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Targeted deletion of Hand2 in cardiac neural crest-derived cells influences cardiac gene expression and outflow tract

development

Kristen L. Holler1, **Tyler J. Hendershot**1, **Sophia E. Troy**1, **Joshua W. Vincentz**2, **Anthony B. Firulli**2, and **Marthe J. Howard**1,Φ

¹ Department of Neurosciences and Program in Neurosciences and Degenerative DiseaseΨ Health Sciences Campus University of Toledo, 3000 Arlington Ave, Toledo, OH 43614-1007

² Riley Heart Research Center, Herman B Webb Center for Pediatric Research, Indiana Medical School, 1044 W. Walnut, St., Indianapolis, IN 46202-5225, USA

Abstract

The basic helix-loop-helix DNA binding protein *Hand2* has critical functions in cardiac development both in neural crest-derived and mesoderm-derived structures. Targeted deletion of *Hand2* in the neural crest has allowed us to genetically dissect Hand2-dependent defects specifically in outflow tract and cardiac cushion independent of Hand2 functions in mesoderm-derived structures. Targeted deletion of *Hand2* in the neural crest results in misalignment of the aortic arch arteries and outflow tract, contributing to development of double outlet right ventricle (DORV) and ventricular septal defects (VSD). These neural crest-derived developmental anomalies are associated with altered expression of Hand2-target genes we have identified by gene profiling. A number of Hand2 direct target genes have been identified using ChIP and ChIP-on-chip analyses. We have identified and validated a number of genes related to cell migration, proliferation/cell cycle and intracellular signaling whose expression is affected by *Hand2* deletion in the neural crest and which are associated with development of VSD and DORV. Our data suggest that *Hand2* is a multifunctional DNA binding protein affecting expression of target genes associated with a number of functional interactions in neural crest-derived cells required for proper patterning of the outflow tract, generation of the appropriate number of neural crest-derived cells for elongation of the conotruncus and cardiac cushion organization. Our genetic model has made it possible to investigate the molecular genetics of neural crest contributions to outflow tract morphogenesis and cell differentiation.

Keywords

Hand2(dHand); neural crest; bHLH; heart; outflow tract; cardiac cushion; cell type-specific gene regulation; gene microarray

ΦAuthor to Whom Correspondence Should Be Sent: Marthe J. Howard, Department of Neurosciences, University of Toledo, 3000 Arlington Ave., Toledo, OH 43614, marthe.howard@utoledo.edu.

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Introduction

Proper function of the four-chambered mammalian heart requires that oxygenated blood and deoxygenated blood be routed, respectively, to the systemic and pulmonic circulatory systems. The elegantly structured cardiac outflow tract (OFT) fulfills this task. The OFT is composed of cells derived from both neural crest and splanchnic mesoderm that collectively generate a properly septated ascending aorta and pulmonary artery, which is required for postnatal survival. Cardiac morphogenesis initiates at E7.5 in the mouse where anterior lateral plate mesoderm migrates towards the midline forming a linear heart tube. This early tube gives rise to both right and left atria, the left ventricle, and is defined as the primary heart field. During looping of the primary heart tube, a second distinct population of pharyngeal mesodermderived cells, termed the secondary heart field, contribute to both myocardial and endocardial components of the right ventricle, interventricular septum, venous pole and base of the OFT where it connects to the great vessels. In addition to the mesoderm-derived cells, the OFT acquires a substantial contribution from the cardiac neural crest. A more complete understanding of OFT formation might therefore be based in determining the function of regulatory molecules expressed in both mesoderm-derived and neural crest-derived components. The basic helix-loop-helix (bHLH) DNA biding protein Hand2 is of particular interest as it has critical functions in both mesoderm-derived and neural crest-derived components of the developing heart.

Hand2 is expressed within cells of both the primary and secondary heart fields, as well as cardiac neural crest-derived cells. Cells from the primary heart field contribute to heart formation beginning at the cardiac crescent stage (E7). Transcripts encoding *Hand2* are present in the cardiac crescent but as cardiac looping initiates (E8) *Hand2* expression within the primary heart field down-regulates to be replaced by expression within the forming right ventricle and OFT (McFadden et al., 2000; Firulli, 2003; Barnes and Firulli, 2009). Although systemic knock-out of *Hand2* does not directly affect development of primary heart field-derived structures, phenotypic anomalies are observed within the OFT and right ventricle (Srivastava et al., 1995; Srivastava et al., 1997; Thomas et al., 1998) suggesting a requirement of Hand2 for proper development of these structures. Expression of Hand2 is also crucial for the formation of several neural crest-derived structures including sympathetic chain ganglia (Howard, 2005; Hendershot et al., 2008; Schmidt et al., 2009), cranio-facial elements (Funato et al., 2009) and the enteric nervous system (Wu and Howard, 2002; Hendershot et al., 2007; Morikawa et al., 2007).

A constellation of congenital heart defects (CHDs) including ventricular septal defects, aortic arch artery patterning defects, and aortic and pulmonary valvular defects, is attributed to cardiac neural crest dysfunction; these disorders represent a substantial proportion of all observed CHDs (reviewed in Srivastava and Olson, 2000; Snider et al., 2007; and Waldo et al., 1998; Stoller and Epstein, 2005; Lie-Venema et al., 2007; Mitchell et al., 2007; Obler et al., 2008; Snarr et al., 2008). These late-stage phenotypes cannot be assessed in systemic *Hand2* knockout mice due to early developmental lethality. To better dissect the molecular mechanisms of Hand2 function specifically in neural crest-derived cells important for OFT morphogenesis and independent from its function in mesoderm-derived components, we took advantage of our floxed-*Hand2* mice (Hendershot et al., 2007; Hendershot et al., 2008). We targeted deletion of *Hand2* in the neural crest using the *Wnt1-Cre* driver line of mice (Danielian et al., 1998; Jiang et al., 2002). Our results demonstrate that targeted deletion of Hand2 in the cardiac neural crest results in embryos with defects in OFT and aortic arch artery development, which phenocopy documented neural crest-dependent defects (reviewed in Stoller and Epstein, 2005; Mitchell et al., 2007; Hutson and Kirby, 2007). Our studies confirm and extend previously published aortic arch artery defects in a similar but independently generated *Hand2* conditional allele (Morikawa and Cserjesi, 2008), by additional analysis of OFT

phenotypic anomalies and analyses of differential patterns of gene expression. We show that targeted deletion of Hand2 results in double outlet right ventricle (DORV) and accompanying ventricular septal defects (VSD). By combining targeted deletion of *Hand2* with microarray and ChIP-on-chip analyses, we have identified a number of transcriptional regulators and signaling molecules within the neural crest whose expression is modulated by loss of *Hand2*. Moreover, we show that Hand2 directly regulates a number of genes expressed within the neural crest suggesting a role for Hand2 in mediating transcriptional programs that orchestrate cell-cell interactions within the forming OFT and which affect cell cycle regulation.

Materials and Methods

Targeting Strategy and Mouse Strains

All breeding procedures, animal care and experimental protocols were approved by the Medical University of Ohio (renamed University of Toledo Health Sciences Campus) animal care and use committee prior to initiation of this work. Our strategy for targeting *Hand2* is published elsewhere (Hendershot et al., 2007). Our analysis of *Hand2fl/fl* mice demonstrates that the mice are fertile, viable and phenotypically normal; introduction of *lox*P sites in the 5′ UTR therefore does not negatively impact transcription at the targeted locus. Normal levels and patterns of *Hand2* expression have been confirmed by *in-situ* hybridization and qRT-PCR (Hendershot et al., 2007; Hendershot et al., 2008). Both systemic and targeted deletion of *Hand2* is embryonic lethal (Srivastava et al., 1997; Hendershot et al., 2008). To generate embryos older than E11 (Hendershot et al., 2008) pregnant dams are fed water containing a cocktail of catecholamines (100 μg/ml L-phenylephrine, 100 μg/ml isoproterenol, and 2 mg/ml ascorbic acid) beginning at embryonic day (E) 8. This pharmacological approach allows us to rescue *Hand2flneo/flneo* and *Hand2fl/fl;Wnt1-Cre* embryos from pre-term death, a phenotype observed in other mouse models in which norepinephrine is absent or expression curtailed (Hendershot et al., 2007; Lim et al., 2000; Thomas et al., 1995; Zhou and Palmiter, 1995). Reporter mice were generated as previously described (Hendershot et al., 2007). Homozygous R26RYFP females were mated to hemizygous Wnt1-Cre males and *Hand2fl/+;Wnt1-Cre* males were mated to *Hand2fl/fl;R262YFP* females.

Immunocytochemistry and Histology

The antibodies used for the current studies are detailed in Table 1. Embryos were prepared and treated as previously described (Hendershot et al., 2007;Hendershot et al., 2008). Briefly, embryos or tissues are emersion fixed in 4% paraformaldehyde overnight at 4°C, extensively washed in PBS and stored in 30% sucrose unless otherwise stated. For analysis of heart development, hearts were removed in ice-cold PBS, fixed in 10% neutral buffered formalin, dehydrated through a graded series of ethanol and embedded in paraffin. Paraffin sections were prepared and stained with hematoxylin and eosin (University of Texas, Southwestern Medical center, Molecular Pathology Core Laboratory) using standard procedures. Frozen sections of *Hand2wt/wt* and *Hand2flneo/flneo* E11 hearts were stained using a modified hematoxylin and eosin stain (Sanderson, 1994). Briefly, tissue sections were post-fixed in 4% paraformaldehyde (10 minutes), rinsed in PBS (2×5 minutes) and stained in hematoxylin (1 minute), followed by washing in running water, differentiation in 1% HCl in 70% EtOH, bluing in Scott's tap water (3.6 g sodium bicarbonate, 20g magnesium sulphate/litre) with counterstaining in Eosin Y for 10–20 seconds, followed by dehydration and mounting in Permount (Fisher Scientific, Fair Lawn, NJ) or EUKITT (O. Kindler, Germany). Immunostaining was done according to our established procedures (Howard et al., 1999,Wu and Howard, 2002,Liu et al., 2005,Hendershot et al., 2007). Tissue sections were blocked in a solution containing 0.1M Tris, pH. 7.5, 1.5% NaCl, 0.3% Triton-X 100 and 10% horse serum for 3×5 minutes. Primary antibody is applied in this same solution but containing 4% horse serum and incubated overnight at 4°C. Following washing in 0.1M Tris pH 7.5, 15% NaCl, secondary antibody was

applied in this same solution with 4% horse serum for three hours at room temperature. Sections are then washed 3×5 minutes in 0.1M Tris, pH 7.5, 15% NaCl and mounted in Vectashield (Vector Laboratories, Burlingame, CA) or Fluoromount-G (Southern Biotech, Birmingham, Al). To view sites of antibody binding in whole embryos (Young et al., 1999;Hendershot et al., 2007;Hendershot et al., 2008) samples were incubated overnight in PBS containing 0.3% Triton-X 100 with gentle shaking at 4° C. We used our standard blocking step and primary antibody was applied as described above and incubated for two days at 4°C with gentle shaking. Following extensive washing secondary antibody was applied and embryos incubated overnight at 4°C with shaking. Whole embryos were mounted on depression slides in Fluoromount-G (Southern Biotech, Birmingham, AL).

Confocal Microscopy

All confocal images were acquired using a Leica Microsystems multiphoton confocal microscope (TCS SP5). Confocal Z-stack images were captured using either a 10X objective $(n.a.= 0.3)$ or 63X (oil immersion) objective $(n.a.= 1.4)$ with 4X software magnification at 0.5 μm (tissue sections) or 1.0 μm (whole-mount) steps. To acquire images, fluorescent FITC– coupled, TRITC–coupled and AlexaFlour 647-coupled secondary antibodies were excited using argon, HeNE or 633 laser lines, respectively.

Gene Array Analysis

For this array (Affymetrix 430 2.0, mouse array) total cellular RNA was isolated from *Hand2wtwt* and *Hand2fl/del;Wnt1-Cre* E10.5 hearts using RNeasy (Qiagen, Valencia, CA) according to manufacturer's directions. The RNA samples were submitted to the NIH Neurosciences Microarray Consortium (Phoenix, AZ, USA;

<http://arrayconsortium.tgen.org/np2/home.do>;) for array preparation and analysis. The screen included three biological replicates for each genotype (3 chips/genotype). Expression and statistical analyses were compiled using GeneSpring (GX V 7.3.1 (Agilent Technologies, Santa Clara CA). The signal intensities from each probe set were normalized at the 50th percentile for each chip; the expression level for each gene was normalized to the median level across the arrays. The linear regression for all replicates was greater than 0.9 indicating that replicates correlated very well. A list of significantly expressed genes was generated based on a comparison of 3 arrays derived from control and three arrays derived form mutant hearts. The list was filtered first for the absent genes, secondly for a fold change cutoff of 2, and thirdly for p value of ≤ 0.05 using Welch's t-test. Genes whose expression was significantly altered by *Hand2* deletion in the neural crest were subjected to network analysis using GeneGo (St. Joseph, MI), MetaCore and amiGO to group genes based on function and/or pathway analysis. We chose genes for further analysis based on their fold change, relationship to *Hand2* and cardiac development.

Real-Time RT-PCR

Expression of transcripts encoding proteins identified in the gene array analysis were verified using quantitative reverse transcription-real time PCR as previously described (Zhou et al., 2004; Liu et al., 2005b; Hendershot et al., 2008). Briefly, we calculated the levels of transcript from the Ct values and normalized them based on expression of GAPDH. A detailed description of the protocol and equations can be found in Zhou et al., (2004) or Liu et al., (2005b). Each triplicate sample was analyzed (in triplicate) using cDNA corresponding to 10 ng of input RNA isolated from either *Hand2wt/fl* or *Hand2fl/del;Wnt1-Cre* E10.5 hearts. cDNA was mixed with Taqman Gene Expression master mix (Applied Biosystems, Foster City, CA, USA) and premixed (20X) Taqman probes (5 μM) and primers (18 μm). Forward and reverse primers (1.1 μM final concentration) with 6-FAM (carboxyfluorescein reporter dye) and a nonfluorescent quencher dye incorporated at 5′ and 3′ ends, respectively were used for all assays.

Primers and probes were purchased from Applied Biosystems (Foster City, CA, USA) from their inventory of pre-developed Taqman® Gene Expression Assays. Triplicate PCR reactions were run and analyzed using a 7500 Fast Real-Time PCR sequence detection system (Applied Biosystems); the increase in product is monitored directly and reflects the threshold number of cycles (*C*) required for a detectable change in fluorescence based on release of probe. The efficiency of each primer and probe set was determined in separate reactions and based on the slope of *C* versus input cDNA dilution. Efficiency values were ≥ 1.875 for all transcripts.

Chromatin Immunoprecipitation (ChIP) Assay

To determine whether *Cx40* is a direct *Hand2* target, we employed **Ch**romatin **I**mmunoprecipitation (ChIP) on genomic DNA cross-linked, immunoprecipitated and purified from H92c cells, using a protocol modified from manufacturer's guidelines (Millipore Corp., Billerica, MA). Briefly, \sim 1 \times 10⁶ H9c2 cells (\sim 75% confluent, 10 cm dish) were cross-linked in 1% formaldehyde for 10 minutes at **RT.** Cross linking was quenched in 1 ml of 1.25 M glycine in PBS with shaking for 5 minutes at RT. Cells were rinsed twice with ice-cold PBS, collected to 1.5 ml Eppendorf tubes, and pelleted at $2,000\times g$ for 4 minutes at 4^oC. Cell pellets were washed with 1 ml of PBS containing protease inhibitors (1 mM phenylmethylsulfonyl fluoride, 1 μg/ml aprotinin, 1 μg/ml pepstatin, Roche Inc., Madison WI) then resuspended in 200 μl of lysis buffer (1% SDS, 10 mM EDTA, 50 mM Tris-Cl, pH 8.1, 1 mM phenylmethylsulfonyl fluoride, 1 μ g/ml aprotinin, 1 μ g/ml pepstatin). The cell suspensions were incubated on ice for 10 minutes and sonicated on ice 4 times, 30 seconds each, at 1 minute intervals using a setting of 3 (Fisher Scientific Model: 550 Sonic Dismembrator). The appropriate sonication protocol was determined empirically. Lysates were centrifuged (1000 rpm, 7 minutes) and supernatants were added to 1.8 ml of dilution buffer (1.1% Triton X-100, 0.01% SDS, 1.2 mM EDTA, 167 mM NaCl, 16.7 mM Tris-HCl, pH 8.1, 1 mM phenylmethylsulfonyl fluoride, 1 μg/ml aprotinin, 1 μg/ml pepstatin), then pre-cleared with 50 μl of Protein A/G PLUSAgarose (Santa Cruz Biotechnology, Inc., Santa Cruz, CA) for 1 hour with end-over-end rotation at 4°C. Protein A/G PLUSAgarose was removed by brief centrifugation and immunoprecipitation was performed overnight at 4°C with 5 μg of Hand2 antibody.

The following day, an additional 50 μl of Protein A/G PLUSAgarose was added and the incubation continued for 2 hours at 4°C. Immuno-complexes were collected by centrifugation at 1000 rpm for 1 minute at 4°C and subjected to sequential 5 minute washes in 1 ml of each of the following buffers: low salt immune complex wash buffer (1% Triton X-100, 0.1% SDS, 2 mM EDTA, 20 mM Tris-HCl, pH 8.1, 150 mM NaCl), high salt immune complex wash buffer (1% Triton X-100, 0.1% SDS, 2 mM EDTA, 20 mM Tris-HCl, pH 8.1, 500 mM NaCl), and LiCl immune complex wash buffer (0.25 M LiCl, 1% NP-40, 1% deoxycholate, 1 mM EDTA, 10 mM Tris-HCl, pH 8.1). Precipitates were washed twice with TE buffer, then eluted twice in 250 μl of elution buffer $(1\%$ SDS, 0.1 M NaHCO₃) for 15 min at RT. The eluates were pooled and 20 μl of 5M NaCl was added; the eluates were subsequently heated at 65°C for 4 hours. Protein was removed from the samples by addition of 10 μl 0.5 M EDTA, 20 μl 1M Tris-HCl, pH 6.5 and 1 μl of 20 mg/ml Proteinase K for 1 hour at 45°C. DNA fragments were recovered by phenol/chloroform extraction and ethanol precipitation; 10 μg of yeast tRNA was added to each eluate to aid in recovery of DNA pellets. Following centrifugation (12,000 \times g, 20 minutes at 4° C), the pellets were resuspended in 50 μl of TE; 1 μl was used per PCR reaction. PCR primers were as follows: **Cx40(F)ChIP[−1144]**: 5′-ACTGTCCCTCAGTTTCCCTG-3′, **Cx40(R)ChIP[−814]**: 5′-ACAGAGGGTCAAGGACATGG-3′, **Cx40(F)ChIP[−655]**: 5′- GAACACTCTGATTGGTGGGG-3′, **Cx40(R)ChIP[−283]**: 5′-

CCAGTGGCTGTCCTTGTTTT-3′, **GAPDH**: 5′-TCCCACTCTTCCACCTTC-3′, **GAPDH**: 5′-CTGTAGCCGTATTCATTGTC-3′. PCR was conducted using GoTaq polymerase

(Promega Corp., Madison, WI) with a 58°C, 1 min annealing step over 40 cycles of amplification.

ChIP-on-chip

Neural crest-derived cells were FACsorted from 20 E10.5 hearts isolated from wild-type Wnt1 reporter embryos and samples were prepared according to the manufacturer's ChIP-chip user's guide (Roche NimbleGen, Inc. Madison, WI). Briefly, hearts were extensively washed in icecold PBS, incubated in 0.5% trypsin in CMF-PBS for 15 minutes and dissociated into single cells using a fire-polished Pasteur pipette. The cells were washed one time in PBS, pelleted, and cross linked in 1% formaldehyde for 8 minutes at RT. The cross linking was quenched in 125 mM glycine. The washed cells were filtered through a 70 μm nylon cell strainer (Falcon #352350, Bedford, MA) and subjected to FACsorting (BDFacs Aria, BD Biosciences, San Jose, CA); the non-neural crest cells from the non-sorted sample were used for the controls. The sorted neural crest cells were lysed and sonicated (Fisher Scientific Model: 550 Sonic Dismembrator) for 4 rounds for 15 seconds on a setting of 3 with a one minute incubation on ice in between each round. The appropriate settings for sonication were determined on genomic DNA and the fragment size verified in a 1% agarose gel. For IP, we used our Hand2 antibody (raised in rabbit) coupled to Dynabeads (M-20, Invitrogen, Carlsbad, CA). Complexes were allowed to form for 2 hours at 4° C with rotation and isolated on a magnetic table. Sample DNA was amplified according to manufacturer's directions (WGA3, Sigma-Aldrich, St. Louis, MO) and sent to Roach for processing $(2.1 \times 10^6$ Deluxe Promoter Array).

Transfection and Luciferase Reporter Assay

HEK293a cells were grown in DMEM plus 10% FBS supplemented with l-glutamine, Napyruvate and 100 mg/ml Pen/Stryp (Invitrogen, Carlsbad, CA); cells were transfected when 50% confluent using CaPO4 according to our published procedures (Firulli et al., 1994). In each well, 2.5 μg of pcDNA-Hand2 and/or pCS2-MT+E12 (plus empty pcDNA vector to make a total of 5 μg of plasmid DNA) was co-transfected with 5 μg of *Cx40* luciferase reporter (Cx40 +pXP2) and 150 ng of Renilla reporter (pRL-CMV) as a transfection control. Cultures were harvested 48 hours after transfection and diluted ~10 fold in lysis buffer; 50 μl of this lysate was assayed. Luciferase assays were performed as previously reported (Xu et al., 2003) using the dual luciferase assay kit (Promega Corp., Madison, WI) according to the manufacturer's protocol. Luciferase activities were read using a 96-well microtiter plate luminometer (Thermo Labsystems, Franklin, MA). Assays were done in triplicate.

Statistics

Data are presented as the mean \pm S.E.M unless otherwise stated. Statistical significance was determined using Student's unpaired t test or ANOVA and Bonferroni post hoc test, unless otherwise stated. At least four embryos were examined for each condition from at least three different matings unless otherwise stated. For microarray analysis, three biological replicates were run and each in triplicate.

Results

Hand2 Misexpression Affects Heart Development

In order to define the molecular basis for defects in heart development associated with loss of *Hand2* function, we generated a conditional targeted mutation of *Hand2* (Hendershot et al., 2007; Hendershot et al., 2008). For the current studies, we targeted deletion of *Hand2* specifically within neural crest-derived cells by intercrossing our compound heterozygous floxed-*Hand2* (*Hand2f/ldel*) mice with mice expressing Cre recombinase under control of the *Wnt1* promoter. The *Wnt1*-*Cre* transgene allows floxed genes to be targeted in the neural crest

(Danielian et al., 1998; Jiang et al., 2000; Brewer et al., 2004; Hendershot et al., 2008). Because loss of *Hand2* expression within neural crest-derived cells is embryonic lethal between E10-E11 due to insufficient levels of norepinephrine (Lim et al., 2000; Hendershot et al., 2008), we routinely feed pregnant dams a cocktail of catecholamine intermediates beginning at E8 (Lim et al 2000; Thomas et al., 1995; Zhou and Palmiter, 1995; Kaufman et al., 2003; Hendershot et al., 2008). This pharmacological rescue makes it possible to bring embryos to term, allowing assessment of *Hand2* loss-of-function within neural crest-derived cells and tissues, throughout late stage OFT morphogenesis and up to birth. *Wnt1*-mediated deletion of *Hand2* results in developmental defects in all neural crest-derived structures dependent upon *Hand2* for early aspects of their development (Hendershot et al., 2007; Hendershot et al., 2008). Briefly, there is significant loss of sympathetic and enteric neurons, alterations in neurotransmitter expression and regulation, as well as a number of defects in cranio-facial development, consequent to the loss of *Hand2*.

We previously reported that embryos harboring the neomycin cassette (*Hand2flneo/flneo*) in their genome misexpress *Hand2* resulting in a *Hand2* hypomorph (Hendershot et al., 2008). Interestingly, the impact of *Hand2* under-expression was observed to be tissue dependent demonstrating both gene dosage and penetrance effects. The importance of *Hand* gene dosage for cardiac development is well documented (McFadden et al., 2005) and this current study confirms and extends these previous findings. At E18 we observe differential effects on heart development in *Hand2flneo/flneo* embryos compared to *Hand2fl/fl* embryos (Fig. 1). In *Hand2* hypomorphic (*Hand2flneo/flneo*) embryos, under-expression of *Hand2* could be affecting neural crest-derived structures, primary heart field myocardium, and/or intraventricular septal myocardium derived from the second heart field. In the cardiovascular system, the neural crest makes contributions to the aortic arch arteries and OFT. In *Hand2flneo/flneo* embryos, there were no obvious developmental defects within the aortic arch arteries (one embryo had patent ductus arteriosus); ~50% of hearts examined showed membranous VSDs (not shown). Development of the right ventricle is dependent upon *Hand2* but independent of the neural crest (Srivastava et al., 1997; Thomas et al., 1998; Srivastava, 1999). Development of the right ventricle was affected in all hypomorphic hearts examined (Fig. 1B). In these hearts ventricular hypoplasia and disorganized trabeculations indicates likely effects on proliferation and/or apoptosis of cardiomyocytes. We have previously reported that *Hand2* regulates proliferation of neural crest-derived noradrenergic neural precursor cells and neuroblasts (Hendershot et al., 2008) suggesting a potential function for *Hand2* in the heart (see data below). The hypomorph phenotype thus recapitulates some of the defects observed in *Hand2* systemic null embryos (Thomas et al., 1998; McFadden et al., 2005). Since we were interested in dissecting the function of *Hand2* in cardiac neural crest-derived structures and given this potential complication, we employed SYCP-*Cre* mice to remove the neomycin-resistance cassette within our *Hand2* conditional allele and completely rescue these hypomorphic defects (Hendershot et al., 2008).

Targeted Deletion of Hand2 Results in Outflow Tract Defects

Based on the expression pattern of *Hand2* within the cardiac neural crest, we predicted that deletion of *Hand2* specifically within this cell population would adversely effect proper development of outflow tract structures. Analysis of *Hand2* loss-of-function within the neural crest consistently demonstrated effects on ventricular septation and alignment of the great vessels (Fig. 2). We consistently observed double outlet right ventricle (DORV) with an associated membranous VSD in all *Hand2fl/fl;Wnt1-Cre* embryos (Fig. 2B & C) examined. This neural crest-derived anomaly (DORV) was never observed in *Hand2* hypomorphic embryos. Analysis of *Hand2wt/wt;Wnt1-Cre;R26R/YFP* mice (Fig. 2D) and *Hand2fl/fl;Wnt1-Cre/R26R/YFP* mutant reporter mice (Fig. 2E) demonstrates that *Hand2* will be recombined in the neural crestderived cells which contribute to the conotruncus and aortic arch arteries. The effect of

misexpression of *Hand2* within the heart therefore depends heavily upon *Hand2* expression levels as well as the identity of the cells where expression of Hand2 is affected.

Defects in Neural Crest Migration and Proliferation

Between days E10 and E13 in the mouse, the neural crest not only remodels the pharyngeal arches, contributing to the smooth muscle of the forming arch arteries, but also invades the OFT to form the septum that divides the aorta and pulmonary trunk (Conway et al., 1997; Conway et al., 2000; Snarr et al., 2008). We considered that aberrant migration of neural crestderived cells into the OFT might contribute to the VSDs observed in all embryos deficient in expression of *Hand2* in neural crest-derived cells. At E10.5 neural crest-derived cells have migrated into the OFT traversing approximately 50% of this developing structure (Fig. 3). The patterns of migration observed in *Hand2wt/wt;R26R/Wnt1-Cre* and *Hand2fl/del;R26R/Wnt1-Cre* E10.5 reporter embryos is strikingly affected by deletion of *Hand2*. At this stage of development it is possible to observe the "prongs" of neural crest-derived cells that will migrate into the truncal cushions (arrows in Fig. 3A, C) and which contribute to septation of the aorta and pulmonary arteries. Visual inspection suggests that loss of *Hand2* negatively impacts cell migration and/ or cell numbers. There is a dramatic reduction in the number of neural crest-derived cells within the OFT (compare Fig. 3B and D). The migration of neural crest-derived cells into the aortic sac and truncus is not only required for septation but elongation of the conotruncus as well (Waldo, 1999; Epstein et al., 2000; Snider et al., 2007). Examination of neural crest-derived cells (Fig. 3B and 3D) actively migrating into this region suggested that cell-cell interactions are important. It appears that neural crest-derived cells migrate as a coherent sheet of cells with the majority of cells being in contact with their neighbors. In the

Hand2fl/del;Wnt1-Cre;R262RYFP hearts we observe not only fewer cells that have reached the truncus but in addition they appear to migrate more as individual cells with fewer contacts with neighboring cells.

Based on our previous findings indicating that *Hand2* is required to generate the appropriate number of neuronal precursor cells and neuroblasts in developing noradrenergic sympathetic ganglia (Hendershot et al., 2008) we counted the number of cells expressing the proliferation marker Ki-67 at E10.5 in *Hand2wt/wt;Wnt1-Cre;R26RYFP* and *Hand2fl/del;Wnt1-CreR262RYFP* hearts. The number of GFP (YFP) marked neural crest cells that labeled for immunoreactivity to Ki-67 was determined on 20 μm frozen cross sections at the level of the OFT (Fig. 4). Cells were counted in the entire outflow tract of control and mutant hearts. In control embryos, the mean number of proliferating neural crest-derived cells per tissue section in the OFT was 47 ± 6 compared to 10 ± 2 (P < 0.01) in the mutant OFT. Based on the number of neural crest-derived cells (GFP) migrating into and within the OFT, we observed a significant reduction in the proportion of proliferating cells based on the total neural crest cell population (Fig. 4G); in control embryos 41% + 4 of neural crest-derived cells were proliferating compared to 24% \pm 3 (P < 0.05) in the mutant outflow tract. This result suggested possible impact of loss of *Hand2* on mitosis. To determine if the number of cells in mitosis was affected by loss of Hand2, we counted the number of mitotic figures in the OFT of *Hand2wt/wt;Wnt1-Cre;R26RYFP* and *Hand2fl/del;Wnt1-CreR262RYFP* E10.5 embryos labeled with Ki-67 (Fig. 4E, F, H). The percentage of mitotic figures in *Hand2fl/del;Wnt1-CreR262RYFP* OFT was not significantly different from control indicating that equivalent numbers of neural crest-derived cells initiated mitosis; this likely results from problems in either G1 or S phases of the cell cycle. This suggests that the effects we observe on proliferation likely reflect alterations in the regulatory mechanisms controlling transit through the various stages of mitosis.

Outflow Tract Defects Associated with Altered Gene Expression

In order to directly address the genetic basis of the OFT phenotype we observe in *Hand2fl/del;Wnt1-*Cre embryos, we compared the panel of transcripts whose expression was

altered in mutant compared to wild-type embryos. We screened gene arrays using RNA isolated from E10.5 *Hand2wt/wt* and *Hand2fl/del;Wnt1-Cre* whole hearts. The screen included three biological replicates for each genotype (9 chips/genotype). Our results demonstrate that targeted deletion of *Hand2* in neural crest-derived cells in the heart significantly ($P < 0.05$) affected expression of 309 identified transcripts and 56 unidentified transcripts regulated minimally by greater than or equal to a 2-fold change (Fig. S1). The identified transcripts were associated with 11 functional classes (Fig. 5A; GeneGo Inc., St. Joseph, MI). Of the identified regulated genes, 68 were up-regulated (Fig. 5B) and 162 were down-regulated (Fig. 5C). Of the identified genes (309) 79 proteins were identified but which have no known function. Of the genes regulated, a notable fraction are known to affect heart development and/or cardiac function (Table 2). We validated a set of genes whose apparent function is associated with some aspect of the morphological or functional consequences of *Hand2* deletion; a subset of these genes were validated using RNA purified from cardiac neural crest-derived cells FACsorted from E10.5 heart showing that our transcript analysis yielded alterations in expression of neural crest-specific genes (* in Table 2.).

Genes associated with neural crest cell migration

In our initial analysis of the consequences on heart development due to targeted deletion of *Hand2* in the neural crest, we observed that population of the conotruncus by neural crestderived cells was aberrant (Fig. 3). Several genes associated with migration were differentially expressed in the *Hand2fl/del;Wnt-Cre* mice. These regulated genes fall into at least three functional networks (Table 2). We validated genes from each group using qRT-PCR and/or immunostaining. We observed a decrease in transcript encoding Pdgf (↓3.89); Pdgf has been associated with neural crest cell migration (Orr-Urtreger and Lonai, 1992). Interestingly, defects in the Pdgf receptor results in developmental defects in the OFT as well as other neural crest-derived structures (Morrison-Graham et al., 1992). We validated *integrin α9* (*Itga9*; ↑2.94) because it has an important function in angiogenesis (Brooks et al., 1994), which is adversely affected in *Hand2* mutant embryos (and see Morikawa et al., 2008) as well as a potential role in cell migration (Young et al., 2001;Huang et al., 2005). The *integrin subunit* α *4* is a component of the α 4b1 fibronectin-binding domain, which is one important interaction regulating cell migration (Sheppard et al., 1994;Pinco et al., 2001), and was up-regulated (↑3.83) in response to deletion of *Hand2*. Interestingly, *Adam19* (↓2.16) inhibits migration mediated by α4b1 integrins (Huang et al., 2005). Deletion of *α4 integrin* is embryonic lethal in part due to defective development of the epicardium and coronary vessels (Yang et al., 1995).

The regulation of connexin 40 (Cx40; Gja5) was of particular interest because it is heartspecific (Dahl et al., 1995; Kirchhoff et al., 2000) and associated with both adhesion and cellcell communication. Based on the array results, transcripts encoding Cx40 were downregulated 24-fold in *Hand2fl/del;Wnt1-Cre* hearts compared to control; this transcript was not detectable by qRT-PCR in the linear range of amplification (transcript became detectable at 40 cycles of amplification). We used immunostaining and confocal microscopy to visualize sites of Cx40 expression on tissue sections of E10.5 OFT (Fig. 6). Interestingly, although we could not detect transcripts encoding Cx40 there remains a small number of cells (Fig. 6B, D, F, arrows) that express Cx40 protein in the mutant OFT. The majority of neural crest-derived cells express Cx40 in the wild-type OFT, a novel finding in itself (Fig. 6A, C, E). Connexin 40 is also expressed in some cells in the second heart field-derived component of the conotruncus in both the control and mutant hearts (white arrows Fig. 6A, B) validating that neural crest specific deletion of *Hand2* is revealing a cell autonomous effect. We further assessed regulation of Cx40 because it has not previously been reported in the neural crest and is affected cell autonomously by *Hand2* loss-of-function.

Hand2 binds to and *trans***-activates the Cx40 promoter**

We sought to validate that a subset of the genes identified in our microarray screen represent direct Hand2 transcriptional targets (Table 2 (+); Fig. 7). Based on ChIP (*Cx40*) or ChIP-onchip we identified a number of putative Hand2-target genes. We analyzed *Cx40* in some detail because *Cx40* null mice display membranous VSDs phenotypically similar to those occurring in our *Hand2*-loss-of-function embryos (Kirchhoff, 2000) and we found that *Cx40* is expressed in cardiac neural crest cells (Fig. 6). A putative *cis*-regulatory region for the *Cx40* gene has been previously identified (Seul et al., 1997) and spans bases from −1190 to +121. Initially, we analyzed this region for E-box (bHLH consensus) binding sites (CANNTG) and identified a total of 9 sites 5′ to the *Cx40* transcription start site 5′ to exon 1 (Fig. 7A). For ChIP analysis we used H9c2 cells, a rat cardiomyocyte cell line which endogenously expresses *Hand2* (Zang et al., 2004) and which upregulates Cx40 expression in response to *Hand2* overexpression (Togi et al., 2006). PCR amplification of DNA fragments immunoprecipitated using our Hand2 antibody revealed that DNA fragments overlapping the proximal (−655 to −283 bp), but not distal (−1144 to −814 bp) E-box-containing DNA sequences co-immunoprecipitated with Hand2 (Fig. 7B); the two sets of primers we designed and tested flank all 9 of the E-boxes in the *Cx40* 5′ putative *cis*-regulatory region. The PCR product band was TOPO®cloned (InVitrogen, Carslbad, CA) and the sequence verified as the appropriate region of *Cx40* (not shown). These data suggest that Hand2 directly binds to the putative *Cx40* promoter at E-box sites located between −655 and −283 bp relative to the transcription start site. In order to directly demonstrate that this binding could have functional relevance, we conducted *trans*-activation studies to confirm that Hand2 can regulate *Cx40* transcriptional activity. We used a reporter construct (generously provided by Benoit Bruneau, Gladstone Institute of Cardiovascular Disease) containing sequence corresponding to bases −1068 to +121 of the *Cx40* gene, that drives expression of a *luciferase* cassette (Bruneau et al., 2001); this was co-transfected with *Hand2* and/or *E12* expression constructs into HEK 293 cells (Fig. 7C). While expression of neither Hand2 nor E12 alone was sufficient to modulate *Cx40*-luc reporter expression, expression of both factors together up-regulated *Cx40*-luciferase activity (~2.2 fold) over baseline ($p < 0.001$). These results indicate that Hand2 and its bHLH dimer partner, E12, can positively regulate gene expression from the *Cx40* promoter. Together, these results indicate that *Cx40* is a direct target of Hand2 within the cardiac neural crest.

We validated (qRT-PCR) an additional subset of genes, identified in the array screen, that appear in one or more of the functional pathways associated with cell-cell interaction and/or adhesion (Table 2; validated genes are shown in blue). In addition to qRT-PCR and ChIP, we also performed a ChIP-on-chip screen on FACsorted cardiac neural crest-derived cells (Table 2; direct Hand2 target genes indicated by a +) to determine which of the validated genes are potentially neural crest-specific Hand2 direct transcriptional targets.

Genes associated with neural crest cell cycle

We validated a subset of genes associated with cell cycle because of the significant reduction in proliferation that we observe (Fig. 4) when *Hand2* is deleted in neural crest-derived cells (Hendershot et al., 2008). The genes that we validated, based on qRT-PCR and ChIP-on-chip, include *cdk6* (↑2.53), *Insm1* (↓20.7 array Vs undetectable qRT-PCR); each of these genes is associated with some aspect of cell proliferation and/or known *Hand2*-related genes. Several classical "cell cycle" genes are differentially regulated in the absence of *Hand2*. The most highly regulated gene in the array was *cyclin B1 interacting protein 1* (*Ccnb1ip1*; ↑37.42). The protein coded for by this gene is a Ring finger protein. It is a member of a group of B-type cyclin E3 ubiquitin ligases that also has cyclin-dependent kinase and cyclin-dependent phosphorylation sites (Toby et al., 2003). This protein interacts with cyclin B1, an important checkpoint regulator of cell cycle progression, and causes its degradation (Toby et al., 2003). Overexpression of *Ccnb1ip1* substantially reduces expression of cyclin B1 resulting in

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decreased proliferation (Toby et al, 2003). This protein also has been shown to affect migration and metastasis (Singh et al., 2007) potentially linking cell cycle with migration; functions which are both effected by deletion of *Hand2*. It is interesting to note that none of the cyclin proteins were identified in our array but several cyclin-dependent kinases or interacting proteins were. *Cyclin dependent kinase 6* (↑2.53) was increased. Cyclin dependent kinase 6 is a serine/threonine protein kinase that is activated by cyclin D1, D2 and D3, and which is required for the G0 to G1 transition (Malumbres and Barbacid, 2005). Additional genes coding proteins that have a role in cell cycle regulation were also identified. *Insm1* (*Insulinoma associated 1*) was of particular interest because it has been identified as a member of a transcription factor network that regulates development of neural crest-derived noradrenergic sympathetic ganglion neurons (Howard, 2005;Hendershot et al., 2008;Wildner et al., 2008;Parlier et al., 2008). *Hand2* has a central role in this pathway and likely functions upstream or in parallel with Insm1 (Hendershot et al., 2008;Lan and Breslin, 2009). Targeted deletion of *Hand2* or *Insm1* results in a marked decrease in proliferation of the pool of neural precursor cells (Hendershot et al., 2008;Wildner et al., 2008). A potential site of Insm1 activity in the heart is at cyclin D1. Insm1 binds to cyclin D1 (Liu et al., 2006) which competes for the interaction of cyclin D1 with Cdk4. The Cdk4/cyclin D1 interaction is required for proper phosphorylation of retinoblastoma protein Prb necessary for cell cycle progression from G0 to G1 (Zhang et al., 2009). The putative Insm1 interacting protein, CAP1 (adenylate cyclase associated protein; Xie et al., 2002) transcript level was also increased (↑7.21) in the mutant hearts. Interestingly, transcripts encoding peripherin, a neuron specific gene and associated with *Hand2* and *Insm1* was down-regulated 6.41 fold in the mutant hearts. Expression of several histones was also affected by deletion of *Hand2* including: 1) *histone cluster 1* (↓2.1; Hist1h2bb); 2) *H2a histone family member V* (\downarrow 2.27); 3) *H3 histone family 3B* (\downarrow 2.55); and 4) *histone cluster 2 H2be* (\downarrow 2.67). Although the nucleosome (core histone complex) is associated with multiple aspects of DNA replication, repair and chromatin structure, one of these histones, H2be, is required for Cdk6 to bind its substrate, retinoblastoma protein Prb, required for initiation of cell cycle progression from G to S phase (Zhao et al., 2000;Marzluff et al., 2002;Jaeger et al., 2005;Ewen, 2009). Although we found a significant reduction in cell proliferation (Fig. 4G), there is no significant reduction in the percentage of cells that initiate mitosis (Fig. 4H) suggesting a potential role for Hand2 in regulation of cell cycle progression.

Genes associated with VSD and DORV

A number of genes have been identified as risk loci for VSDs and DORV. We were very interested in these genes since targeted deletion of *Hand2* in the neural crest results in these defects in 100% of embryos examined (Fig. 2). *Sox11* (↓2.5), a member of the SRY-related HMG-box family of transcriptional regulatory genes is an important regulatory gene affecting neural, skeletal and cardiac development. *Sox11* null embryos develop VSDs with 100% penetrance (Sock et al., 2004). These embryos also display DORV and other OFT defects similar to that which we observe in the *Hand2* mutant embryos. *Gli3* (↓2.54; C2H2-type zinc finger protein) was identified as a VSD susceptibility gene in a study focused on 12q13 chromosomal region previously identified as associated with occurrence of VSD in humans (Qui et al., 2006). Although the molecular mechanism of Gli function in the cardiac neural crest is not yet known, Gli-family members mediate sonic hedgehog (Shh) signaling (te Welscher et al., 2002a;te Welscher et al., 2002b;Stamataki et al., 2005) and Zic3 translocation to the nucleus appears dependent upon Gli3 (Bedard et al., 2007). Disruption of *Zic3* expression results in a number of developmental defects related to left-right asymmetry including conotruncal malformations (Mégarbané et al., 2000;Ware et al., 2006). Mutations in *Foxc1* (↓3.73; forkhead-winged helix transcription factor family) are associated with abnormal vasculogenesis, coarctation of the aortic arch, and VSDs, related to Notch signaling (Kume et al., 2001;Seo et al., 2006); the only linkage to Notch signaling in our array is Hey1 (\downarrow 2.0). Notch signaling has been implicated as important for septation and Hey 1 (hairy and enhancer

of split-related family of bHLH-type transcriptional repressor) is a Notch-downstream target gene (Sakata et al., 2002). In conjunction with Hey 2 (not regulated) Hey 1 has been implicated in Notch-mediated epithelial-mesenchymal transition (EMT) (Fischer et al., 2004;Niessen and Karsan, 2008). This occurs via a decrease in expression of several metalloproteinases (MMP; Brou et al., 2000). Two MMPs, MMP14 (\downarrow 2.02) and Adam 19 (\downarrow 2.16) were down-regulated in response to deletion of *Hand2* in the cardiac neural crest suggesting that the reduction in migration and invasion of the cardiac cushions could be related to decreased expression of these genes.

Valvulogenesis

The cardiac cushions develop in the distal portion of the common OFT and between the common atria and ventricle (reviewed in Barnett and Desgrosellier, 2003; Goodwin et al., 2005). Cushion formation requires production of extracellular matrix material that becomes situated between the endocardial and myocardial layers of the developing heart. Neural crestderived cells contribute to the truncal cushions that eventually form the aorticopulmonary septum and semilunar valves (de Lange et al., 2004). A number of genes associated with aspects of cardiac cushion formation, valvulogenesis and/or septation have been identified and which are affected by targeted deletion of *Hand2* in the neural crest. In the absence of *Nf-ATc* (nuclear factor of activated T-cells) heart valves fail to form and embryos die (de la Pompa et al., 1998, Zhou et al., 2002). Expression of *NF-ATc1* is induced by *NF-ATc2* (↓2.29) that is downregulated in response to *Hand2*-deletion in the neural crest. *Collagen type XI a1* (*Col11a1*, ↓2.1 array Vs. ↓1.43 PCR) is expressed in the non-cartilaginous tissue of the heart. Mutations in the *Col11a1* gene have been implicated in valve defects. *Adam 19* (↓2.16 array Vs. ↓2.63 qRT-PCR) is a metalloprotease that is expressed in the endocardial cushion. *Adam 19* null mice die within a week of birth due to thickened and misshapen semilunar valves, tricuspid valves, and membranous VSD (Horiuchi et al., 2005; Komatsu et al., 2007). EMT is required for proper cardiac cushion transformation (Barnett and Desgrosellier, 2003) and is stimulated by members of the transforming growth factor-β family of proteins. Deletion of latent transforming growth factor beta binding protein 1 (Ltbp1, \downarrow 2.13) results in multiple heart defects including improper septation of the OFT, patent truncus arteriosus (PTA), and interrupted aortic arch (IAA) (Todorovic et al., 2007).

Discussion

The consequences to loss of *Hand2* in the heart have been a focus of much investigation (Srivastava et al., 1997; Yamagishi et al., 2001; McFadden et al., 2005; Morikawa and Cserjesi, 2008; Reviewed in Firulli and Thattaliyath, 2002; Firulli, 2003; Bhattacharya et al., 2006; Olson, 2006; Srivastava, 2006). *Hand2* contributes to normal development of both the OFT and right ventricular myocardium. Cells from the secondary heart field and cardiac neural crest contribute to the OFT by migrating into the formed heart tube, whereas only secondary heart field cells contribute to right ventricle myocardium development (reviewed in Kelly and Buckingham, 2002; Garry and Olson, 2006; Hutson and Kirby, 2007). Since *Hand2* is expressed within the second heart field myocardium and cardiac neural crest it has been difficult to separate effects of *Hand2* deletion in neural crest-derived or mesoderm-derived components of the heart because of the close interaction of cells from these two lineages. This problem has been addressed in the current studies. We employed an intercross of our mice harboring a *Hand2* conditional allele with the *Wnt1-Cre* transgenic mouse line to interrogate the consequences to loss of *Hand2* specifically in the neural crest. Our analysis of OFT development combined with gene profiling data demonstrate that *Hand2-*target genes fall into a variety of functional categories including cell adhesion, cell cycle regulation, proliferation control and intracellular signaling. We have identified a large number of genes as likely or

identified loci for neural crest-dependent cardiac developmental anomalies, including VSD, DORV, and valve malformation.

Hand2 Gene Dosage and Cardiac Development

Conditional inactivation of *Hand2* in secondary heart field cells presages malformation of the right ventricle manifesting as ventricular hypoplasia and in the conotruncus resulting in membranous ventricular septal defects (VSDs. Specific deletion of *Hand2* in neural crestderived cells that contribute to the conotruncus results in developmental abnormalities expected based on expression pattern. Interestingly, occurrence of OFT anomalies and VSDs was not predicted to be prevalent in *Hand2fl/fl;Wnt1-Cre* embryos because of potential redundant function with *Hand1*, which is co-expressed with *Hand2* in a subset of cardiac neural crest cells (McFadden et al., 2005). Deletion of *Hand1* in neural crest-derived cells does not affect phenotypically, embryonic development. However, in these same mice, *Hand2* haploinsufficiency, due to deletion in the ventrolateral branchial arch, shows novel defects in craniofacial neural crest-derived structures (Yanagisawa et al., 2003; Barbosa et al., 2007). We observe clear defects in the cardiac neural crest of *Hand2fl/fl;Wnt1-Cre* embryos, where *Hand1* is coexpressed, suggesting that Hand2 function would be dominant to that of Hand1; these closely related transcriptional regulators do not appear to have redundant functions related to migration or differentiation of cardiac neural crest. Our data suggest that *Hand1* and *Hand2* have unique functions in the cardiac neural crest.

The defect we observe in right ventricle development in the *Hand2* hypomorph embryos, disorganized trabeculations and thin ventricular wall, was predictable given that *Hand1* expression is low (albeit undetectable) within the right ventricle. The phenotypic characteristics we describe in both *Hand2fl/fl;Wnt1-Cre* and *Hand2flneo/flneo* embryos supports previous data suggesting that Hand2 protein function is dosage dependent (McFadden et al., 2005; Firulli et al., 2005; Hendershot et al., 2008). The phenotype observed in the *Hand2* hypomorph embryos compared to loss of *Hand2* in neural crest-derived cardiac structures underscores the complex functional interactions that the *Twist* family of bHLH proteins exhibits *in-vivo* (Firulli et al, 2003; Firulli et al., 2005; Barnes and Firulli, 2009); our data indicate a complex pattern of interaction between different cell types that express *Hand2*. The nature of the defects we observe in both the *Hand2* hypomorph embryos and *Hand2* neural crest-specific knockout embryos attests to the utility of our genetic model. Because of the acknowledged importance of gene dosage in *Hand* factor function, the *Hand2* hypomorph allele will provide a valuable tool for future examination of phenotypic response to misexpression of *Hand2* as well as genetic and functional interaction of Hand2 with other transcriptional regulators.

Outflow Tract Defects

In the mouse, formation of the septum and valves initiates between E9.5 and E10 (Webb et al., 1998; Jiang et al., 2000). Cells in the OFT undergo an EMT and migrate into the cardiac jelly (Song et al., 2000; Camenisch et al., 2002; Gitler et al., 2003; Person et al., 2005); these cells derive from the neural crest and the endocardium (Runyan and Markwald, 1983; Camenisch et al., 2000; Kisanuki et al., 2001). In conjunction with cells located in the AV canal (endocardial origin), neural crest-derived cells will form the membranous septum and valves (Runyan and Markwald, 1983, Poelmann et al., 1998; De Lange et al., 2004). Formation of the aorticopulmonary septation complex involves neural crest-derived cells and non-neural crestderived cells in the conotruncal ridges (Waldo et al., 1998). Neural crest-derived cells form the prongs of the aorticopulmonary septum at around E9.5 (Waldo et al., 1998; Waldo et al., 1999; Kisanuki et al., 2001); these prongs of cells eventually form parts of both the aorta and pulmonary structures, valves and arterial smooth muscle (Waldo et al., 1998). Our results indicate that formation of the AP prongs is affected by loss of *Hand2* in the neural crest-derived cells that migrate into the OFT. The reduced cell density and apparent decrease of cell-cell

contact between the migrating cells is reflected in altered expression of a number of genes associated with these functions.

Formation of the OFT septum begins at around E10.5 (Jiang et al., 2002). The defects that we observe in septation and OFT vessel formation likely result from a substantial reduction in the number of neural crest-derived cells that form the AP prongs. At E9.5 neural crest-derived cells are located proximal to the aortic sac and initiating migration through it to the truncus arteriosus (Jiang et al., 2002). Later in development, the OFT will form the aortic and pulmonary OFT vessels as the aorticopulmonary septum forms at the distal pole and the conotruncal septum forms at the proximal end. The outflow valves form concurrently. Our results demonstrate that *Hand2* is required for these morphogenetic events to proceed properly. As described previously (Morikawa and Cserjesi, 2008), the absence of *Hand2* in cardiac neural crest causes misalignment of the aortic arch arteries and OFT. Our data suggest that cell-cell communication, mediated by gap-junctions, constitutes a critical component of signaling in the cardiac neural crest; anomalies in vessel alignment and septation appear to occur, in part, as a result of decreased expression of Cx40 in neural crest-derived cells. The reduction in Cx40 expression appears to be a direct result of Hand2 loss-of-function since our data indicate that *Cx40* is a direct target of Hand2.

Cell-cell contact and intercellular signaling via gap junctions has been associated with cell cycle control (Lowenstein, 1988) and neural crest migration (Huang et al., 1998; Lo et al., 1997). Both gain-of-function of *connexin 43* in the neural crest and systemic loss-of-function of *connexin 43* results in defective development of the conotruncus as well as the pulmonary outflow vessel (Reaume et al., 1995; Ewart et al., 1997; Huang et al., 1998; Sullivan et al., 1998; Simon et al., 2004). It is clear from these previous studies that expression levels of connexin proteins in the neural crest is critical and contributes to development of the OFT; Cx40 has not previously been reported in the neural crest. Importantly, it appears that Hand2 directly affects expression of Cx40 in the developing OFT. It will be interesting in future studies to validate other Hand2 targets or interacting molecules identified in our mRNA screen and validated as potential direct Hand2 targets in out ChIP-on-chip screen that are involved in cellcell contact and migration in OFT formation. Based on our gene profiling, expression of connexin 43 was not altered by deletion of *Hand2* in the neural crest. We provide evidence that in the absence of *Hand2*, neural crest-derived cells migrating within the OFT do not migrate as a coherent sheet of cells as they normally do (Bancroft and Bellairs, 1976; Davis and Trinkaus, 1981; Raible and Eisen, 1996; Gammill et al., 2006; Kuriyama and Mayor; 2008). It is interesting to note however, that the extent of migration is not adversely affected by *Hand2* deletion. Although we did not detect any notable difference in the length of the OFT in *Hand2* mutant embryos, the apparent reduction in the numbers of neural crest-derived cells coupled with the decreased adhesion that we observe in the AP prongs suggests that it is aberrant remodeling affecting proper generation of the aorticopulmonary septation complex that contributes to generation of DORV and VSD.

Interestingly, in about 20% of the hearts that we examined, we found neural crest-derived cells that had tracked incorrectly and migrated into the ventricles. Although we did not observe any apparent effect of these mis-located neural crest-derived cells on ventricle development, abnormal ventricle development has been reported previously using an independently derived line of floxed *Hand2* mice (Morikawa and Cserjesi, 2008). These authors concluded that lossof-*Hand2* function in cardiac neural crest induced proliferation of cardiomyocytes resulting in enlarged right ventricles (Morikawa and Cserjesi, 2008). This was an intriguing finding because neural crest-derived cells do not normally contribute to myocardial development although expression of *Hand2* in cardiomyocytes is necessary for right ventricle development (Srivastava, 1997). The hypertrabeculations observed in these mutant hearts is reminiscent of what we observe in our *Hand2* hypomorphic embryos. It is likely that this reported aberrant

right ventricle development was in fact a primary hypomorphic phenotype within the cardiomyocytes, as this group observed hypomorphic phenotypes from their floxed *Hand2* mice (Morikawa et al., 2007); in these embryos, although targeted *Hand2* deletion in the neural crest would give rise to the expected defects in OFT development, misexpression of *Hand2* in the right venrtricle would be expected to affect trabeculations and cell cycle.

Neural Crest Cell Cycle Regulation

In the peripheral nervous system *Hand2* is required for neurogenesis and neurotransmitter specification and expression in noradrenergic sympathetic ganglion neurons (Howard et al., 1999; Howard et al., 2000; Wu and Howard, 2002; Xu et al., 2003; Hendershot et al., 2007; Hendershot et al., 2008; Morikawa et al., 2008; Schmidt et al., 2009). One important function of *Hand2* is in cell cycle regulation; in the absence of *Hand2* neural precursor cells cease to divide (Hendershot et al., 2008). We identified a number of cell cycle and proliferation genes regulated downstream of *Hand2* in the heart. The number of cells that populate the OFT is reduced following targeted deletion of *Hand2* in the neural crest; this could occur as a result of aberrant migration, decreased proliferation and/or abnormal cell cycle regulation; gene profiling suggests that all of these events are affected in the mutant hearts. The appropriate number of cells in the AP prongs of neural crest-derived cells that move into the truncal cushions is required for elongation of the conotruncus and septation into the aortic and pulmonary trunks. The abnormal pattern of migration, decreased cell numbers and loss of cellcell signaling likely contribute to the formation of VSDs observed in all mutant embryos. Normal closure of the membranous septum involves three coordinated morphogenetic events: the muscular septum must grow upwards, the endocardial cushions must form the inlet septum (AV canal) and the formation of outlet septum by growth of the conotruncal ridges must occur (Webb et al., 1998; Lamers and Moorman, 2002; Raid et al., 2009). As part of this fusion, the valves are also formed. We observed in 100% of mutant embryos DORV that may have developed as a consequence of the development of VSD (Yelbuz et al., 2002; Xu et al., 2004) or due to the misalignment of the great vessels (described in Morikawa and Cserjesi, 2008) observed in embryos with neural crest targeted deletions of *Hand2*. Abnormal development of the cardiac cushions could contribute to the generation of these defects. Misregulation of transit through the cell cycle, resulting in a decreased number of neural crestderived cells populating the AP prongs likely affects truncal cushion development and septation. Based on the profile of Hand2-trancriptioanl targets and the array of proliferation/ cell cycle genes modulated by loss of Hand2, we suggest that the cell cycle is arrested/delayed in G1 or S phase, a situation that would account for a reduction in cells while maintaining a stable proportion of cells in M phase.

Cardiac Cushion and Valve Development

Proper patterning and differentiation of the cardiac cushions is required for formation of the semilunar valves and aorticopulomnary septation. In order for the cardiac cushions to develop extracellular matrix must be deposited into the acellular cardiac jelly. In order for valves to form, EMT must occur to remodel the cushions (Wang et al., 2004). Genes associated with transcriptional control (*Nf-ATc; Foxc1, Foxp1*), extracellular matrix composition (*collagen XI a1; Ltbp1*) and protein modifying enzymes (metalloproteases), all associated with aspects of cardiac cushion development, were affected by deletion of *Hand2*; a number of these genes are likely direct Hand2 targets. The cardiac phenotype of mouse embryos deficient in *Nf-ATc*, *FoxP1*, and *Sox11* show nearly identical defects resulting from problems in cardiac cushion development (De la Pompa, 1998, Ranger et al., 1998; Kume et al., 2001; Wang et al., 2004; Sock et al., 2004). Previously none of these genes has been associated with *Hand2. Hand2* is a BMP-downstream target (Howard et al., 2000; Liu et al., 2005b). BMP is an important signaling molecule regulating aspects of OFT development including cushion transition and valvulogenesis (reviewed in Délot, 2003; Jia et al., 2007).

Conditional targeted deletion of *Hand2* in the neural crest resulted in DORV with associated VSD with 100% penetrance. Interestingly, neither DORV nor VSD is reported in a similar line of mice where *Hand2* was conditionally deleted in the neural crest (Morikawa and Cserjesi, 2008). Although this could reflect phenotypic differences between these two *Hand2* conditional alleles, our examination of the mice generated in the Cserjesi laboratory (kindly provided by Peter Cserjesi) demonstrated that these mice also present DORV and associated VSD (Firulli and Vincentz, unpublished observations). Taken together, studies reporting on OFT development (current paper) and aortic arch development (Morikawa and Cserjesi, 2008) define fairly completely consequences on OFT and aortic arch artery development due to targeted deletion of *Hand2*. Additionally, we identified a number of *Hand2*-target genes correlated with or identified as susceptibility loci for VSD and/or DORV (Zhang et al., 2006). We identified *Gli3* as a direct Hand2 target and whose expression is significantly reduced as a consequence to loss of Hand2 in cardiac neural crest. The decreased expression of *Gli3* is of interest as *Gli3* has been identified as a susceptibility locus for VSD (Qui et al., 2006) and is associated with *Hand2* expression and function in the developing limb bud (te Welscher et al., 2002a; te Welscher et al., 2002b; Liu et al., 2005a; Barnes and Firulli, 2009). Gli3 represses sonic hedgehog (SHH), a signaling molecule associated with second heart field development but not associated with OFT morphogenesis (Dyer and Kirby, 2009). *Sox11* is another gene associated with DORV and VSD (Sock et al., 2004) that we identified as a direct Hand2 target and which is downregulated in the neural crest in response to deletion of *Hand2*. A number of genes regulated in response to deletion of *Hand2*, which we showed to be direct Hand2 targets and regulated in the neural crest, (Hey1, Foxc1, Mmp14) are related to the Notch signaling pathway. Notch signaling is implicated in migration and development of the cardiac cushions. Expression of *Hand2* in neural crest-derived neural precursor cells down-regulates expression of Hes1 (Howard unpublished result); Hey 1 is regulated in the cardiac neural crest. This intriguing result raises the possibility of identifying networks of Hand2 target genes that affect cell cycle and cell-cell contact in a number of differentiating

differentiation. The constellation of birth defects associated with disrupted development of the OFT appears to represent a common phenotype susceptible to multiple sites of regulation. The results of our gene profiling have identified *Hand2* as a critical transcriptional regulatory factor that directly or indirectly impacts expression of genes associated with all aspects of OFT development. Given the complexity of OFT morphogenesis and the role Hand2 plays in multiple cell lineages that contribute to its formation, our studies add insight into the cell autonomous neural crest

cell types but which together mediate morphogenetic movements, proliferation and

Supplementary Material

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contributions to OFT morphogenesis.

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Fig. 1. Cardiac development is affected in *Hand2* **hypomorph embryos**

Whole hearts were removed from *Hand2fl/fl* [A] and *Hand2flneo/flneo* [B] E18.5 embryos and 20 μm frozen transverse sections were visualized following staining with Hematoxylin and Eosin. Misexpression of *Hand2* [B] consistently results in enlarged hypoplastic right ventricles and disorganized trabeculations. Abbreviations: right ventricle (RV), left ventricle (LV), aorta (Ao). Scale bars indicate 100 μm [A,B] and 25 m [A1,A2,B1,B2].

Fig 2. Conditional deletion of *Hand2* **effects outflow tract development**

Whole hearts were removed from *Hand2fl/fl* (WT) [A] and *Hand2fl/fl;Wnt1-Cre* (−/−) [B, C] embryos, embedded in paraffin and 20 μm transverse sections stained with Hematoxylin and Eosin. In all mutant embryos examined (5) we observed membranous VSDs (B, black arrow) and double outlet right ventricle (C, DORV, arrows). Based on the expression pattern of GFP in comparable cryosections of E18.5 *Hand2wt/wt;Wnt1-Cre/R26RYFP* reporter [D] and *Hand2fl/fl;Wnt1-Cre/R26RYFP* mutant reporter [E–F] embryo hearts, the defects that we observe following deletion of *Hand2* can be ascribed to neural crest-derived anomalies. Abbreviations: conotruncus (CT), aorta (Ao).

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Fig 3. Conditional Deletion of *Hand2* **effects cell-cell interactions and migration of cardiac neural crest-derived cells**

Migrating neural crest-derived cells were visualized in E10.5 *Hand2wt/wt;Wnt1-cre/R26RYFP* reporter [A–B] and *Hand2fl/del;Wnt1-cre/R26RYFP* mutant reporter hearts [C–D] immunostained for YFP expression in whole-mount. Confocal optical sections (1μm) were compressed into a z-stack (~400 sections) and analyzed. The pattern of migrating neural crest-derived cells observed within the outflow tract of mutant embryos was strikingly affected by loss of *Hand2* expression. Neural crest-derived cells failed to migrate as a sheet and individual cells appear to have lost apparent contact with their neighbors routinely observed in control migrating cells (compare panels B and D). The number of neural crest-derived cells localized within the conotruncus (white arrows) appears reduced (see Figure 4) and the patterning of the AP Prongs is defective in the mutant compared to control hearts. The insets show high power views of the area indicated by the (*).

Fig 4. *Hand2* **affects proliferation of cardiac neural crest-derived cells**

To determine if the apparent decrease in neural crest-derived cells migrating within the outflow tract of *Hand2fl/del;Wnt1-Cre;R26R/YFP* hearts was due to a defect in cell proliferation, we determined the number of neural crest-derived cells expressing the proliferation marker, Ki-67 in E10.5 outflow tract of *Hand2wt/wt;Wnt1-cre/R26RYFP* reporter [A, C, E] and

Hand2fl/del;Wnt1-cre;R26RYFP mutant reporter hearts [B, D, F]. Cells were visualized, following immunostaining for Ki-67 (red) and GFP (green; marks YFP expressing neural crest-derived cells) in 20 μm frozen cross sections. There was no apparent difference in the number of cells expressing Ki-67 localized to second heart field. Visual inspection of the entire set of sections encompassing the developing outflow tract indicated a substantial decrease in the number of neural crest-derived cells (green) and neural crest-derived cells proliferating (green + red) within the outflow tract of mutant $[B, D]$ compared to control $[A, C]$ hearts. This result was quantified [G] by counting the number of cells co-expressing GFP and Ki-67 over the entire length of the outflow tract (N=3 for each genotype). There was a significant (P < 0.001) decrease in the number of proliferating neural crest-derived cells in response to targeted deletion of *Hand2*. [H] The percentage of mitotic figures did not differ significantly in control compared to mutant OFT.

Fig 5. Functional classes of differentially regulated genes in the heats of *Hand2wt/wt* **and** *Hand2fl/del;wnt1-cre* **embryos**

Regulated genes were grouped into 11 functional categories and graphed as a percentage of the total based on their GeneGo designation. 309 genes were differentially regulated based on analysis of the array data; 79 of these regulated genes did not have any known function and were eliminated from this analysis [A]. Of the regulated genes, 68 were up-regulated [B] and 162 were down-regulated [C]. A number of down-regulated genes (17) are associated with cell adhesion; none of the genes in this category were up-regulated. The cytoskeletal-related transcript category contained 16 down-regulated genes and 2 up-regulated genes. In the transcription factor category, a notable difference in the number of transcripts down-regulated (17) and up-regulated (4) related to *Hand2* was observed. In the kinases/phosphatases category a significant number of transcripts (11 up-regulated, 11 down-regulated) were affected by *Hand2* deletion. We have identified a number of Hand2-target genes in each of these categories that are associated with the developmental anomalies that we observe in embryos with targeted deletion of *Hand2* in the neural crest.

Fig 6. Connexin 40 expression is decreased by targeted deletion of *Hand2* **in the neural crest** Expression of Connexin 40 was examined in the outflow tract of

Hand2wt/wt;Wnt1-Cre;R26R/YFP [A,C,E] and *Hand2fl/del;Wnt1-Cre;R26R/YFP* [B,D,F] E10.5 cryostat sections encompassing the entire outflow tract. Tissue sections (20 μm) were immuno-labeled for expression of YFP (reporter, green) and Connexin 40 (red). The entire outflow tract was imaged at 20X and 63X; 0.5 μm optical sections were compressed in the z-plane and assessed. Ten pairs of matched sections were examined. The majority of neural crest-derived cells migrating within the outflow tract in control embryos [A,C,E] express connexin 40 at the cell surface and in contact with neighboring cells. In *Hand2* mutant embryos, only a few neural crest-derived cells maintain expression of Connexin 40 [B,D,E]; the expression pattern appears normal in those few cells that maintain Connexin 40. The arrows point to connexin 40 expressed in the myocardium. Abbreviation: Cx40 (Connexin 40).

Fig. 7. Connexin 40 is a direct Hand2 target gene

[A] Schematic representation of the E-boxes (consensus: CANNTG) located within genomic region 5′ to Exon 1 (ExI) of the *Cx40* gene. Amplicons of the distal (Cx40 [−1144 – −814]) and proximal (Cx40 $[-655 - -283]$) primer sets (arrows) used for ChIP assays are denoted. [B] ChIP using a Hand2-specific antibody co-immunoprecipitates DNA fragments overlapping the proximal (Cx40 [−655 – −283]), but not distal (Cx40 [−1144 – −814]), primer sets, indicating that Hand2 binds to the *Cx40* promoter within this region. [C] Luciferase-reporter transactivation assays in HEK 293 cells show that Hand2 and E12 cooperatively up-regulate expression of a *Cx40*-luciferase reporter (Cx40-pX2P) \sim 2.2 fold over baseline (n = 3, p < 0.001). Individually, Hand2 and E12 do not significantly influence Cx40-pX2P expression.

Table 1

Antibodies used for immunostaining in this study are listed. The antigen, dilution, and secondary antibodies are listed. The source for each antibody is listed. Antibodies used for immunostaining in this study are listed. The antigen, dilution, and secondary antibodies are listed. The source for each antibody is listed.

Table 2

Hand₂ Target Genes Hand2 Target Genes

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indicative of a potential binding site. Within the three confidence levels, FDR ≥ 0.05 represents the highest confidence levels, FDR ≥ 0.1 is the second highest confidence level and FDR $> 0.1 \leq 0.2$ is the

lowest confidence level where binding remains statistically significant. The primary data for the heart micro-arrays is available at the NIH microarray consortium website

http://np2.ctrl.ucla.edu/np2/home.do. To navigate to the data, click on the "navigate repository" button and write Howard under investigator and click go. The project will come up and the data can be accessed
by clicking o NIH-PA Author ManuscriptNIH-PA Author Manuscript

by clicking on the magnifying glass.

<http://np2.ctrl.ucla.edu/np2/home.do>. To navigate to the data, click on the "navigate repository" button and write Howard under investigator and click *go*. The project will come up and the data can be accessed