saline (TBS)-Tween (0.1%) containing 5% (wt/wt) milk, followed by overnight incubation at 4 °C with rabbit anti-GFP (1:50,000; ab290; Abcam). Goat anti-rabbit HRP-conjugated secondary antibody was incubated for 1 h at RT before visualization with ECL detection substrate. After detection of GFP, the blots were washed extensively with TBS-Tween (0.1%). Blots were blocked as described above before incubation with mouse anti- β -actin diluted 1:5,000 in TBS-Tween (0.1%). Goat anti-mouse HRP-conjugated secondary antibody was incubated for 1 h at RT before visualization with ECL substrate. The densitometric analysis was performed using LI-COR Image Studio Lite software.

Quantification and Statistical Analysis. All cell quantifications represent the average values of experiments performed on multiple animals originating from multiple litters, using at least three sections per brain. Data indicate the means with SD unless stated otherwise. GFP-labeled cortices were subdivided into three bins covering the cortical plate, intermediate zone, and ventricular zone. All GFP-positive cells were counted per bin manually. Statistical analysis was performed by Student's *t* test (two-way, unpaired). Bonferroni correction was applied to correct for multiple testing. Sox2, Tbr2, Ctip2, Satb2, and Cux1 countings were performed digitally with ImageJ software (NIH) on 400× magnified thresholded images using optimized parameters. Sox2 quantifications are represented as the percentage of Sox2-positive cells per DAPI-positive cells for a set region covering the VZ and SVZ; *n* = 4 was used for each condition. Tbr2 quantifications are represented as the percentage of Tbr2-positive cells per DAPI-positive cells for a set region covering the SVZ; *n* = 4 was used for each condition. Ctip2 quantifications are represented as the percentage of Ctip2-positive cells per DAPI-positive cells for a set region covering layers V and VI; *n* = 4 was used for each condition. Satb2 quantifications are represented as the percentage of Satb2-positive cells per DAPI-positive cells for a set region covering upper- and deep-layer Satb2-positive pyramidal neurons; *n* = 4 was used for each condition. Cux1 quantifications are represented as the percentage of Cux1-positive cells per DAPI-positive cells for a fixed region covering upper layers II, III, and IV; *n* = 3 was used for each condition. qPCR results represent the average values of experiments performed on four biological samples for each condition, originating from multiple litters.

Data indicate means with SD unless stated otherwise. Statistical analysis was performed by Student's *t* test (two-way, unpaired). The average width of the cortex was measured using ImageJ software. Previously performed DAPI counterstaining was used to identify the width of the indicated layers in E18 *FoxO6^{+/+}* and *FoxO6^{-/-}* brains. The width of the cortex was measured at three fixed locations per section and then averaged per section.

In Utero Electroporation. C57BL/6J mice at 14 d of gestation were anesthetized by injecting a mixture of ketamine/medetomidin/atropine intraperitoneally. The uteruses of these mice were exposed and a mixture containing 1 μ g/ μ L siRNA (siFoxO6-1 or siScrambled), 1 μ g/ μ L expression vector containing GFP and/or *Plxna4*, and 0.01% Fast Green dissolved in 20 mM Tris-HCl (pH 8) was injected through pulled glass capillaries, using a microinjector, into one lateral ventricle of each embryo. We used custom-made siRNAs (Thermo Scientific) to silence *FoxO6* (siFoxO6-2, 5'-NNACAUGCCAGUCAGACUAC-3'; siFoxO6-1, 5'-NNCAUCAGCCUUAUGACGCU-3') and a nontargeting siRNA (Thermo Scientific; D-001210-03-20). The expression vectors used were pCAG-AFP 2424 (GFP) and pcDna3.1 (-) Plexin-A4 (kindly provided by R. J. Pasterkamp, Brain Center Rudolf Magnus, University Medical Center Utrecht). Next, electric pulses were generated by an ECM 830 Square Wave Electroporation System and applied to the cerebral wall at five repeats of 33 V for 50 ms with an interval of 950 ms. After the electroporation the uterus was placed back and the mouse was taken care of. Depending on the analysis, the embryos developed to stage E16.5 or stage E18 before they were killed.

Microarray and qPCR Validation. Embryos were electroporated at E14.5 with either siFoxO6-2 or siFoxO6-1, both targeting *FoxO6* specifically or the control siScrambled. All cortices were cotransfected with GFP. At E16.5, embryos were harvested, and the cortex was isolated and dissociated using a papain kit according to the manufacturer's protocol (Worthington Biochemical) followed by FACsorting. Sorted cells were subjected to RNA isolation using the phenol-chloroform extraction method. Electroporated cortices from different nests were pooled to minimize effects due to electroporation differences and nest differences. All RNA samples were analyzed using a 2100 Bioanalyzer (Agilent Technologies) to ensure the quality of the RNA. Microarray analysis was performed using four experimental samples (three cortices pooled) in utero electroporated with siFoxO6-1 and four experimental samples in utero electroporated with siFoxO6-2. These samples were compared with a reference pool of RNA derived from eight siScrambled electroporated cortices. Per siRNA, two samples were labeled normal and two in a dye swap against the common reference, and hybridized on a mouse expression microarray. Agilent Mouse Whole Genome Gene Expression Microarray V1 (Agilent Technologies) sets were used for all hybridizations, in a 4 × 44 K layout. Hybridized slides were scanned on an Agilent scanner (G2565BA) at 100% laser power, 30% photomultiplier. Data were analyzed using ANOVA (R version 2.2.1/MAANOVA version 0.98-7; www.r-project.org/). *P* values were determined by a permutation F2 test. Genes with *P* < 0.05 after family-wise error correction (or Benjamini–Hochberg correction/FDR control) were considered significantly changed. The data discussed in this publication have been deposited in the National Center for Biotechnology Information's Gene Expression Omnibus (GEO) and are accessible through GEO Series accession no. GSE71954 (www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE71954). Top hits from the array were validated with qPCR. Analysis was performed on a LightCycler 480 II (Roche) using a One Step SYBR Green Kit (Qiagen) according to the manufacturer's protocol. Independent in utero electroporated samples were FACsorted, followed by RNA isolation. Water was used as a nontemplate control. All samples were normalized to an 18S reference. Primers used for qPCR amplification were as follows: 18S (60 °C): forward 5'-aaacggctaccacatccaag-3', reverse 5'-ctccaatggatcctctgta-3'; *FoxO6* (60 °C): forward 5'-agagcgcggcggacaagaga-3', reverse 5'-gccgatggaggtcttcagcc-3'; *Plxna4* (60 °C): forward 5'-gaggatgaacca-gaatggagc-3', reverse 5'-ttctcatgctggcagcgat-3'; *Nfia* (60 °C): forward 5'-agt-gagccgagtggaatg-3', reverse 5'-tggggcagaagtgtctcaatg-3'; *Pik3ca* (60 °C): forward 5'-ttagaacctagtaggaacccg-3', reverse 5'-gctatgagcaggttagatcc-3'; *Xlr3c* (60 °C): forward 5'-gaaacatgcggaacactct-3', reverse 5'-agcgatggaaattccccga-3'; and *Lmx1a* (60 °C): forward 5'-aacacgcgagccaagatgaa-3', reverse 5'-cccgcattccccactaccat-3'.

ChIP-qPCR. In vivo ChIP was performed on E16.5 *FoxO6^{+/+}* and *FoxO6^{-/-}* cortices. Cells were cross-linked using 1% PFA followed by sonication (optimized for neuronal tissue to generate 400- to 1,000-bp DNA fragments). Cross-linked samples were incubated overnight with either rabbit anti-FoxO6 (kindly provided by A. Brunet) or rabbit IgG control followed by incubation with preblocked protein A agarose beads. Beads were extensively washed (nine times) in different salt washing buffers, and DNA–histone complexes were eluted using elution buffer (1% SDS, 0.1 M NaHCO₃). Samples were

reverse-cross-linked and treated with proteinase K. DNA was purified and used for qPCR analysis. qPCR primers are specified below. In vitro ChIP was performed similarly. In short, different mouse cell lines (3T3) were transfected with either pcDNA3.1 FoxO6-Myc or N1-FoxO6-GFP using Lipofectamine 2000 according to the manufacturer's protocol (Thermo Fisher Life Technologies). Cells were cross-linked using 1% PFA followed by sonication. The latter sonication procedure was optimized for each cell line to generate DNA fragments of 200 to 1,000 bp. Cross-linked samples were incubated overnight with either rabbit anti-Myc or rabbit anti-GFP and the rabbit IgG control followed by incubation with pre-blocked protein A agarose beads. Beads were extensively washed in different salt washing buffers, and DNA-histone complexes were eluted using elution buffer (1% SDS, 0.1 M NaHCO₃). Samples were reverse-cross-linked and treated with proteinase K. DNA was purified and used for qPCR analysis. In silico search for FoxO DBEs revealed four FoxO DBEs in the 10 kb surrounding the transcription start site (TSS) of *Plxn4*. Primers used for qPCR amplification were as follows: DBE_plexina4_2.2kbupstream_TSS (60 °C): forward 5'-tgacattcttccctgtca-3', reverse 5'-gtgtgtgtcctgatggatgg-3'; DBE_plexina4_8.6kbdownstream_TSS (60 °C): forward 5'-tcaatctcttccctgatgttg-3', reverse 5'-accacacgtcaatactaccacc-3'; DBE_Plexina4_8.3kbupstream_TSS (60 °C): forward 5'-cccttctctctctcaca-3', reverse 5'-gccactagtgtaagctggaa-3'; DBE_Plexina4_10.4kbdownstream_TSS (60 °C): forward 5'-cccagctcacttaagaaa-3', reverse 5'-ccttcattccccctgaggat-3'; 6xDBE (60 °C): forward 5'-tacggctgtagcccg-3', reverse 5'-gcaggatcaagcttactt-3'.

Luciferase Assay. A total of ~400 bp surrounding the DBEs was independently cloned into the pGL3 promoter vector. 3T3 cells (neuronal cell lines died when overexpressing FoxO6) were grown in 24-well plates and transfected

with 2 µg plasmid DNA per well (N1-FoxO6-GFP or N1-GFP) and cotransfected with 0.2 µg pGL3 promoter vector and 0.02 µg CMV-Renilla using Lipofectamine 2000 (Invitrogen). Twenty-four hours after transfection, cells were serum-starved to activate FoxO6. After 8 h, cells were lysed and assayed for luciferase activities using a Dual-Luciferase Reporter Assay (Promega; E1910) according to the manufacturer's protocol. All experiments were performed in triplicate. Data represent average values.

FACS sorting. Freshly dissected cortices were dissociated using a papain dissociation system (Worthington Biochemical), and cells were sorted on a Cytopeia Influx or BD FACSAria III cell sorter. Sort gates were set on forward scatter versus side scatter (live cell gate), forward scatter versus pulse width (elimination of clumps), and forward scatter versus fluorescence channel 1 (528/38 filter; GFP fluorescence). Cells were sorted using a 100-µm nozzle at a pressure of 15 or 20 psi (BD FACSAria III) with an average speed of 5,000 cells per s and collected in TRIzol-LS reagent (Invitrogen).

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