A Dimerized HMX1 Inhibits EPHA6/epha4b in Mouse and Zebrafish Retinas

Fabienne Marcelli¹,², Gaëlle Boisset¹, Daniel F. Schorderet¹,²,³

IRO – Institute for Research in Ophthalmology, Sion, Switzerland, 1 Faculty of Life Sciences, Swiss Federal Institute of Technology (EPFL), Lausanne, Switzerland, 2 Faculty of Biology and Medicine, University of Lausanne, Lausanne, Switzerland

Abstract

HMX1 is a homeobox-containing transcription factor implicated in eye development and responsible for the oculo-aural syndrome of Schorderet-Munier-Franceschetti. HMX1 is composed of two exons with three conserved domains in exon 2, a homeobox and two domains called SD1 and SD2. The function of the latter two domains remains unknown. During retinal development, HMX1 is expressed in a polarized manner and thus seems to play a role in the establishment of retinal polarity although its exact role and mode of action in eye development are unknown. Here, we demonstrated that HMX1 dimerized and that the SD1 and homeodomains are required for this function. In addition, we showed that proper nuclear localization requires the presence of the homeodomain. We also identified that EPHA6, a gene implicated in retinal axon guidance, is one of its targets in eye development and showed that a dimerized HMX1 is needed to inhibit EPHA6 expression.

Introduction

Homeobox-containing transcription factors represent an important class of factors involved in the regulation of embryogenesis and other molecular programs. HMX1 is a homeobox-containing transcription factor implicated in eye development. In 1992, Stadler et al. described a new homeobox gene called GH6. This gene was later renamed HMX1 and was assigned to the NKX5 family, the reason why HMX1 is also known as NKX5-3 [1]. Later, further members were identified: HMX2 (NKX5-2), HMX3 (NKX5-1) and, in chicken, zebrasfish and medaka, SOHo-1 [2–4]. The NKX5/HMX family of transcription factors contains a unique homeobox region that is phylogenetically conserved. HMX1, HMX2 and HMX3 contain two other conserved domains called SD1 and SD2, located immediately C-terminally to the homeobox [5]. The function of these domains is still unknown.

Whereas Hmx2 and Hmx3 play a role in inner ear development, Hmx1 and SOHo-1 are mainly implicated in eye development. In the mouse eye, Hmx1 expression can be detected as early as E10.5, and transcripts are more specifically present in the lens and in the antero-medial part of the neural retina [4–8]. In the developing chicken eye, it is expressed in the dorsal neural retina and lens epithelium as well as in the optic nerve [9]. HMX1 expression starts 40 hours into development (stage 11) in the surface ectoderm surrounding the optic vesicle. At optic cup invagination (stage 14–15), it is expressed in the anterior/nasal side of the early retina [10]. In zebrafish, hmx1 is first expressed in the entire eye at 10 somite-of-stage (ss), and is then repressed in the dorsal part at 18 ss. At 24 hours post fertilization (hpf), it is restricted to the nasal retina and, one day later, expression is restricted to the nasal part of the ganglion cell layer (GCL). At four and five days post fertilization, signal is also observed in the nasal part of the inner nuclear layer (INL). In the developing lens, expression is observed from 24 to 72 hpf [11,12].

We recently reported a family with a 26-bp deletion in exon 1 of HMX1 leading to the oculo-aural syndrome of Schorderet-Munier-Franceschetti (OMIM: 612109), characterized by microphthalmia, microcornea, nystagmus, cataract, coloboma, optic nerve dysplasia, RPE abnormalities, rod-cone dystrophy and deformation of the ear lobe [12,13]. A mouse model containing a mutation in Hmx1 has been described [6]. It shows laterally protruding ears, subtle changes in cranial bone morphology, perinatal semi-lethality, reduced body mass and microphthalmia with low-grade keratoconjunctivitis sicca and entropion. The eyes show no evidence of microcornea, anterior segment dysgenesis, cataract, coloboma, retinal detachment or retinal dysplasia. Quina et al. observed a significant reduction of geniculate ganglion neurons [7]. In vitro, HMX1 binds to a 5’-CAAGTG-3’ sequence, represses transcription from a luciferase reporter containing this binding site and can antagonize Nkx2.5, a cardiac homeo protein, which is activating this same reporter construct [14]. Nks2.5 is also known to dimerize at its homeodomain and other regions in the C-terminus [15].

In this study, we showed that HMX1 acted as a dimer and that the homeobox and the conserved domain SD1 were needed for dimerization to occur. SD2 was not involved in the dimerization process. We also identified EPHA6 as a target of HMX1 and showed that HMX1 repressed the EPHA6 promoter in vitro. The inhibitory activity of HMX1 was associated with the presence of the SD1 and homeobox domains. Whereas the EPHA6 inhibition was lost with mutants of each of these 2 domains, the SD2 mutant showed a small activation of the EPHA6 promoter. Mutation of...
the three CAAG(TG) sequences of the promoter attenuated the repression by HMX1. This inhibition was confirmed in vivo in zebrafish embryos.

Materials and Methods

Plasmid Constructions
Subcloning was performed according to standard protocols. Mutagenesis was performed using the QuickChange II Site-Directed Mutagenesis Kit (Stratagene, Agilent Technology AG, Basel, Switzerland). The sequence of the primers used in this study is available from the authors.

Cell Culture and Transfection
Human embryonic kidney (HEK) 293T cells were cultured at 37°C and in 5% CO2 atmosphere, in Dulbecco’s Modified Eagle’s Medium (DMEM) high glucose with stable glutamine (GE-Healthcare, Glattbrugg, Switzerland), supplemented with 10% FBS (Lonza, Basel, Switzerland), 100 U/ml penicillin and 100 μg/ml streptomycin (Invitrogen, Basel, Switzerland). Trans-
Infection was performed using the Calcium Phosphate method (ProFection Mammalian Transfection System, Promega, Dubendorf, Switzerland).

BRET2

200'000 HEK 293T cells in DPBS were distributed into black 96-well microplates for fluorescence quantification. Filter sets were adapted to 485 nm for GFP excitation and 510 nm for emission. Cells expressing BRET2 donor (RLUC) alone were used to determine the fluorescence background. 200'000 cells with comparable fluorescence levels were distributed into white 96-well microplates for luminescence quantification. The luciferase substrate Coelenterazine 400A, DeepBlueC (Chemie Brunschwig, Basel, Switzerland) was added to a final concentration of 5 mM. Filter sets were adapted to 410 nm for luciferase emission and 515 nm for GFP emission. The emitted fluorescence and luminescence were measured using an Envision 2103 Multilabel Reader (PerkinElmer, Schwerzenbach, Switzerland), and analyzed with the Wallac Envision Manager V1.12 software (PerkinElmer, Schwerzenbach, Switzerland).

Co-immunoprecipitation

200 μg of proteins were immunoprecipitated overnight at 4°C on a rotating wheel with 2.5 μl anti-Renilla Luciferase antibody (MAB4400, Millipore, Zug, Switzerland). 20 μl of washed protein G plus agarose beads (Santa Cruz, LabForce AG, Nunningen, Switzerland) were added and incubated 2 hrs at 4°C on the rotating wheel. After centrifugation at 4°C, the supernatants were kept as controls. The pellets were resuspended in 25 μl 2x SDS loading buffer and loaded on a 12% SDS-page gel, alongside with 20 μl of supernatant and 40 μg of proteins.

Western Blot

Proteins were extracted from cell cultures using RIPA (50 mM Tris-HCl pH 8.0, 150 mM NaCl, 1% NP-40, 0.5% sodium deoxycholate, 0.1% SDS) and concentrations measured using the Micro BCA Protein Assay Kit (Thermo Fisher Scientific, Reinach, Switzerland) on a Multiplate Reader Synergy HT (Bio-Tek, Luzern, Switzerland) with the KC4 software. The following antibodies were used: HA-Tag (6E2) Mouse mAb #2367 (Cell Signaling, LabForce AG, Nunningen, Switzerland), GFP N-terminal G1544 (Sigma, Buchs, Switzerland), PARP (46D11) Rabbit mAb #9532 (Cell Signaling, Labforce AG, Nunningen, Switzerland), Ub (A-5) sc-166553 (Santa Cruz Biotechnology, LabForce AG, Nunningen, Switzerland) and α-Tubulin Clone B-5-1-2 T5168 (Sigma, Buchs, Switzerland).
Native Western Blot

Cells were lysed in a non-denaturing lysis buffer (20 mM Tris-HCl, 137 mM NaCl, 10% glycerol, 1% Triton X-100, 2 mM EDTA, pH 8.0). The protein concentrations were measured as described above. 5 μg were loaded on a Mini-Protean TGX precast gel 4–15% (BioRad Laboratories AG, Cressier, Switzerland) with non-denaturing loading buffer (300 mM Tris-HCl pH 7.8, 30% glycerol, 0.6% bromophenol blue) and migrated without denaturation in a running buffer without SDS.

GFP² Fluorescence Imaging and Nuclei Isolation

Cells were analyzed 48 hrs post-transfection under a Zeiss Axiovert 200 microscope with filters adapted for excitation and emission at λex = 480 nm and λem = 510 nm, respectively, and the AxioVision 4.2 software. For nuclei isolation, cells were counted and resuspended at 10⁶ cells/ml Nuclei Isolation Buffer (250 mM sucrose, 20 mM Hepes pH 7.8, 10 mM KCl, 1.5 mM MgCl₂, 0.5 mM spermidin). Cells were then homogenized with a Potter and spread on a slide.

In Silico Search for a Nuclear Localization Signal (NLS)

The mouse HMX1 sequence was entered in the NLS-Mapper software that can be found at http://nls-mapper.iab.keio.ac.jp/cgi-bin/NLS_Mapper_form.cgi.

Immunofluorescence

Immunofluorescence was performed 24 hrs post transfection. When necessary, 50 μM chloroquine were added for 16 hrs. The primary antibody (LC3B #2775, Cell Signaling, LabForce AG, Nunningen, Switzerland) was diluted in 1x PBS +2% NGS +0.2% Triton X-100 and incubated overnight at 4°C in a humid chamber. The secondary antibody (Alexa Fluor 594 goat α-rabbit IgG (H+L) (A11012), Molecular Probes, LubioScience, Luzern, Switzerland) was diluted in the same buffer, and incubated 1 hr at
RT in a humid chamber in the dark. Nucleic acids were stained with 100 μM DAPI (4,6-diamidino-2-phenyl-indole HCl) (1/1'500 in 1x PBS) for 10 min in a humid chamber in the dark. Cells were then mounted with Citifluor AF1 (Citifluor Ltd, Leicester, UK), and conserved at 4°C. The slides were analyzed under an Olympus BX61 microscope and the CellM software (Olympus, Volketswil, Switzerland).

Hoechst-PI Staining
20 mg/ml bisBenzimide H 33342 trihydrochloride (Sigma, Buchs, Switzerland) and 1 mg/ml Propidium Iodide (Fluka, Buchs, Switzerland) were diluted 1/29000 into the culture medium. Cells were analyzed under a Zeiss Axiovert 200 microscope and the AxioVision 4.2 software.

Luciferase Assays
48 hrs post transfection cells were washed with 1x PBS, and 300 μl luciferase assay lysis buffer (100 mM K2HPO4, pH 7.8, 0.2% Triton X-100) were added. Cells were scraped at 4°C and centrifuged 3 min at 12'000 rpm at 4°C. 5 μl of supernatant were transferred to a transparent 96-well plate containing 50 μl 2x β-gal buffer (120 mM Na2HPO4, 80 mM NaH2PO4, 2 mM MgCl2, 100 mM β-mercaptoethanol). 50 μl of 2x ONPG (1.33 mg/ml 2-nitrophenyl-B-D-galactopyranoside) were added and the plate read at 412 nm of absorbance on a Multiplate Reader Synergy HT (Bio-Tek, Luzern, Switzerland) with KC4 software. If the values were constant in all conditions, 5 μl of supernatants were transferred to a white 384-well plate, 20 μl of Luciferase Assay Reagent (Promega, Dubendorf, Switzerland) were added and luminescence measured on the Multiplate Reader Synergy HT every 3 minutes until the peak of luciferase activity was reached. The obtained values were normalized using a β-gal reporter under the control of a CMV promoter. A mean between the 3 highest values was used for the luciferase/β-gal ratio. Each experiment was performed three times in duplicates. Only transfections with stable β-gal values between the different conditions, indicating similar transfection efficiency, were used. Two-tailed Student’s T-tests with unequal variance were used to determine statistical differences between the conditions.

Chromatin Immunoprecipitation
All experiments involving live animals were authorized by the Veterinary Service of the State of Valais under authorizations N° VS-13 and VS-19. The litter of four was housed with the mother and was anesthetized with isoflurane prior to being euthanized by cervical dislocation. Retinas from four 2-week-old wild-type C57Bl/6j mice were dissected, fixed, and homogenized. Glycine was added to a final concentration of 0.125 M before centrifugation.
Figure 5. Action of HMX1 on the EPHA6 promoter. Schematic representation of the subcloned fragment of the EPHA6 promoter with the three binding sites (red arrows and black characters), and the mutated sequences (red characters) (A). Luciferase assay on the wt EPHA6 promoter with HMX1, HMX1 del SD1, HMX1 del SD2 and HMX1 del HD. HMX1 inhibits the promoter by 42%, HMX1 del SD1 and HMX1 del HD have no effect, and
HMX1 del SD2 slightly activates the promoter (B). Chromatin immunoprecipitation on 2-week-old CS78I/6J retinas demonstrated the physical interaction between HMX1 and the EphA6 promoter. 5% BSA was added in the control conditions instead of the Hmx1 antibody. TIC = total input chromatin (C). Mutation of the HMX1 binding sites attenuates the effect of HMX1 and HMX1 del SD2 but does not completely abolish it (D). Data points represent the mean of three experiments +/- SD. **: P<0.01, *: P<0.05 (Student’s T-test).
doi:10.1371/journal.pone.0100096.g005

Whole-mount in situ Hybridization

Standard one-color whole-mount in situ hybridization was performed at various stages. Hybridization reaction was done at 68°C for 14-18 hrs. Washing steps and antibody incubation were performed in an in situ machine (BioLane HTI, Holle&Huttner, Tubingen, Germany). Templates used to generate DIG-labeled RNA probes included zebrafish hmx1 (ID: 797503), epha4b (ID: 64270) and pax6 (ID: 60639). In situ transcription was done with the Roche RNA Labeling Kit (Roche Applied Science, Basel, Switzerland).

Results

HMX1 dimerizes through the SD1 and homeobox domains

The ability to homo- or heterodimerize has been demonstrated for many transcription factors, including the NKX member NKKX2-5, a cardiac homeobox gene that dimerizes through its homeodomain (HD) [15]. We therefore investigated, using a BRET2 approach, whether HMX1 behaved similarly and formed dimers or oligomers. The BRET2 technique is based on the energy transfer occurring between the renilla luciferase (RLUC) and the green fluorescent protein (GFP) when they are in close proximity. The principle of the technique is to generate fusion proteins between proteins of interest and the RLUC and the GFP, and measure the energy transfer in culture conditions to determine if the proteins of interest are interacting.

When plasmids expressing fusion proteins between RLUC and HMX1, and GFP2 and HMX1 were mixed and transfected in HEK293T cells, a robust increase in the BRET2 ratio was observed with increasing concentrations of GFP2-HMX1, indicating that HMX1 dimerized (Figure 1A). We confirmed dimerization of HMX1 by co-immunoprecipitation (co-IP) using an RLUC antibody for immunoprecipitation and a GFP antibody for blotting. Out of the six conditions tested, the only condition in which immunoprecipitation occurred was when the two different HMX1 fusion proteins were present (Figure 1B). Non-denaturing electrophoresis was also used to further confirm this result. As no western-blot suitable antibody against HMX1 existed, we tagged HMX1 with an HA-tag and used antibodies against HA to visualize the fused HA-HMX1 protein. HA-tagged wild-type HMX1 proteins were loaded on a non-denaturing native electrophoresis gel and sizes were compared to a denaturing gel after western blot analysis with an anti-HA antibody. The size of HMX1 proteins were loaded on a non-denaturing native electrophoresis gel and sizes were compared to a denaturing gel after western blot analysis with an anti-HA antibody. The size of HMX1 proteins were loaded on a non-denaturing native electrophoresis gel and sizes were compared to a denaturing gel after western blot analysis with an anti-HA antibody.

In order to determine the dimerization domain of HMX1, we generated deletions of various portions of the protein. HMX1 is composed of two exons, with three conserved domains in exon two: the homeobox (HD), and two domains called SD1 and SD2, located 3’ to the HD and whose function is presently unknown. We deleted each of these domains separately by site-directed mutagenesis and repeated the BRET2 experiments. As shown in figure 2A, deletions of HD or SD1 led to the loss of dimerization, whereas deletion of SD2 had no effect. This suggested that HD and SD1 were implicated in the dimerization of HMX1. In order

---

Generation of the Zebrafish Hsp70-HMX1 Transgenic Line

AB zebrafish were raised and kept under standard laboratory conditions at 28.5°C. Transgenesis was performed by generating Tol2 transposon constructs using the to2kit [16]. The zebrafish hmx1 coding sequence was cloned downstream of the hsp70 promoter and the DNA construct together with the transposase mRNA were injected at the one-cell stage. Fish were raised to adulthood and the cardiac GFP expression was used as a marker for germline transmission. Experiments were done on F3 obtained from F2 that were intercrossed in order to increase the number of larvae carrying the transgene. Tg [hsp70: hmx1] and wt were heat shocked at 1 dpf during 30 min at 39°C, euthanized and fixed 4 hrs after.

---

Whole-mount in situ Hybridization

Standard one-color whole-mount in situ hybridization was performed at various stages. Hybridization reaction was done at 68°C for 14-18 hrs. Washing steps and antibody incubation were performed in an in situ machine (BioLane HTI, Holle&Huttner, Tubingen, Germany). Templates used to generate DIG-labeled RNA probes included zebrafish hmx1 (ID: 797503), epha4b (ID: 64270) and pax6 (ID: 60639). In situ transcription was done with the Roche RNA Labeling Kit (Roche Applied Science, Basel, Switzerland).

Results

HMX1 dimerizes through the SD1 and homeobox domains

The ability to homo- or heterodimerize has been demonstrated for many transcription factors, including the NKX member NKKX2-5, a cardiac homeobox gene that dimerizes through its homeodomain (HD) [15]. We therefore investigated, using a BRET2 approach, whether HMX1 behaved similarly and formed dimers or oligomers. The BRET2 technique is based on the energy transfer occurring between the renilla luciferase (RLUC) and the green fluorescent protein (GFP) when they are in close proximity. The principle of the technique is to generate fusion proteins between proteins of interest and the RLUC and the GFP, and measure the energy transfer in culture conditions to determine if the proteins of interest are interacting.

When plasmids expressing fusion proteins between RLUC and HMX1, and GFP2 and HMX1 were mixed and transfected in HEK293T cells, a robust increase in the BRET2 ratio was observed with increasing concentrations of GFP2-HMX1, indicating that HMX1 dimerized (Figure 1A). We confirmed dimerization of HMX1 by co-immunoprecipitation (co-IP) using an RLUC antibody for immunoprecipitation and a GFP antibody for blotting. Out of the six conditions tested, the only condition in which immunoprecipitation occurred was when the two different HMX1 fusion proteins were present (Figure 1B). Non-denaturing electrophoresis was also used to further confirm this result. As no western-blot suitable antibody against HMX1 existed, we tagged HMX1 with an HA-tag and used antibodies against HA to visualize the fused HA-HMX1 protein. HA-tagged wild-type HMX1 proteins were loaded on a non-denaturing native electrophoresis gel and sizes were compared to a denaturing gel after western blot analysis with an anti-HA antibody. The size of HMX1 proteins were loaded on a non-denaturing native electrophoresis gel and sizes were compared to a denaturing gel after western blot analysis with an anti-HA antibody. The size of HMX1 proteins were loaded on a non-denaturing native electrophoresis gel and sizes were compared to a denaturing gel after western blot analysis with an anti-HA antibody.

In order to determine the dimerization domain of HMX1, we generated deletions of various portions of the protein. HMX1 is composed of two exons, with three conserved domains in exon two: the homeobox (HD), and two domains called SD1 and SD2, located 3’ to the HD and whose function is presently unknown. We deleted each of these domains separately by site-directed mutagenesis and repeated the BRET2 experiments. As shown in figure 2A, deletions of HD or SD1 led to the loss of dimerization, whereas deletion of SD2 had no effect. This suggested that HD and SD1 were implicated in the dimerization of HMX1. In order

---

Generation of the Zebrafish Hsp70-HMX1 Transgenic Line

AB zebrafish were raised and kept under standard laboratory conditions at 28.5°C. Transgenesis was performed by generating Tol2 transposon constructs using the to2kit [16]. The zebrafish hmx1 coding sequence was cloned downstream of the hsp70 promoter and the DNA construct together with the transposase mRNA were injected at the one-cell stage. Fish were raised to adulthood and the cardiac GFP expression was used as a marker for germline transmission. Experiments were done on F3 obtained from F2 that were intercrossed in order to increase the number of larvae carrying the transgene. Tg [hsp70: hmx1] and wt were heat shocked at 1 dpf during 30 min at 39°C, euthanized and fixed 4 hrs after.

---

Whole-mount in situ Hybridization

Standard one-color whole-mount in situ hybridization was performed at various stages. Hybridization reaction was done at 68°C for 14-18 hrs. Washing steps and antibody incubation were performed in an in situ machine (BioLane HTI, Holle&Huttner, Tubingen, Germany). Templates used to generate DIG-labeled RNA probes included zebrafish hmx1 (ID: 797503), epha4b (ID: 64270) and pax6 (ID: 60639). In situ transcription was done with the Roche RNA Labeling Kit (Roche Applied Science, Basel, Switzerland).
**Figure 6. Regulation after hmx1 misexpression in zebrafish.** Expression of hmx1 after heat shock in wt and tg (hsp70:hmx1) embryos (A, B). Hmx1, normally restricted to the nasal retina, lens and ear (arrows in A), was broadly expressed in the transgenic embryo (B). Epha4b expression after heat shock in wt and Tg (hsp70:hmx1) embryos (C–F). Dissected eye showed a strong reduction of epha4b expression in the temporal retina when HMX1 was co-expressed (E, F). Pax6 regulation after HMX1 misexpression in zebrafish. Ocular expression of pax6 in wt (G) was not modified by overexpression of HMX1 in Tg(hsp70:hmx1) embryos (H). Scale bars: 100 μm.

doi:10.1371/journal.pone.0100096.g006
to confirm these results and to show that the HMX1 C-terminal region was not involved in dimerization, in contrary to that of NKX2-5, we generated serial deletions of the C-terminal part of the protein. None of these constructs prevented dimerization as shown by BRET2 (Figure 2B).

The Entire HD is Necessary for Correct Nuclear Localization of HMX1

Fusing HMX1 to the GFP2 reporter allowed us to visualize its cellular localization. GFP alone localized to the cytoplasm (Figure 3A), whereas GFP-HMX1 localized to the nucleus (Figure 3B). All of the generated mutants retained this nuclear localization except one (Figure 3C–J). The HD deletion mutant was expressed in a punctate manner in the nucleus as well as in the cytoplasm (Figure 3K, L, O, P). This punctate phenotype could possibly be due to the loss of a nuclear localization signal located in the homeobox. We therefore tested the sequence for potential nuclear localization signals (NLS) using NLS mapper, a bioinformatic tool available online. The analysis of HMX1 revealed the presence of a monopartite NLS -RGGRRKKTRTVF-, with KKTRTVF corresponding to the very beginning of the HD, with a score of 9.5. Deleting this signal could thus explain why the GFP-HMX1 del HD protein lost its nuclear localization. To test this hypothesis, we reintroduced the seven-amino acid KKTRTVF into the HMX1 del HD sequence. However, reintroducing these amino acids did not modify the punctate expression and localization of this mutant (Figure 3M). To verify if the predicted NLS needed additional amino acids to be functional, we generated a new mutant with a deletion of the C-terminal half of HD (30 amino acids). However, this construct was still expressed in a punctate manner similar to the deletion of the entire HD (Figure 3N).

The homeobox of HMX1 is of helix-turn-helix-loop-helix type. It is likely that removal of any part of this structure prevents the correct folding of the protein and that it activates clearance mechanisms. In an effort to determine the nature of the aggregates generated by the GFP-HMX1 del HD mutant, we tested several hypotheses. First, the shape, size and distribution of the aggregates suggested that they could be autophagosomes induced by the abundant expression of aberrant proteins. We therefore verified if GFP-HMX1 del HD colocalized with LC3B by immunofluorescence, but this was not the case (Figure 4A, B, C). Even after treating cells with chloroquine to visualize autophagosomes, GFP-HMX1 del HD did not colocalize with autophagosomes. To confirm this result, we also tested whether p62 and ubiquitin expression was increased in the presence of GFP-HMX1 del HD. The role of ubiquitin is to clear abnormal proteins by targeting them for degradation by the 26S proteasome. Poly-ubiquitinated protein aggregates are also sequestered in inclusion bodies containing p62, and the aggregates are cleared via autophagy. In our experiments, we did not observe any increase in expression of these two proteins, indicating that these mechanisms were not activated (Figure 4D). To determine whether the cells were suffering from the presence of GFP-HMX1 del HD aggregates, we looked for the presence of increased apoptosis by PARP cleavage assay. Cleavage of PARP by Caspase-3 is a step in the cascade leading to apoptosis. However, we failed to show any such increase (data not shown). Moreover, no increased cell death was observed when performing a Hoechst-PI staining for dying cells (data not shown). The exact nature of these punctae could thus not be determined and we do not know at this time whether they represent pure HMX1 aggregates or a more complex structure.

HMX1 Binds to the Promoter of EPHA6/epha4b and Inhibits its Expression

HMX1 and SOHO-1 are defining the EPHA3 expression domain in the developing chick retina [10]. Ephrins act as topographically specific repulsive guidance cues for ganglion cell axons. EPHA3 is expressed in a temporal>nasal gradient in the developing chick retina and is present on ganglion cell axons during the time of target innervations. HMX1 and SOHO-1 are expressed in an inverted gradient to that of EPHA3 (nasal>temporal), and when HMX1 and SOHO-1 are expressed ectopically, EPHA3 expression is lost. EPHA3 thus appeared to be a good candidate as a target for HMX1. However, ephrins do not have the same patterns of expression and do not play the same roles between different species. In the ganglion cell layer, where Hmx1 is expressed on the nasal side, EphA5 and EphA6 (P0 mouse) and EPHA3 (chicken) are only expressed on the temporal side [10,17,18]. Chicken EphA5 and EphA6 are uniformly expressed in the chick retina, and EphA3 is not expressed in the mouse retina CGL [17,19–21]. Therefore mouse EphA5 and EphA6 seem to be functional homologs of chicken EPHA3, which suggests that HMX1 could repress the activity of the EphA5 or EphA6 promoter in mouse. In zebrafish, epha4b is expressed in the same temporal pattern as chicken EPHA3 and mouse EphA5 and EphA6, whereas epha6 is not expressed in the eye (not shown).

Amendt et al. showed that HMX1 was preferentially binding to a CAAC(TG) sequence [14]. The EPHA6 promoter contains three such binding sites, the second being conserved between human and mouse (−39 relative to the ATG), whereas the EPHA5 promoter does not contain any. EPHA6 was also identified as a potential target of HMX1 using a predictive promoter model that we recently developed [22]. We therefore analyzed the effect of HMX1 on the human EPHA6 promoter. The technique we used allowed measuring the activity of the promoter by placing a luciferase reporter under its control. We subcloned a fragment spanning from −150 to +150 nucleotides relative to the EPHA6 translation initiation codon into a luciferase reporter vector. This fragment represented the minimal EPHA6 promoter with a 13-fold increased activity compared to pGL3-basic vector. Shorter fragments (−100 to +150 and −50 to +150) displayed reduced promoter activity (five- and three-fold increased activity, respectively, compared to pGL3-basic vector) whereas the +1 to +150 fragment displayed no promoter activity (data not shown). The measured activity values were normalized using a β-gal reporter under the control of a CMV promoter. The subcloned fragment contained three potential binding sites for HMX1: one CAATGTG in the forward direction at position −39, one CAAG in the forward direction (−20) and one CAAG in the reverse direction (−65) (Figure 5A). We performed luciferase assays with wild-type HMX1 and the mutants deleting the HD, SD1 or SD2. As shown in figure 5B, HMX1 was inhibiting the EPHA6 promoter activity by 42%. The physical interaction between HMX1 and the EphA6 promoter was demonstrated by chromatin immunoprecipitation on retinas isolated from two-week-old C57BL/6j mice (Figure 5C), a technique allowing to determine which proteins bind to a DNA fragment by crosslinking them and selecting for the fragments bound to the protein by immunoprecipitation and PCR amplification of the fragment. We also validated this interaction in vivo on the zebrafish epha4b gene, the functional homolog of EPHA6, having two HMX1 binding sites in its promoter. We generated a transgenic fish line expressing a ubiquitous heat-shock activated hmx1 gene and showed by in situ analysis that the aberrant ectopic expression of hmx1 in the temporal retina reduced the expression of epha4b (Figure 6A–F), which was not the case for the control gene pax6 (Figure 6G–H).
We then checked whether dimerization was needed for HMX1 repressive activity. Mutant constructs preventing dimerization, i.e. deletions of HD or SD1, had no activity, indicating that only dimerized HMX1 regulates EPHA6 expression. The mutant with a deleted SD2 domain slightly activated the EPHA6 promoter, suggesting that this region might represent the binding site of a cofactor needed for the inhibitory activity of HMX1 (Figure 5B). In order to confirm that the CAAG/CAAGTG sites represented bona fide binding sites for HMX1, we mutated them into the sequences shown in red in figure 5D. Control experiments showed that these mutations did not affect EPHA6 promoter activity (data not shown). Transfection experiments using these mutated constructs showed a reduction of the inhibitory activity of HMX1 from 42% to 22% (Figure 5D). This indicates that the CAAG/CAAGTG sites represent true binding sites for HMX1. However additional sites might exist, as deletion of CAAG/CAAGTG sites failed to completely abrogate the inhibition.

Discussion

The interest in the HMX1 transcription factor has surged with the discovery in 2008 that it was causing the oculo-auricular syndrome of Schorderet-Munier-Franceschetti [12]. In addition of being expressed in somatosensory organs [23], Hmx1 has been shown to retain a neuronal fate in migrating neural crest cells [24] and to modulate the adrenergic/cholinergic program of sympathetic neurons [25]. It is also well expressed in sensory spinal and cranial ganglia [9]. In C. elegans, the Mbs-2 gene, a member of the HMX family, regulates cytoskeletal organization and cell elongation [26]. However, few contributions have been published about its mode of action in the eye. We therefore investigated its role in eye development.

We showed that HMX1 exerts an inhibitory effect on EPHA6 and that dimerization is necessary for this activity. Luciferase assays are known for producing artefactual results. By increasing the number of replicates and analyzing only the experiments where all conditions showed similar transfection efficiencies, we were able to obtain stable results, which were further confirmed by ChIP and re-experiments in zebrafish. Mutations that removed the dimerization domains of HMX1, i.e. the HD and SD1 domains, abolished its inhibitory efficiency on EPHA6 promoter. Removing the HD also perturbed the cellular localization of HMX1, which was no longer restricted to the nucleus. All other mutants, including deletion of the SD1 domain, maintained a strict nuclear expression indicating that SD1 is involved in dimerization while the HD is necessary both for dimerization and nuclear localization. In addition to ChIP validation in mouse retina, we also showed that ectopic overexpression of HMX1 in the whole eye in a zebrafish transgenic animal in which expression of HMX1 was under a heat-shock-inducible promoter was accompanied by a reduction of epha4b ocular expression, the zebrafish functional homolog of EPHA6.

The role of the SD2 domain remains unknown. We showed that a deletion mutant, which was dimerizing normally, was not inhibiting the EPHA6 promoter like the wild type protein, but was slightly activating it, instead. This conserved domain could thus be an interaction site for a cofactor necessary for the inhibition action of HMX1.

When deleting the homeobox, we observed that the GFP-HMX1 fusion protein lost its specific nuclear localization, and became expressed in a punctate manner in the nucleus as well as in the cytoplasm. Our first hypothesis was that the protein lacking the homeobox was misfolded, and therefore activated clearance mechanisms, either by autophagy or by the proteasome degradation system. The homeobox of HMX1 has a well defined helix-loop-helix-turn-helix tertiary structure type. It is possible that deleting it entirely or part of it changes the three-dimensional structure enough to activate the clearance mechanisms for misfolded proteins. However, we could not detect any indication that these mechanisms were triggered. One of the main components of autophagosomes is LC3B, and we therefore tested whether it colocalized with GFP-HMX1 del HD. This was, however, not the case even after blocking autophagy using a chloroquine treatment. We did not observe an increase in expression of ubiquitin and p62, confirming that the HMX1 aggregates were not autophagosomes, and that the proteasome was not activated. The HMX1 del HD aggregates did not induce cell death either as we observed no increase in PI-stained cells compared to other transfections (not shown). Moreover, PARP was not cleaved by Caspase 3 in GFP-HMX1 del HD transfections, indicating an absence of apoptosis (not shown). Thus, we do not know at this time what is the exact nature of these GFP-HMX1 del HD aggregates and if they represent pure HMX1 aggregates or a more complex structure.

In a previous study, we showed that a morpholino-based knockdown of zebrafish hmx1 had no effect on retinal patterning [11], which is in contradiction to the results obtained previously in the chick retina [10] and the results presented here. However, in chicken the relationship between HMX1 and EFHA3 was shown by overexpressing HMX1 on the temporal side of the retina where it is not expressed normally. The same procedure was used in the current work. In our previous study, hmx1 was knocked-down on the nasal side of the retina, whereas the temporal part was unaffected by this procedure, and epha4b was able to play its role in the temporal retina.

In summary, we showed that HMX1 exerts its inhibitory activity through a dimer and identified EPHA6 as a target of HMX1. Identifying other targets will allow us to further understand the role of HMX1.

Acknowledgments

We thank Des Nathalie Allaman-Pillet, Sandra Cottet, Pascal Escher and Raphael Roduit from the Institute for Research in Ophthalmology for their help and critics, Mr Cédric Schöpfer, Mrs Tatiana Favez and Céline Agosti for technical help, and Dr Nicole Renner and Mrs Suzan Houghton for editing the manuscript.

Author Contributions

Conceived and designed the experiments: FM GB DFS. Performed the experiments: FM GB DFS. Analyzed the data: FM GB DFS. Wrote the paper: FM GB DFS.

References

1. Stadler HS, Padanilam BJ, Bartos K, Murray JC, Solursh M (1992) Identification and genetic mapping of a homeobox gene to the 4p16.1 region of human chromosome 4. Proc Natl Acad Sci U S A 89: 11579–11583.
2. Adamska M, Wolff A, Kreuder M, Wittbrodt J, Braun T, et al. (2001) Five Nkx5 genes show differential expression patterns in anlagen of sensory organs in medaka: insight into the evolution of the gene family. Dev Genes Evol 211: 338–349.
3. Stadler HS, Murray JC, Loya EJ, Goodfellow PJ, Solursh M (1995) Phylogenetic conservation and physical mapping of members of the H6 homeobox gene family. Mamm Genome 6: 383–388.
4. Wang W, Lo P, Frasch M, Lufkin T (2000) Hmx: an evolutionary conserved homeobox gene family expressed in the developing nervous system in mice and Drosophila. Mech Dev 99: 125–137.
5. Yoshiura K, Leysens NJ, Reiter RS, Murray JC (1998) Cloning, characterization, and mapping of the mouse homeobox gene Hmx1. Genomics 50: 61–68.
6. Munroe RJ, Prabhu V, Acland GM, Johnson KR, Harris BS, et al. (2009) Mouse H6 Homeobox 1 (Hmx1) mutations cause cranial abnormalities and reduced body mass. BMC Dev Biol 9: 27.
7. Quina LA, Tempest L, Hsu YW, Cox TC, Turner EE (2012) Hmx1 is required for the normal development of somatosensory neurons in the geniculate ganglion. Dev Biol 356: 152–163.
8. Wang W, Van De Water T, Lafkin T (1998) Inner ear and maternal reproductive defects in mice lacking the Hmx1 homeobox gene. Development 125: 621–634.
9. Stadler HS, Solursh M (1994) Characterization of the homebox-containing gene GH6 identifies novel regions of homeobox gene expression in the developing chick embryo. Dev Biol 161: 231–262.
10. Schulte D, Cepko CL (2000) Two homeobox genes define the domain of EphA3 expression in the developing chick retina. Development 127: 5033–5045.
11. Boisset G, Schorderet DF (2012) Zebrafish hmx1 promotes retinogenesis. Exp Eye Res 105: 34–42.
12. Schorderet DF, Nichini O, Boisset G, Polok B, Tisler L, et al. (2008) Mutation in the human homeobox gene NKX5-3 causes an oculo-auralic syndrome. Am J Hum Genet 82: 1178–1184.
13. Vaclavik V, Schorderet DF, Borrut FX, Munier FL (2011) Retinal dystrophy in the oculo-auralic syndrome due to HMX1 mutation. Ophthal Genet 32: 114–117.
14. Amendt BA, Sutherland LB, Russo AF (1999) Transcriptional antagonism between Hmx1 and Nkx2.5 for a shared DNA-binding site. J Biol Chem 274: 11633–11642.
15. Kasahara H, Usheva A, Ueyama T, Aoki H, Horikoshi N, et al. (2001) Characterization of homo- and heterodimerization of cardiac Cox/Nkx2.5 homeoprotein. J Biol Chem 276: 4570–4580.
16. Kwan KM, Fujimoto E, Grabber C, Mangum BD, Hardy ME, et al. (2007) The Tol2kit: a multistate gateway-based construction kit for Tol2 transgenesis constructs. Dev Dyn 236: 3088–3099.
17. Brown A, Yates PA, Burrola P, Orruto D, Vaidya A, et al. (2008) Topographic mapping from the retina to the midbrain is controlled by relative but not absolute levels of EphA receptor signaling. Cell 102: 77–88.
18. Feldheim DA, Vanderhaeghen P, Hansen MJ, Friens J, Lu Q, et al. (1998) Topographic guidance labels in a sensory projection to the forebrain. Neuron 21: 1303–1313.
19. Cheng HJ, Nakamoto M, Bergemann AD, Flanagan JG (1995) Complementary gradients in expression and binding of ELF-1 and Mek4 in development of the topographic retinotectal projection map. Cell 82: 371–381.
20. Connor RJ, Menzel P, Pasquale EB (1998) Expression and tyrosine phosphorylation of Eph receptors suggest multiple mechanisms in patterning of the visual system. Dev Biol 193: 21–33.
21. Memm L, Mozan P, Matthies W, Carase PW (1997) Lerk2 (ephrin-B1) is a collapsing factor for a subset of cortical growth cones and acts by a mechanism different from AL-1 (ephrin-A5). Mol Cell Neurosci 9: 314–328.
22. Boulding A, Wicht L, Schorderet DF (2013) Identification of HMX1 target genes: a predictive promoter model approach. Mol Vis 19: 1779–1794.
23. Adamkova M, Leger S, Brand M, Hadrys T, Braun T, et al. (2000) Inner ear and lateral line expression of a zebrafish Nkx5-1 gene and its downregulation in the ears of FGFr8 mutant, ace. Mech Dev 97: 161–165.
24. Adameyko I, Lallemend F, Aquino JR, Preveira JA, Topilko P, et al. (2009) Schwann cell precursors from nerve innervation are a cellular origin of melanocytes in skin. Cell 139: 366–379.
25. Furlan A, Luhke M, Adameyko I, Lallemend F, Errners P (2013) The transcription factor Hmx1 and growth factor receptor activities control sympathetic neurons diversification. EMBO J 32: 1613–1625.
26. Aboul-Saab I, Stone CE, Murray JJ, Sundaram MV (2012) The Nkx5/HMX homeodomain protein MLS-2 is required for proper tube cell shape in the C. elegans excretory system. Dev Biol 366: 296–307.