Cell type specific, traceable gene silencing for functional gene analysis during vertebrate neural development

Citation for published version:
https://doi.org/10.1093/nar/gkr628

Digital Object Identifier (DOI):
10.1093/nar/gkr628

Link:
Link to publication record in Edinburgh Research Explorer

Document Version:
Publisher's PDF, also known as Version of record

Published In:
Nucleic Acids Research

Publisher Rights Statement:
This is an Open Access article distributed under the terms of the Creative Commons Attribution Non-Commercial License (http://creativecommons.org/licenses/by-nc/3.0), which permits unrestricted non-commercial use, distribution, and reproduction in any medium, provided the original work is properly cited.

General rights
Copyright for the publications made accessible via the Edinburgh Research Explorer is retained by the author(s) and / or other copyright owners and it is a condition of accessing these publications that users recognise and abide by the legal requirements associated with these rights.

Take down policy
The University of Edinburgh has made every reasonable effort to ensure that Edinburgh Research Explorer content complies with UK legislation. If you believe that the public display of this file breaches copyright please contact openaccess@ed.ac.uk providing details, and we will remove access to the work immediately and investigate your claim.
Cell type specific, traceable gene silencing for functional gene analysis during vertebrate neural development

Nicole H. Wilson and Esther T. Stoeckli*

Institute of Molecular Life Sciences, University of Zurich, Winterthurerstrasse 190, CH-8057 Zurich, Switzerland

Received June 15, 2011; Revised July 15, 2011; Accepted July 18, 2011

ABSTRACT

Many genes have several, sometimes divergent functions during development. Therefore, timing of gene knockdown for functional analysis during development has to be done with precise temporal control, as loss of a gene’s function at early stages prevents its analysis later in development. RNAi, in combination with the accessibility of chicken embryos, is an effective approach for temporally controlled analysis of gene function during neural development. Here, we describe novel plasmid vectors that contain cell type-specific promoters/enhancers to drive the expression of a fluorescent marker, followed directly by a miR30-RNAi transcript for gene silencing. These vectors allow for direct tracing of cells experiencing gene silencing by the bright fluorescence. The level of knockdown is sufficient to reproduce the expected pathfinding defects upon perturbation of genes with known axon guidance functions. Mixing different vectors prior to electroporation enables the simultaneous knockdown of multiple genes in independent regions of the spinal cord. This permits complex cellular and molecular interactions to be examined during development, in a fast and precise manner. The advancements of the in ovo RNAi technique that we describe will not only markedly enhance functional gene analysis in the chicken, but also could be adapted to other organisms in developmental studies.

INTRODUCTION

The function of a gene can only be understood in the context of the entire organism, as it is cell type specific and context dependent. The analysis of gene function is particularly demanding during development, as temporal control of gene expression adds additional complexity to the experimental approach. While functional gene analysis in vertebrates is most commonly performed by gain-of-function and loss-of-function experiments in the mouse, many genes involved in the development of the nervous system cannot be studied. Since many genes play additional roles in early developmental processes, classical gene knockouts may cause early embryonic lethality that precludes their analysis in later stages of development. In such circumstances, perturbation of gene function in a spatially and temporally defined manner is required. To some extent, this can be achieved by sophisticated gene knockout techniques (1). However, the high cost and the large amount of time required for the production of multiple transgenic mice is rate limiting in the quest to gain insight into the functional role of a gene of interest. Because of its easy accessibility in the egg, the chicken embryo has been the model of choice for developmental in vivo studies. In recent years, our lab and others have used RNA interference (RNAi) and electroporation as a powerful method to perturb gene expression during development. In the chicken, spatiotemporal control of gene knockdown can be achieved by electroporation of long double-stranded RNA (dsRNA) (2–7), DNA plasmids encoding short hairpin RNAs (shRNAs) (8) or artificial microRNAs (miRNAs) (9). These approaches harness the naturally occurring miRNA biogenesis pathway, a post-transcriptional gene regulatory mechanism that activates the RNA-induced silencing complex to evoke gene silencing in response to dsRNA [reviewed in (10)]. We have previously used long dsRNA to identify new roles for multiple genes during neural development (2–7). Alternatively, Das and colleagues (9) described a plasmid in which a ubiquitously active chicken U6 promoter was used to drive the expression of an RNA transcript that mimicked the stem–loop and flanking structure of microRNA30, to ensure its successful processing by Drosha and Dicer. Following electroporation, the authors reported the silencing of various target genes in the chicken neural tube. However, the lack of cell type specificity of these current methods can be problematic since (i) targeting of distinct subpopulations

*To whom correspondence should be addressed. Tel: +41 44 635 4840; Fax: +41 44 635 6879; Email: Esther.Stoeckli@imls.uzh.ch

© The Author(s) 2011. Published by Oxford University Press.
This is an Open Access article distributed under the terms of the Creative Commons Attribution Non-Commercial License (http://creativecommons.org/licenses/by-nc/3.0), which permits unrestricted non-commercial use, distribution, and reproduction in any medium, provided the original work is properly cited.
of cells located in the same area is not possible using manual targeting techniques; (ii) it is difficult to down-regulate different genes in independent subpopulations of cells; and (iii) identification of cells experiencing gene knockdown is often indirect (for example, by the co-electroporation of a fluorescent reporter). Here, we describe an improvement on current techniques to markedly enhance the precision and combinatorial possibilities of in ovo RNAi.

Several recent studies have reported successful gene knockdown by RNAi in a cell type-specific manner, either in vitro [for example (11–13)] or in adult mice using viral vectors or by crossing transgenic lines [for example, (11,14,15)]. In these studies, a cell type-specific RNA polymerase II promoter/enhancer is used to drive the transcription of shRNA. However, many of the approaches described to date do not allow the transfected cells to be directly identified within the target tissues. While this feature may not be necessary for in vitro studies or in therapeutic settings (in which the specific cells experiencing gene knockdown may not need to be identified in the context of the whole tissue or organ), it is crucial for developmental studies. Electroporation of the neural tube results in a transfection efficiency of ~60% of the cells in the electroporated area (2). Therefore, accurate analysis of novel gene function in vivo is only possible when the genetically perturbed cells are identifiable in their environment.

Recent in vitro studies have reported that it is possible for miRNA targeting specific genes to be expressed simultaneously with protein-coding genes (16–18). Thus, we decided to directly couple an artificial miRNA to a fluorescent indicator that would enable the faithful identification of the transfected cells in vivo. We made plasmids driven by RNA polymerase II promoters/enhancers that are specifically active in different subpopulations of cells in the neural tube. These enhancers drive the expression of a single transcript encoding both a fluorescent protein and a single transcript encoding both a fluorescent protein and a sequence (21–22 bp). Novel miRNAs were amplified by PCR using Pfu DNA polymerase and cloned into the first or second hairpin insertion sites in pRFPRNAiC, following the protocol provided with the vector. The target sites for the miRNAs used in subsequent experiments are shown in Table 1.

**Table 1.** Target sequences and vector insertion sites for miRNAs used in this study

<table>
<thead>
<tr>
<th>Name</th>
<th>Gene</th>
<th>Target sequence</th>
<th>Vector insertion site</th>
</tr>
</thead>
<tbody>
<tr>
<td>miShh (9)</td>
<td>Sonic hedgehog</td>
<td>ACAAGAAACTCCGAGAGATTTA</td>
<td>Second</td>
</tr>
<tr>
<td>miLuc (9)</td>
<td>Firefly luciferase</td>
<td>TGCTGTGGGTTGCAACCTATTT</td>
<td>First</td>
</tr>
<tr>
<td>mi2Luc (21)</td>
<td>Firefly luciferase</td>
<td>CGTGATTAGTAGGGGAGGCG</td>
<td>First or Second</td>
</tr>
<tr>
<td>mi1AX1</td>
<td>Axonin1</td>
<td>AAGTACACCAGCTCTAGTGAGA</td>
<td>First</td>
</tr>
<tr>
<td>mi2AX1</td>
<td>Axonin1</td>
<td>AAGTACGCGGCTCTCAGTGAGA</td>
<td>First</td>
</tr>
<tr>
<td>mi3AX1</td>
<td>Axonin1</td>
<td>AAGGACCTTATTGAGTGCAGG</td>
<td>First</td>
</tr>
<tr>
<td>mi4AX1</td>
<td>Axonin1</td>
<td>AAGGAGGCCACCAGGCAACATC</td>
<td>First</td>
</tr>
<tr>
<td>mi5AX1</td>
<td>Axonin1</td>
<td>AAGGAGGCCACCAGGCAACATC</td>
<td>First</td>
</tr>
<tr>
<td>mi6AX1</td>
<td>Axonin1</td>
<td>AACCTTGCATAGCGGAGCAGG</td>
<td>First</td>
</tr>
<tr>
<td>miGPC1</td>
<td>Glypican1</td>
<td>AAGCCTGTAGTGAGAAGTTGAGCC</td>
<td>First</td>
</tr>
<tr>
<td>miWnt17a</td>
<td>Wnt17a</td>
<td>AAGGACGATTCAGGCAACCAG</td>
<td>Second</td>
</tr>
</tbody>
</table>

**Materials and Methods**

**Plasmids**

pRFPRNAiC. pRFPRNAiC (9) was obtained from ARK Genomics and used for many of the initial assessments of miRNA activity in the chicken spinal cord. Artificial miRNAs targeting firefly Luciferase and Shh were previously published (9,21). For each of our other genes of interest, we used the online GenScript siRNA Target Finder to generate at least three new miRNA target sequences (21–22 bp). Novel miRNAs were amplified by PCR using Pfu DNA polymerase and cloned into the first or second hairpin insertion sites in pRFPRNAiC, following the protocol provided with the vector. The target sites for the miRNAs used in subsequent experiments are shown in Table 1.

**Backbone fluorescent constructs.** Vector components were synthesized in a stepwise manner and inserted into a pBluescript SK+ backbone (pBSK+; Stratagene). An SV40 polyA sequence flanked by KpnI and XhoI sites was amplified from pIRES (Clontech) using the following primers: 5'-AATCTCGAGGAGACATGAGAATTACATTGCTTCTACCATGAGGACAGGCC and 5'-AAAGGTACCTGGATTTTACCACATTTTACCACTACTAGTAGGACAGGCC and 5'-AAAGGCACTTATGAGTGCGAGG First or Second. EGFP was amplified from pIRES using primers that incorporated SpeI and EcoRI flanking sites and a β-globin basal promoter (22) upstream of a Kozak start site. The primers used were: 5'-AACTAGTGAGCGTCCGATCGGACGCACTGCACTCAGTGAGAGGTACCTGGATTTTACCACATTTTACCACTACTAGTAGGACAGGCC and 5'-AAAGGCACTTATGAGTGCGAGG First or Second. Enhanced blue fluorescent protein 2 (EBFP2) (23) was excised from pBad-EBFP2 (Addgene plasmid 14891) using XhoI (blunted) and EcoRI, and cloned into the Smal and EcoRI sites of pA/BSK+, creating EBFP2/pA/BSK+. Humanized Renilla GFP (hrGFPII; Stratagene) was cloned into pA/pBSK+ on a BamHI/EcoRV fragment, creating hrGFPII-pA/pBSK+. To enable visualization of axon projections, we used a membrane-targeted (farnesylated) EGFP...
Constructs were bilaterally electroporated at 0.5–1.0 kV cm⁻¹ and TATA box were excised using NotI and SpeI. Constructs were unilaterally electroporated at 0.2–0.5 μg/μl.

We obtained the Hoxa1enIII-TATA-Cre-IRESPApA construct from A. Klar. The 0.85 kb Hoxa-1 promoter element II (containing retinoic acid response elements) and TATA box were excised using NotI and SpeI. Constructs were electrooporated at 0.5–1.0 μg/μl.

The 1.5 kb Math1 enhancer (24) (mouse Atonal homolog; Genbank accession AF218258) was provided in pMath1-Gal4/AD by S. Arber, and was subsequently cloned upstream of the β-globin basal promoter, the EGFP-coding region and SV40 polyadenylation sequences (in EGFP-pA/pBSK⁺). Constructs were electroporated unilaterally at 0.7 μg/μl.

**In ovo electroporation**

Fertilized eggs obtained from a local supplier were incubated at 38.5°C for 3 days prior to windowing. Embryos were staged according to Hamburger and Hamilton (25). Extra-embryonic membranes were carefully removed at HH17-18.5, and the central canal of the spinal cord was injected with a mix of plasmid DNA and Trypan blue to enable visualization of the injection solution. Unilateral transfection was achieved by electroporation (five pulses of 25 V, 50 ms duration) as described previously (2) using a BTX ECM830 square wave electroporator (Harvard Instruments). Bilateral electroporation was performed using five pulses of 18 V, 50 ms duration and then switching the polarity of the electrodes, repositioning and repeating the electroporation. Efficiency and accuracy of electroporation were assessed by the expression of the relevant fluorescent protein/s.

**Immunohistochemistry**

Staining of 25-μm-thick cryostat sections was carried out as described previously (26). Primary antibodies used in this study were as follows: mouse anti-Shh (5E1), mouse anti-myc (9E10), mouse anti-Nkx2.2 (74.5A5), mouse anti-Pax7, mouse anti-Islet1 (40.2D6), mouse anti-HNF3β (4C7), rabbit anti-Axonin1, goat anti-NgCAM, rabbit anti-cleaved caspase-3 (Asp175; Cell Signaling Technology) and goat anti-GFP-FITC (Rockland). All monoclonal antibodies were obtained from Developmental Studies Hybridoma Bank. Fluorophore-conjugated secondary antibodies were purchased from Jackson ImmunoResearch Laboratories (donkey anti-mouse IgG-Cy3, goat anti-mouse IgG-Cy3, donkey anti-rabbit IgG-Cy3 and goat anti-rabbit IgG-Alexa488) or Invitrogen (donkey anti-mouse IgG-Alexa488, donkey anti-goat IgG-Alexa488 and goat anti-mouse IgG-Alexa488).

**Fluorescence intensity recordings**

Electroporated embryos were processed in parallel and immunolabelled for Shh as described. Digital images of red and green fluorescent channels were captured using the same camera settings for all sections within the same experiment. Only those sections where the floor-plate region was successfully targeted (as evident by the expression of RFP or EGFP as appropriate) were used. We analyzed 7–12 sections per condition, taken from at least four different embryos. Fluorescent intensity was calculated using the Gray Value (Mean) function within Adobe Photoshop CS3 software. The area to be sampled was defined by drawing a triangle, starting at the midline just dorsal to the floor plate and extending out to the motor nerve exit points and along the ventral border of the spinal cord. A background recording for each image was taken from the region adjacent to the dorsal root entry zone, where Shh is not expressed. The background recording was subtracted, to give a final fluorescence intensity value for each section. The data were subjected to statistical analysis (see below).

**Efficiency of bilateral in ovo electroporation**

The efficiency of bilateral electroporation was assessed on the basis of expression of hrGFPII. Embryos were bilaterally electroporated with βactin-hrGFPII (0.25 μg/μl) at HH17-18 and sacrificed at HH26. The electroporated area of the spinal cord was removed and dissociated into a single-cell suspension, and the percentage of green cells was determined in a Neubauer chamber. The percentage of hrGFPII-expressing cells was calculated from an average of four embryos ± SEM.

**Quantification of axon guidance phenotypes**

Embryos were sacrificed at HH25-26 and the trajectory of commissural axons was analyzed in an open-book configuration (Figure 4F). In the embryos electroporated with Math1-EGFP constructs, the pathfinding behavior of commissural axons was directly visualized by the expression of membrane-bound EGFP. The scoring of phenotypes was performed by analysis of the caudal part of each open book. The open books were classified as abnormal if a bundle of more than five green axons was present along the ipsilateral floor-plate border. At least six embryos were examined in each condition. Alternatively, Fast-DiI (5 mg/ml; Molecular Probes) was applied by focal injection of DiI into dorsal commissural neurons, as previously described (27). Labeled axons at the midline were documented by confocal microscopy (Leica SP2, or Olympus DSU coupled to BX61 microscope). Only DiI injections sites that were in the appropriate location in the dorsal-most part of the spinal cord, and within the region expressing fluorescent protein, were included in the analysis. As it was impossible to count axons at individual injection sites, the percentage of axons displaying
In situ screening of candidate miRNAs

The specific downregulation of the target genes was verified by performing in situ hybridization using probes produced from chicken cDNAs that were previously generated in the lab, or ESTs obtained from Geneservice. The sequences used corresponded to a 1.5-kb fragment from the 5′-UTR and ORF of Axonin1 (Genbank accession NM_001004395), a 1.4-kb fragment of the 3′-UTR of Glypican1 (similar to ChEST 98608; Genbank accession CR394121.1), a 0.7-kb fragment of Glypican4 (ChEST 335n21), a 0.9-kb fragment of Wnt4 (ChEST 17914) and a 0.7-kb fragment from the ORF and 3′-UTR of Wnt7a (ChEST 543m22; bp 491–1228 of Genbank accession NM_204292). In situ hybridization was carried out essentially as described (28), using digoxigenin-labeled in situ probes.

In vitro screening of candidate miRNAs

We used a multiple transfection protocol, modified from that of Yamamoto and colleagues (29), to maximize the number of cultured cells expressing each miRNA, and then assessed the ability of each miRNA to suppress the expression of its target gene. This approach approximated the production of a ‘stable cell line’ expressing each miRNA, allowing different miRNAs to be rapidly tested side by side. Since the cell machinery required for miRNA biogenesis is highly conserved in eukaryotes (10), we reasoned that COS-7 cells should provide a relevant readout of our chicken sequences. Confluent COS-7 cells were diluted 1:10 in DMEM (Gibco) with 10% fetal calf serum and plated into eight-well Labtek dishes (300 μl/well). Cells were grown at 37°C with 5% CO2 for 60 h before the first transfection. Lipofectamine reagent (0.6 μl/well in 25 μl of OptiMEM) was mixed with each pRFPRNAi vector (125 ng/well in 25 μl of OptiMEM), incubated for 30–45 min, and then added to the cells in 200 μl of DMEM + 10% FCS. Transfected cells were incubated for 6 h, and then transfected again. The following morning, the transfection process was repeated a third time, but this time we also co-transfected a ‘reporter’ pAX1-myc/his construct (50 ng/well), in which a CMV promoter drives the expression of myc/his-tagged chicken Axonin1. Cells were incubated for another 24 h before fixation in 4% PFA and immunolabeling for myc. Briefly, fixed cells were washed in PBS, treated for 20 min with 0.1% Triton-X in PBS, blocked in 2% BSA and incubated for 2 h with 9E10 antibody against myc. Cells were washed in PBS, and then incubated for 1 h with goat anti-mouse IgG, conjugated to Alexa488, before washing and mounting in Mowiol reagent.

We used the expression of RFP in each well to assess the relative transfection rates of the miRNA-expressing vectors and found that ~70% of the cells in each well were transfected. We observed no differences in the transfection rates between different wells within the same experiment. The level of myc staining was used as an indicator of the efficiency of each miRNA to knockdown Axonin1, using the ‘empty vector’ condition (pRFPRNAi without a miRNA) as a reference. In addition to the six specific miRNAs targeted against Axonin1, we used a miRNA against Glypican1 (an unrelated gene) as a negative control. Myc staining in each well was visually scored as ‘no knockdown’, ‘weak knockdown’ or ‘strong knockdown’. miRNAs eliciting ‘weak’ or ‘strong’ knockdown were tested further in vivo. All experiments were batch processed in parallel to minimize variability between wells and repeated three times to confirm the results.

Statistical analysis

For the Shh immunofluorescence intensity recordings, we analyzed 7–12 sections, from at least four embryos per condition. The average fluorescence intensity of Shh was calculated for each condition and the data were subjected to ANOVA analysis and the Tukey’s HSD test. Average percentage values provided in the text were normalized to the untreated control condition (100%), and are expressed ± SEM.

For analysis of axon guidance phenotypes observed in DiI injection sites, raw data were subjected to a one-tailed Fisher’s exact test and chi-square test of association. At least 6 embryos and 50 DiI injection sites were analyzed for each condition.

We used the VassarStats Website for Statistical Computation (Vassar College; ©Richard Lowry 1998–2011; http://faculty.vassar.edu/lowry/VassarStats.html).

RESULTS

RNA polymerase II promoters can drive effective, traceable gene knockdown in vivo

In chicken, previous studies have introduced synthetic shRNAs via plasmids driven ubiquitously by RNA polymerase III (9,30,31). While the effective gene silencing reported in these studies is encouraging, the use of RNAi vectors described to date do not allow for directly identifiable, cell type-specific expression of shRNAs. Thus, we tested whether an RNA pol II promoter could be used to drive the effective expression of a single transcript encoding both a fluorescent protein (to directly mark transfected cells) and artificial miRNA (Figure 1A).

We first tested this idea using a ubiquitously active (RNA pol II-driven) βactin promoter, and a previously published miRNA against Sonic Hedgehog (9) (miShh: Figure 1A). The miShh and flanking regions were placed within the context of a miRNA30-based sequence (9) and
cloned within the 3′-UTR of red fluorescent protein (RFP). Shh is a secreted protein that is produced and released by floor-plate cells, thus forming a gradient in the ventral spinal cord (Figure 1B and C). When a control β-actin-RFP plasmid was injected into the central canal of Hamburger–Hamilton (HH) stage 18 chicken embryos followed by bilateral electroporation, we saw no significant change in the average intensity of Shh immunofluorescence at HH26 compared to untreated embryos (Figure 1E; higher by 7 ± 10.8% SEM). However, when β-actin-RFP-miShh was electroporated, we saw a 42 ± 12.5% reduction in Shh (Figure 1G; \( P = 0.0015 \)). Electroporation with either the β-actin-RFP or β-actin-RFP-miShh plasmid produced strong red fluorescence making it easy to identify transfected cells in the spinal cord (Figure 1D and F). We calculated that the transfection efficiency of our bilateral electroporations is 51.8% ± 1.3%. Therefore, an average knockdown of 42% at the protein level represents ~80% of the theoretical maximum that could be achieved. Since any Shh expressed prior to electroporation would be unaffected, the results suggest that our miRNA approach very effectively downregulates Shh in the targeted cells.

To test whether the knockdown of Shh was due to the specific activity of the miRNA rather than global gene downregulation, we immunolabeled sections with antibodies against an array of spinal cord markers, including HNF3b, Nkx2.2, Islet1 and Pax 7 (32,33). Although Shh is a morphogen that specifies cellular differentiation during early neural tube development, perturbation of Shh signaling after HH18 no longer affects spinal-cord patterning (3). Indeed, we found that the expression patterns and staining intensities of the markers that we used were indistinguishable between the different groups (Figure 2). Our results showed that the forced, widespread expression of miShh after HH18 did not cause gross developmental abnormalities, and that miShh downregulates its target gene specifically.

We further assessed the versatility of our plasmid design by expressing different fluorescent proteins [EBFP2, hrGFPII and EGFP (data not shown)] and novel miRNAs against a variety of endogenous target genes (Figure 3). In each case, the plasmids expressed strong fluorescence and elicited knockdown of their targets. The expression of a miRNA against firefly luciferase (9) had no effect on the levels of the endogenous genes that we tested (Figure 3G and H), indicating that our harnessing of the cellular miRNA processing pathway did not in itself elicit broad physiological toxicity. In support of this, immunolabeling for activated caspase-3 showed no differences in cell death between control and miRNA-treated tissues (data not shown). Thus, our plasmids elicit traceable, effective gene knockdown in vivo.

Cell type-specific, traceable gene knockdown in the floor plate

We next tested whether a cell type-specific RNA Pol II promoter/enhancer could be used to drive traceable gene knockdown. We used the floor-plate-specific Hoxa-1 enhancer element III (34) to drive the expression of EGFP ± miShh (Figure 4A). Following bilateral electroporation, both Hoxa1-EGFP and Hoxa1-EGFP-miShh

Figure 1. RNA polymerase II-driven plasmids can be used to express a single transcript encoding both a fluorescent protein and an effective artificial miRNA in chicken embryos. (A) Schematic of the plasmid constructs used in this study. The stem–loop structure of the miRNA30-like cassette is shown for an artificial miRNA against Shh (miShh). The target (sense) sequence of Shh is indicated in red. (B) Immunolabeling of cryosections of untreated chicken spinal cords at HH26 with the 5E1 antibody shows normal distribution of Shh in the notochord (nc), floor plate (asterisks) and adjacent ventral spinal cord. The boxed area is shown in higher power in (C), and equivalent regions are shown in (D–G). (D and F) RFP-positive cells in the floor plate and ventral spinal cord of embryos bilaterally electroporated with the control β-actin-RFP construct (D) or β-actin-RFP-miShh (E). The corresponding images showing Shh expression in each condition are shown in (F) and (G) (compare to C). Scale bar: 100 μm.
embryos displayed green fluorescence specifically in the floor plate (Figure 4B and insets). Relative to untreated embryos (Figure 4C), we found no significant change in the average Shh immunofluorescence intensity in the Hoxa1-EGFP group (Figure 4D; higher by 10 ± 5.6%), whereas the average Shh intensity was reduced by 40% ± 19.1% following Hoxa1-EGFP-miShh treatment (Figure 4E; *P* = 0.0006). Immunolabeling for other spinal cord markers showed no differences between the three groups (data not shown). Taken together, these results suggest that the Hoxa-1 enhancer III can drive both bright fluorescence and active miRNA to elicit traceable gene knockdown in the chicken floor plate.

In order to assess their effectiveness in developmental studies, we next asked whether this level of knockdown was sufficient to perturb biological function. After its function in neural tube patterning, Shh plays two roles during the guidance of spinal commissural axons: first, it collaborates with Netrin1 to attract axons toward the floor plate (35) and second, it repels post-crossing axons...
along the rostrocaudal axis (3). We examined the guidance of commissural axons using open-book preparations (Figure 4F and G) from embryos electroporated with Hoxa1-EGFP ± miShh. In untreated control embryos, 81.3% of all DiI injection sites (n = 166, from 17 embryos) displayed the normal phenotype in which commissural axons crossed the floor plate before turning sharply at the contralateral floor-plate border and extending rostrally (Figure 4H). This normal phenotype was also seen in the majority of DiI sites from Hoxa1-EGFP embryos (67.0%; n = 115, from 20 embryos). However, the normal phenotype was observed in only 39.7% of DiI sites from Hoxa1-EGFP-miShh embryos (n = 58, from 13 embryos; \( P = 0.0005 \) compared to Hoxa1-EGFP). The specific phenotypic abnormalities that we saw in the miShh group were...
consistent with the loss of Shh during commissural axon guidance: 24.1% of injection sites showed ipsilateral pathfinding errors (compared to 9.6% in Hoxa1-EGFP embryos; \( P = 0.0109; \) Figure 4I and K), and post-crossing axons failed to make a turning decision into the longitudinal axis in 31.1% of injection sites (compared to 15.7% in Hoxa1-EGFP controls, \( P = 0.0169; \) Figure 4I–K). In contrast to these results, the bilateral electroporation of Hoxa1-EGFP-miLuc did not cause axon pathfinding abnormalities compared to controls electroporated with Hoxa1-EGFP alone (65.9% normal phenotype; \( n = 141 \) DiI sites from 24 embryos; \( P = 0.4865 \)), demonstrating that the phenotypes were not due to non-specific toxicity. Thus, the cell type specific, targeted knockdown that is elicited by the plasmids is sufficiently potent to perturb physiological gene function.

**Commissural neuron-specific gene knockdown**

To establish the efficiency of the miRNA-expressing vectors on other gene targets and in other cell types, we used an enhancer of the *Math1* (mouse Atonal homolog) gene to drive expression of a membrane-targeted EGFP, followed by a miRNA against Axonin1 (AX1; also known as TAG-1 and Contactin2). The 1.5 kb *Math1* enhancer drives specific expression in the commissural dI1 interneurons of the spinal cord (24,36). The majority of *Math1*-labeled axons (~85%; *Math1*\textsuperscript{comm}) cross the floor plate (24) and project rostrally within the medial longitudinal fascicle (MLF), before being deflected diagonally and eventually re-converging within the intermediate longitudinal fascicle (ILF). Another later-born class of *Math1*-labeled neurons (~15%; *Math1*\textsuperscript{ipsi}) occupies a more ventral position. Their axons project ipsilaterally, turning rostrally in the intermediate spinal cord without contacting the floor plate (36,37) (Figure 5).

We have previously demonstrated that commissural axons require Axonin1/TAG-1 (AX1) for correct pathfinding (2,38). Axonally expressed AX1 interacts with midline-derived NrCAM to enable axons to enter the floor plate. When AX1 function is perturbed by dsRNA-induced downregulation or by the addition of function-blocking antibodies, >20% of commissural axons stall or turn at the ipsilateral floor-plate border (2,38). Thus, we used this well-characterized phenotype as an indicator of efficient downregulation of AX1 specifically in commissural axons using miRNAs driven by the *Math1* enhancer.

We first identified miRNAs that successfully targeted AX1. We designed six miRNAs (mi1AX1-mi6AX1) against the ORF of AX1, using an online algorithm. These were cloned into pRFPRNAi (9) and first screened ***in vitro*** to rapidly identify effective candidates (Figure 6). We used a multiple transfection protocol (29) to approximate a ‘stable cell line’ of cultured cells expressing each miRNA. Based on the expression of an RFP reporter, we estimated that 70% of cells were transfected and we saw similar transfection rates in all wells. Next, a myc-tagged AX1 construct was introduced. Cells were incubated for 24 h before immunolabeling for myc and comparing the expression levels to that observed in the control ‘empty’ pRFPRNAi condition. Of the six miRNAs that we tested, three showed strong suppression of AX1 (mi1AX1, mi2AX1 and mi5AX1), two showed weak knockdown (mi3AX1 and mi4AX1) and one was ineffective ***in vitro*** (mi6AX1). A miRNA against an unrelated gene (miGPC1) did not affect AX1 expression.

For ***in vivo*** testing, we unilaterally electroporated pRFPRNAi-AX1 vectors, or pRFPRNAi-miLuc as a control. By ISH (Figure 7A and B), we found that mi3AX1 elicited the most significant knockdown of AX1, while mi1AX1, mi2AX1 and mi4AX1 were moderately effective (data not shown). Surprisingly, mi5AX1, which strongly suppressed AX1 ***in vitro***, appeared to be ineffective ***in vivo***. The expression of miLuc had no impact on AX1 levels. Immunolabeling for AX1 yielded the same results (Figure 7C and D), suggesting that mi3AX1 was an effective miRNA to knockdown AX1 ***in vivo***. The expression of NgCAM (another immunoglobulin superfamily protein expressed by commissural axons) was unaffected by either miLuc or mi3AX1, demonstrating the specificity of knockdown (Figure 7E and F).

We next tested whether the knockdown of AX1 by mi3AX1 was adequate to elicit axon guidance effects.
in vivo, and further, whether the specific population of commissural axons labeled by the Mathl enhancer requires AX1 for floor-plate crossing (Figure 8). We cloned mi3AX1 or miLuc into the 3'UTR of EBFP2, in a β-actin-driven plasmid. This was co-electroporated with a plasmid containing the Mathl enhancer driving a membrane-bound EGFP (EGFPf), to genetically label dI1 axons (Figure 8A). In the control conditions in which Math1-EGFPf was co-electroporated with β-actin-EBFP2 or with β-actin-EBFP2-miLuc, the majority of embryos displayed Math1-labeled (green) axons that grew normally across the floor plate before turning longitudinally (6/7 and 12/13 embryos, respectively; data not shown). In contrast, we found that the co-electroporation of Math1-EGFPf with β-actin-EBFP2-mi3AX1 caused many Math1-labeled axons to stall or turn before crossing the floor plate, creating a fascicle of green axons at the ipsilateral floor-plate border (6/6 embryos). The position of this aberrant longitudinal fascicle was distinct from (and in addition to) that of the Math1ipsi axons that normally extend within the intermediate spinal cord. The result phenocopied the axon guidance defects that we previously reported as a result of perturbations in AX1 activity (2,38). Assessment of individual DiI injection sites in open-book preparations showed ipsilateral errors in 56% of DiI sites (n = 99 from six embryos) in the β-act-EBFP2-mi3AX1 group (Figure 8D), compared to 4% of the DiI sites (n = 51 from six embryos) in the β-act-EBFP2 group or 7.5% of DiI sites (n = 53 from six embryos) in the β-act-EBFP2-miLuc group (P < 0.0001). Taken together, the results show that mi3AX1 efficiently downregulates AX1 in vivo and confirm that a subpopulation of the Mathlcomm axons requires AX1 for correct guidance.

We next tested whether the Mathl enhancer itself could drive sufficient expression of fluorescent protein and mi3AX1, specifically in the dI1 neurons, to recapitulate these guidance defects. We cloned mi3AX1 or miLuc into the 3'UTR of Math1-EGFPf and electroporated these constructs (Figure 8A). In the control conditions electroporated with Math1-EGFPf alone or with Math1-EGFPf-miLuc, the majority of embryos [66.7% (n = nine embryos) and 69.2% (n = 13), respectively] displayed green fluorescent axons with the normal pathfinding phenotype (Figure 8C and E). In contrast, 68.2% of Math1-EGFPf-mi3AX1 embryos (n = 22) displayed green axons that turned aberrantly along the ipsilateral floor-plate border, as described above, thus
recapitulating the expected axon guidance defects arising from perturbation of AX1 function (Figure 8F; $P = 0.0455$ compared to miLuc group).

Interestingly, we observed that the severity of the axon pathfinding defects in the Math1-EGFPf-mi3AX1 embryos was dependent on the age at electroporation: 87% of all moderate/severe phenotypes were observed in embryos electroporated at $<\text{HH17.5}$, while 67% of all normal phenotypes were found in embryos electroporated at $>\text{HH18}$. Within each embryo, we also found that the caudal segment of the open book was more severely affected than the rostral end, reflecting the decreasing ‘maturity’ of the nervous system in the rostral-caudal axis.

Analysis of individual Dil sites in open-book preparations (Figure 8D) revealed that ipsilateral stalling or turning errors (Figure 8G) were observed in $\approx 9\%$ of DiI sites following electroporation with Math-EGFPf ($n = 142$ from 16 embryos) or Math1-EGFPf-miLuc ($n = 61$ from nine embryos). In contrast, 26.9% of DiI sites ($n = 182$ from 20 embryos) in the Math1-EGFPf-mi3AX1 group displayed ipsilateral errors ($P = 0.0011$ compared to the miLuc group). Of note here is that we inject Dil into the cell bodies of the Math1comm, but not the Math1ipsi, subpopulation of dI1 neurons (Figure 5). Additionally, we routinely label only commissural axons that project within the MLF, in contact with the contralateral floor-plate border, and not those that project in the ILF. Thus, our Dil injection sites label only a sub-population of the Math1comm axons, which may explain the lower percentage of ipsilateral errors that we observed in the Dil analysis of Math1-EGFPf-mi3AX1 embryos compared to the $\beta$act-EBFP2-mi3AX1 group.

Dual activity of vectors in vivo

The plasmids contain insertion sites for two different miRNAs. Previous reports demonstrate that both miRNA insertion sites in the parent pRFPRNAi vectors are active against exogenous gene targets in vitro (9,39). In the single knockdown experiments described above, we used either the first miRNA insertion site (for mi3AX1) or the second (for miShh), demonstrating that, individually, both sites can yield miRNAs that are active against endogenous gene targets. To test whether the insertion sites are simultaneously active in vivo, we used a single construct encoding two miRNAs against different target genes (Figure 9). The miRNA cassette was located in the 3′-UTR of hrGFPII, in a $\beta$actin-driven construct.

We arbitrarily chose target genes whose expression patterns partially overlapped in the developing neural tube. The first insertion site contained a miRNA against the heparan sulfate proteoglycan glypican1 (GPC1), while the second contained a miRNA against Wnt7a, yielding $\beta$actin-hrGFPII-miGPC1-miWnt7a. To control for saturation of the miRNA processing pathway, we made dual expression constructs containing a miRNA against luciferase (mi2Luc), rather than an endogenous gene target, in the first or second insertion sites (yielding $\beta$actin-hrGFPII-miGPC1-mi2Luc and $\beta$actin-hrGFPII-mi2Luc-miWnt7a). The constructs were electroporated, and the expression of $\beta$actin, GPC1, Wnt7a and two closely related members of the target gene families (GPC4 and Wnt4) was assessed 2 days later. In each case, we found that the appropriate target gene/s was/were knocked down without any discernible effect on the expression of non-targeted genes. The level of silencing for each gene was comparable to that produced by vectors.
encoding only a single miRNA (Figure 3). Thus, both miRNA insertion sites are simultaneously active in vivo and their concurrent activity can drive effective, specific downregulation of distinct target genes.

Mixing two plasmids before electroporation can additionally be used to knockdown genes in different cell types. Due to the expression of fluorescent markers from the plasmids encoding the different miRNAs, the cells experiencing knockdown of the specific target genes can easily be identified (Figure 10).

**DISCUSSION**

We have designed a simple, vector-based artificial miRNA expression strategy that can be used to knockdown endogenous gene expression in a cell type-specific, traceable manner in vivo. All constructs that we tested strongly express a fluorescent protein marker and can efficiently silence target gene expression in vivo. While RNA pol II-driven shRNAs have previously been shown to elicit cell type-specific knockdown (11,13,15), the current study is the first to demonstrate in vivo that an artificial miRNA directly coupled to a fluorescent protein marker performs effectively during development. These functional tools offer multiple gene silencing, temporal control and cell type specificity, to facilitate the elucidation of complex developmental pathways.

We tested three different RNA polymerase II promoters/enhancers to drive expression either ubiquitously (β-actin promoter) or in a cell type-specific manner.
Math1 enhancer, Hoxa-1 enhancer element III). The success of the cell type-specific approach offers a number of benefits over current techniques. First, enhancer-mediated control of gene knockdown provides an increased precision of cell type-specific targeting compared to manual techniques. Since the artificial miRNA is expressed only in the cell type of interest (and not in surrounding tissues), there is a decreased chance of abnormal phenotypes arising secondarily as a result of local morphological and/or cell fate changes. Second, the use of RNA pol II promoters, which are generally weaker than those driven by RNA pol III, decreases the chance of non-specific toxicity arising from...
saturation of the miRNA-processing pathway (11). Finally, the combined use of constructs driven by different promoters enables the simultaneous knockdown of different genes in independent regions of the neural tube from a single electroporation event, thus enhancing the combinatorial possibilities of in vivo RNAi (Figure 10). In the neural tube, several other enhancers/promoters have been shown to drive cell type-specific expression, for example, in motor neurons [HB9; (40)], primary sensory neurons [CREST3; (41)], subpopulations of dorsal interneurons (37) and other cell types (42). Thus, the plasmid design may facilitate many developmental studies of diverse subsets of neurons and their targets.

Several different fluorescent reporters were all strongly expressed and can be easily exchanged in each construct. This feature facilitates the concurrent use of two or more different constructs in vivo since their simultaneous activity in independent cell types can be identified with different fluorescent markers. This is of critical relevance to developmental studies examining highly intricate cellular and molecular pathways. The ability to simultaneously knockdown multiple different genes in independent cells types, combined with the ability to directly identify the cells experiencing knockdown of particular gene/s, offers a significant advantage over current techniques. The direct coupling of the fluorescent marker with the artificial miRNA within a single transcript also allows more accurate assessment of targeted cells, compared to commonly used designs where the miRNA and marker are encoded by different transcripts that are derived from independent plasmids or promoters. The ability to easily exchange the fluorescent protein that is expressed also enhances the flexibility of the plasmid application in conjunction with other fluorescent labeling methods (e.g. DiI labeling and fluorescent immunohistochemistry). Previous studies have shown that a miRNA precursor located in the 3′-UTR only moderately inhibits expression of a linked, protein-coding ORF (due to processing of the primary transcript by Drosha) (16,17). Consistent with these findings, we had no problems detecting fluorescence in any of our experiments.

A crucial factor in the use of miRNA-based plasmids as functional tools in developmental biology is the selection of effective, specific miRNAs (43). This process should incorporate the identification of several independent miRNAs to confirm an observed phenotype, negative controls (scrambled and/or unrelated miRNAs) and rescue constructs. To facilitate this process, we have described a quick in vitro protocol as a primary screening method of novel miRNAs. The multiple transfection of a given miRNA-expressing plasmid maximizes the number of cells expressing the miRNA, approximating a ‘stable cell line’ into which other plasmids (providing a readout of knockdown efficiency, off-target effects or cross-reactivity) can be co-transfected. We found that one of our miRNAs against AX1 was effective in vitro but not in vivo. While this finding was unexpected, it is consistent with another previous report (44) and underscores that in vitro efficacy does not necessarily apply in vivo. Still, our in vitro protocol provides a platform for further in vivo testing of novel miRNAs.

In our AX1 knockdown experiments, we observed decreasing phenotypic effects with increasing age at electroporation. Importantly, these effects are entirely consistent with the onset of AX1 expression [HH19 in the lumbar spinal cord (38)] and its long half-life >100 h in DRG neurons (45)]. The findings support the idea that any AX1 protein that is expressed prior to the introduction of the miRNA is sufficient and persistent enough to allow correct axon pathfinding. They also illustrate an important consideration for the timing of application of the plasmids in vivo: following transfection, the transcription and processing of ‘active’ miRNAs capable of eliciting gene knockdown may take several hours.

Our results show that two endogenous genes can be targeted from a single vector. This provides a number of advantages for functional analyses in vivo; for example, by addressing redundancy in gene families, or enabling combinatorial gene knockdowns to identify genetic interactions. The simultaneous use of multiple cell type-specific vectors would enable many different genes to be knocked down in independent regions of the spinal cord, to investigate complex cellular and molecular interactions in a manner that is rapid, precise and inexpensive.

Finally, the possibility to fluorescently label a few genetically modified cells in a ‘wild-type’ environment will also provide a basis for time-lapse imaging of navigating axons or migrating cells in organotypic cultures. The combination of the novel tools with ex ovo RNAi (5) will allow for analysis of cell migration, axon guidance and synapse formation in the brain (in particular in the cerebellum) in a temporally controlled and cell type-specific manner. In contrast to such approaches in mice, where cell type-specific promoters driving Cre expression must be used, spatial and temporal control of gene silencing is facilitated in the chicken using these novel tools. Additionally, since our results suggest that, in principle, any RNA pol II promoter could be used, the applicability of the construct design may extend beyond that of the chicken neural tube.

ACKNOWLEDGEMENTS

We thank C. Krull for providing the chicken β-actin promoter, A. Klar for providing the Hoxa1 enhancer III and TATA box elements and S. Arber for providing the Math1 enhancer. We are grateful to I. Andermatt for critical reading of the manuscript.

FUNDING

The Swiss National Science Foundation (to E.S.). Funding for open access charge: Institute of Molecular Life Sciences, University of Zurich.

Conflict of interest statement. None declared.

REFERENCES


