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Functional Attenuation of Ufd1l, a 22q11.2 Deletion Syndrome Candidate Gene, Leads to Cardiac Outflow Septation Defects in Chicken Embryos

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ABSTRACT

Microdeletion of chromosome 22q11.2 is commonly associated with congenital cardiovascular defects that involve development of cranial neural crest cells (NCC) that emigrate through the pharyngeal arches. UFD11 is one of several candidate genes for 22q11.2 deletion syndrome (22q11DS). UFD11 encodes a protein whose yeast counterpart is involved in a ubiquitindependent proteolytic degradation pathway; however, the role of UFD1L in NCC development remains unknown. Mouse embryos that lack Ufd11 die before organogenesis. We have therefore studied the function of Ufd11 in the chick system. Chick Ufd11 encoded a 307-amino acid protein that was highly conserved with mouse and human UFD1L. Chick Ufd11 was expressed in the developing neural tube, NCC, and mesenchyme of the head and pharyngeal arch structures, as well as in the conotruncal region (cardiac outflow tract), consistent with the clinical features of 22q11DS. To determine loss-of-function effects of chick Ufd11 in NCC, we infected cardiac NCC with a retrovirus expressing antisense Ufd11 transcripts in chick embryos before their migration. Morphologic analysis of infected embryos at a later developmental stage demonstrated that functional attenuation of chick Ufd11 in cardiac NCC resulted in an increased incidence of conotruncal septation defects. These data suggest that Ufd11 may play a role in cardiac NCC during conotruncal septation. (*Pediatr Res* 53: 546–553, 2003)

Abbreviations

CVD, cardiovascular defects
del.22q11, microdeletion of chromosome 22q11.2
ET_A, endothelin receptor A
HH, Hamburger and Hamilton
NCC, neural crest cells
VSD, ventricular septal defect
22q11DS, 22q11.2 deletion syndrome

Monoallelic microdeletion of chromosome 22q11.2 (del.22q11) is the most common human chromosomal deletion and results in the 22q11.2 deletion syndrome (22q11DS), with an estimated frequency of 1 in 4000–5000 births (1). 22q11DS encompasses three independently established syndromes, including DiGeorge syndrome, velo-cardio-facial syndrome, and conotruncal anomaly face syndrome, and manifests a variety of

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features, including craniofacial and cardiovascular defects (CVD) and behavioral and psychiatric disorders (2–4). 22q11DS is associated with specific types of CVD that are characterized as malformations of the conotruncus (outflow tract of the heart) or the aortic arch, including persistent truncus arteriosus, tetralogy of Fallot, and interrupted aortic arch, type B (5). The craniofacial defects and CVD in 22q11DS result from defects of embryonic structures populated by migrating cranial and cardiac neural crest cells (NCC) that originate from the caudal hindbrain (6, 7). Ablation of cardiac NCC in chick embryos results in persistent truncus arteriosus, aortic arch malformations, and hypoplasia of thymus and parathyroid glands similar to that observed in patients with 22q11DS (6, 7). This suggests that studies on candidate genes in the 22q11.2

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region that are involved in the development of cardiac NCC may provide insight into the molecular mechanisms underlying the 22q11DS and its associated CVD.

More than 90% of patients with 22q11DS have a similar del.22q11, commonly spanning 1.5-3 megabases in size (8). More than 30 genes have been identified in the deleted region. Recent studies show that haploinsufficiency of *Tbx1*, one of the genes in this region, is likely to be the major determinant of aortic arch defects in patients with 22q11DS (9-11). However, no single gene mutation, including that of TBX1, that results in the 22q11DS phenotype has yet been detected (12, 13). Our screen for mouse genes dependent on dHAND, a transcription factor implicated in NCC development, previously identified Ufd11, whose human ortholog had been mapped to 22q11.2 (14, 15). UFD1L encodes a highly conserved protein from yeast to human that is involved in the degradation of ubiquitinated proteins. Mouse Ufd11 was expressed in many embryonic tissues affected in patients with 22q11DS, including the pharyngeal arches, palatal precursors, the frontonasal region, telencephalon, and the cardiac outflow tract. Human UFD1L was deleted in >180 patients studied with 22q11DS (15). A smaller deletion of exon 1 to 3 of UFD1L and its neighboring gene, CDC45, which encodes a putative cell cycle regulating protein, was found in one individual with the phenotype of 22q11DS. These results suggest that UFD1L may be implicated in some features of 22q11DS, although the very early lethality of mice that lack Ufd11 precludes an understanding of the role of UFD1L in cardiac NCC development (16).

The avian system is well established to study NCC development, as gene constructs can be specifically introduced into the chick NCC lineage using a replication-incompetent retrovirus system (17). To take advantage of the avian system, we cloned the chick ortholog of *UFD1L* and performed expression and functional analyses of chick Ufd11 during embryogenesis, focusing on cardiac NCC development. Chick *Ufd11* was expressed in a variety of developing tissues with enhanced expression in the cardiac outflow tract and neural crest-derived mesenchyme of pharyngeal arches and aortic arch arteries, similar to mouse *Ufd11*. Attenuation of chick Ufd11 expression in premigratory cardiac NCC resulted in an increased number of conotruncal septation defects. These data suggest that Ufd11 may play a role in the development of cardiac NCC that contribute to conotruncal septation.

METHODS

Cloning of chick Ufd11. A stage 12–15 chick embryo library in lambdaZAPII was screened according to the manufacturer's instructions (Stratagene), with chick Ufd11 partial cDNA (741bp) that was obtained from Hamburger and Hamilton (HH) stage 12 chick total RNA by reverse-transcribed PCR. Approximately 1 μ g of total RNA was reverse transcribed using M-MuLV transcriptase (Roche). One microliter of the first-strand cDNA produced was used as template for PCR reactions using specific oligonucleotide amplimers (5'-GGAGAAAGGAGGGAAGATAA-3', 5'-GCTTCATCCTCT-TCAACCTT-3') derived from mouse Ufd11 cDNA. 5' rapid amplification of cDNA ends was then performed to obtain full-length cDNA according to manufacture's instructions (Clontech).

Northern analysis. Total RNA from several chick tissues at embryonic day 18 was electrophoresed, transferred to a nylon membrane, and hybridized with a ³²P-labeled full-length chick Ufd11 cDNA probe. Hybridization was performed in Rapidhyb buffer (Amersham) at 65°C overnight and washed with 2 \times salt-sodium chloride/0.1% SDS at room temperature for 10 min and 1x salt-sodium phosphate-EDTA/1% SDS at 65°C for 15 min. Nylon membrane was exposed to x-ray film (Kodak).

Collection of chick embryos and primary NCC culture experiments. Fertilized eggs were incubated at 37°C to obtain chick embryos of HH stage 10-26. The cranial neural fold from the otic vesicle to somite 3 of stage 9–11 chick embryos was dissected and treated with 0.5 mg/mL collagenase type II in PBS at 37°C for 2-3 min to remove the mesenchyme surrounding the neural tube. The isolated neural tube was washed in Dulbecco's modified Eagle's medium including 10% fetal bovine serum, then placed on gelatin-coated culture plates. The dissected neural tubes were cultured at 37°C for 36-48 h in Dulbecco's modified Eagle's medium including 10% fetal bovine serum to allow NCC to migrate from the neural tubes onto culture plates. NCC were fixed with 4% paraformaldehyde/PBS. In situ hybridization and immunohistochemistry were subsequently performed. All experimental procedures were performed according to the current local guidelines for animal experiments in the University of Texas Southwestern Medical Center and Leiden University Medical Center.

RNA in situ hybridization. RNA in situ hybridization of chick embryos and cultured NCC was performed as previously described (18) with digoxigenin-labeled Ufd11 riboprobes synthesized from a 741-bp fragment of chick Ufd11 cDNA. Briefly, chick embryos of varying stages were isolated, and the pericardium was removed. Embryos were fixed in 4% paraformaldehyde/PBS at 4°C overnight. Embryos were treated with proteinase K (30 μ g/mL) for 3–20 min at room temperature depending on their stage of development. Prehybridization was performed at 61°C for 3 h in solution containing 50% formamide, $4 \times$ salt-sodium chloride, $1 \times$ Denhardt's solution, 500 μ g/mL salmon sperm DNA, 250 μ g/mL yeast tRNA, and 10% dextran sulfate. Hybridization was performed in the same solution for 18 h. After a series of washes, embryos were treated in 2% blocking solution for 2 h and incubated with antidigoxigenin antibody for 1 h at room temperature. Color reaction was performed at room temperature in substrate color reaction mixture for 12–18 h and terminated by fixing embryos in 4% paraformaldehyde/0.1% glutaraldehyde.

Radioactive *in situ* hybridization was performed using ³⁵Slabeled antisense riboprobes synthesized with T7 or SP6 RNA polymerase (MAXIscript, Ambion Inc.) from chicken dHAND or neuropilin-1 full-length cDNA, respectively. Paraffinembedded coronal sections of HH stage 18 control embryos and antisense Ufd11-infected embryos were hybridized with probes as previously described (18). Plasmids for dHAND and neuropilin-1 riboprobe are available upon request.

Fixed NCC were treated with proteinase K (30 μ g/mL) for 5 min at 37°C. After postfixation with 4% paraformaldehyde/

PBS, cells were dehydrated in methanol. *In situ* hybridization was performed at 61°C for 18 h as described above.

Immunocytochemistry. For immunocytochemistry, fixed and dehydrated NCC were incubated with 3% H₂O₂ for 30 min at room temperature. After rehydration, cells were incubated in blocking solution [3% BSA, 0.1% Igepal (Sigma Chemical Co.) in PBS] at room temperature for 30 min and then incubated in primary antibody, anti-HNK-1/N-CAM (Sigma Chemical Co.; 1:60), at room temperature for 60 min. After several washes in PBS, conjugated secondary antibody [antimouse IgM (1:50)] was applied to the cells and incubated at room temperature for 60 min. The cells were then incubated in developing solution [0.03% 3-3'-diaminobenzidine tetrahydrochloride (Sigma Chemical Co.) -0.1% TritonX in PBS] at room temperature for 10 min. Color reaction was performed in 0.03% H₂O₂ at room temperature for 10 min. After several washes in PBS, cells were fixed with 4% paraformaldehyde/ 0.1% glutaraldehyde for 12-24 h.

In vitro transcription and translation. The full-length chick Ufd11 cDNA was cloned in the sense and antisense direction into pBluescript plasmids. Mouse endothelin receptor A (ET_A) was also cloned in antisense direction into pBluescript plasmids as a control. All plasmids were linearized with *SmaI* for *in vitro* transcription and translation reactions using TNT-coupled Reticulocyte Lysate System (Promega) with T7 RNA polymerase. One microgram of sense cUfd11 plasmid and antisense cUfd11 or ET_A plasmid were mixed together in the same tube for transcription and translation. A sense cUfd11 plasmid and pBluescript plasmid mixture was performed simultaneously as a control. Translated products were labeled with [³⁵S] methionine (Amersham) and determined by SDS-PAGE and autoradiography. The signals were quantified using ImageQuant software (Molecular Dynamics).

Retrovirus infection of chick embryos. The full-length chick Ufd11 cDNA was cloned in the sense and antisense direction into the proretroviral pCXIZ vector (gift from Dr. Takashi Mikawa). All retroviral constructs were stably transfected to the D17.2G helper cell line for virus production. *In vivo* infection of premigrating NCC in chick embryos was performed at HH stage 9–11 as described previously (19, 20). The embryos were allowed to develop in ovo until HH stage 31–32 and then were harvested and fixed with 4% paraformal-dehyde for histologic analysis. HH stage 18 embryos were also harvested for section *in situ* hybridization using dHAND and neuropilin-1 riboprobes.

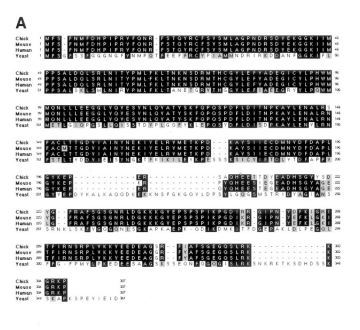
In vitro infection was performed on neural tube dissected from HH stage 9–11 as described above. After NCC migration, β -galactosidase activity was detected by using X-gal as described previously (21).

Histology. Embryos were embedded in paraffin, and serial transverse sections were obtained. The transverse sections of embryos after retrovirus infection or control embryos were stained with hematoxylin and eosin.

RESULTS

Chick Ufd11 *cDNA*. The largest cDNA obtained of chick *Ufd11* by library screening followed by 5' rapid amplification of

cDNA ends contained 1026 nucleotides. Chick *Ufd11* was predicted to encode a 307–amino acid protein and was highly conserved with its murine (94% identical, 96% similar) and human (94% identical, 96% similar) counterparts (Fig. 1A). Although the previously cloned human UFD1L protein (as translated from cDNA, GenBank accession number U64444) (14) showed a tandem repetition of a 36–amino acid domain that was present in the yeast, chicken, and mouse as a single copy, the protein sequences predicted from all human UFD1L cDNAs that we cloned (GenBank accession number



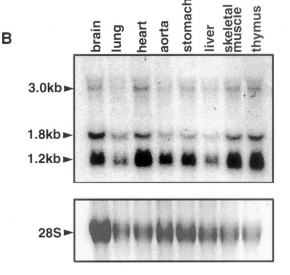


Figure 1. Sequence and expression of chick Ufd11. *A*, Sequence alignment of full-length chick Ufd11 with mouse, human, and yeast Ufd11. Amino acid residues that are identical or similar are shaded in black or gray, respectively. Chick *Ufd11* encodes a 307–amino acid protein and is highly conserved with its murine (94% identical, 96% similar) and human (94% identical, 96% similar) counterparts. It was 36% identical and 50% similar to yeast Ufd1. *B*, Northern blot analysis of chick Ufd11 in embryonic day 18 tissues. mRNA transcripts (1.2, 1.8, and 3.0 kb) were present in all tissues examined. The expression levels of chick Ufd11 in brain, heart, skeletal muscle, and thymus were higher than in other tissues.

AY101594) did not contain such a domain and were more similar to those of other species. Chick Ufd11 was also 36% identical and 50% similar to yeast Ufd1. Nucleotide and protein sequences of chick Ufd11 are available at NCBI (GenBank accession number AF407671).

Expression pattern of chick Ufd11. Northern blot analysis of total RNA from several chick tissues at embryonic day 18 was performed using full-length chick Ufd11 cDNA as a probe. mRNA transcripts (1.2, 1.8, and 3.0 kb) were present in all tissues examined and possibly represented alternatively spliced forms (Fig. 1B). The expression level of chick Ufd11 in each tissue was variable, with expression in brain, heart, skeletal muscle, and thymus relatively higher than in other tissues.

To determine the expression pattern of chick *Ufd11* during embryogenesis, *in situ* hybridization analysis was performed. Chick *Ufd11* was detectable by stage 7 in the neural plate (data not shown). At stage 10, as the neural tube and somites formed, expression throughout the neural tube was apparent (Fig. 2A). At later stages, chick *Ufd11* was expressed in many tissues, with marked expression in the head, pharyngeal arches, frontonasal region, neural tube, and limb buds (Fig. 2, B and C). Whole-mount *in situ* hybridization on the heart dissected from embryos at stage 26, when conotruncal septation takes place, demonstrated strong expression of chick *Ufd11* in the conotruncal region of the heart and in emerging aortic arch arteries (Fig. 2D). The expression of chick *Ufd11* in the neural tube and mesenchymal cells along the pharyngeal arch arteries was detected in transverse sections through chick embryos after whole-mount *in situ* hybridization (Fig. 2, E and F). The overall embryonic expression pattern of chick *Ufd11* was similar to that of mouse (14, 15) and consistent with some of the clinical features of 22q11DS.

To determine whether chick Ufd11 was expressed in migrating NCC, we examined primary chick NCC cultures. NCC that had migrated off chick neural tube in culture exhibited normal expression of the avian NCC marker HNK-1 by immunocytochemistry (Fig. 3A). *In situ* hybridization in the cultured NCC demonstrated chick Ufd11 expression in the migrating NCC, consistent with the result of whole-mount *in situ* hybridization analysis on embryos (Fig. 3, B and C).

Functional attenuation of chick Ufd11 by retroviral infection. To investigate the functional role of chick Ufd11 in cardiac NCC, we used a well-established gene introduction system in chick NCC using a replication-incompetent retrovirus (17). To study loss of function effects *in vivo*, we prepared a Ufd11 antisense retroviral vector that would generate Ufd11 antisense transcripts. To determine first whether the Ufd11 antisense transcripts might interfere with normal translation of Ufd11 sense transcripts, we performed *in vitro* transcription and translation experiments. As shown in Figure 4, the protein

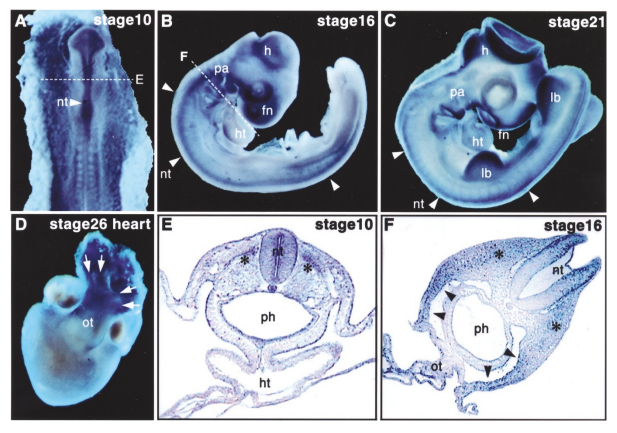


Figure 2. Expression of chick Ufd11 during embryogenesis. *In situ* hybridization of chick *Ufd11* on stage 10, 16, or 21 chick embryos (A–C) and on heart of stage 26 embryo (D). At stage 10, the expression throughout the neural tube (nt) was apparent (A). At later stages, chick Ufd11 expression was enhanced in the head (h), pharyngeal arches (pa), frontonasal region (fn), neural tube (nt, *white arrow heads*), and limb buds (lb) (B and C). D, Strong expression of chick Ufd11 in the outflow tract (ot) of the heart and arch arteries (*white arrows*). Transverse section of embryo in A and B after whole-mount *in situ* hybridization (E and F, respectively) is shown at levels indicated. The expression was observed in the neural tube (nt), cranial mesenchyme (*), aortic arch arteries (*black arrow heads*), and outflow tract (ot) (E and F). ph, pharynx; ht, heart.

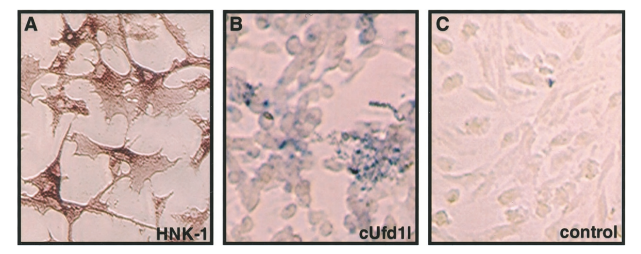


Figure 3. Chick *Ufd11* is expressed in primary neural crest culture. *A*, HNK-1, as a NCC marker, was detected by immunocytochemistry in primary neural crest cultured cells. *B*, Chick Ufd11 transcripts were detected by antisense mRNA *in situ* hybridization in primary cultured cells. *C*, *In situ* hybridization with sense Ufd11 riboprobe as control.

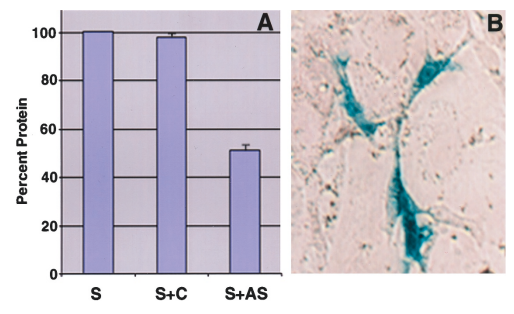


Figure 4. Antisense chick Ufd11 transcripts can attenuate chick Ufd11 protein expression. *A*, Quantitative analysis of *in vitro* transcription and translation reveals a 54% reduction in Ufd11 protein when Ufd11 sense (S) and antisense (AS) transcripts were co-incubated compared with S alone or S with a control (C) antisense (ET_A). The expression of ET_A antisense transcripts did not affect protein expression of chick Ufd11. *B*, Primary NCC culture after neural tube infection with retrovirus containing the AS Ufd11 expression construct linked to β -galactosidase. Survival and migration of infected cells (blue) were unaffected.

expression of chick Ufd11 was reduced ~50% by expression of the antisense chick Ufd11 transcript. The expression of control (ET_A) antisense transcripts did not affect the protein expression of chick Ufd11, suggesting that the effect of antisense chick Ufd11 introduction was specific for the reduction of chick Ufd11 protein expression. To determine whether cells infected with the Ufd11 antisense vector had a nonspecific survival defect, we performed primary neural crest culture from neural tube infected with the Ufd11 antisense retrovirus. Retrovirusinfected cells were marked in blue by β -galactosidase staining, and we found that their survival was unaffected by Ufd11 antisense transcripts (Fig. 4B).

For determining the functional role played by chick Ufd11 in cardiac NCC, the rostral neural tube was infected with retrovirus containing the antisense Ufd11 expression construct before NCC migration. By examining the hearts of a number of embryos at HH stage 31–32, 6 d after the retroviral infection, we found a significant (p < 0.05) increase in septal defects (nine of 12) compared with control embryos (two of 12 and zero of 12, respectively) that were infected with retrovirus containing an empty vector (Fig. 5, A–F). Five of 12 experimental embryos had defects of the upper portion of the ventricular septum (VSD; Fig. 5, B and E). An additional four of 12 embryos had VSD and defects in septation of the aorta and pulmonary artery (conotruncal septal defects; Fig. 5, C and F). This combination was never observed in control embryos (p < 0.05), whereas isolated VSD were observed in two of 12 control embryos. Examination of the aortic arch in all embryos failed to detect any defects at this stage. Results are summarized in Table 1.

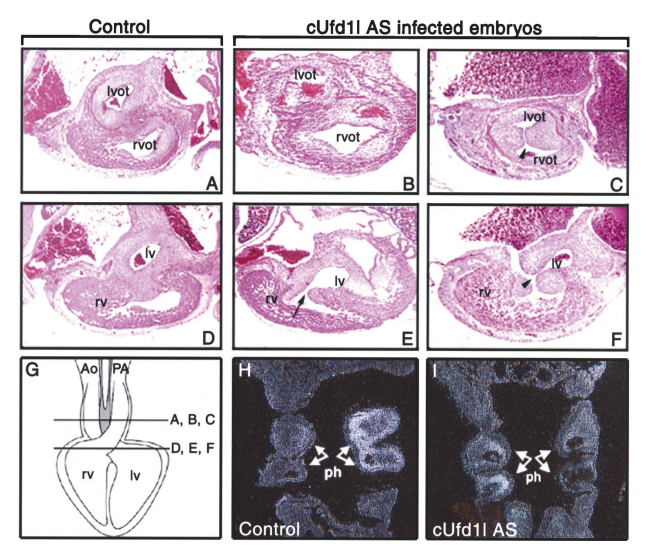


Figure 5. Attenuation of Ufd11 in chick neural crest results in conotruncal defects. Transverse sections of the stage 31-32 chick hearts at the level of the conotruncal septum (*A*–*C*) or ventricular septum (*D*–*F*) in control embryos (*A* and *D*) or retrovirus Ufd11 antisense (AS) infected embryos (*B*, *C*, *E*, and *F*). *G*, Schematic indicating level of section. Ventricular septal defects (*arrow*) were observed in AS-infected embryos (*B* and *E*). Conotruncal septal defects (*arrowhead* in *C*) accompanied with VSD (*arrowhead* in *F*) were observed in AS-infected embryos. Radioactive RNA *in situ* hybridization on coronal sections of HH stage 18 embryos revealed similar expression of dHAND, a marker of neural crest–derived mesenchyme, in control (*H*) and AS-infected (*I*) embryos. Ao, aorta; PA, pulmonary artery; Iv, left ventricle; lvot, left ventricular outflow tract; rv, right ventricle; rvot, right ventricular outflow tract; ph, pharyngeal arch.

Table 1. cUfd11	AS retroviru:	s-infected	embryos	display	cardiac
	outflow	tract defe	cts		

	cUfd11 AS	Control vector	
Septal defects	75% (9/12)*	17% (2/12)	
Conotruncal	33% (4/12)*	0% (0/12)	
Ventricular	42% (5/12)	17% (2/12)	
Aortic arch defects	0% (0/12)	0% (0/12)	
No defects	25% (3/12)*	83% (10/12)	

Percentage of affected embryos are indicated; absolute number of affected/ total embryos are also shown.

* p < 0.05.

To investigate whether NCC were specified and migrated in antisense Ufd1-infected embryos, we examined two molecular markers, dHAND and neuropilin-1, both expressed in the NCC-derived mesenchyme of pharyngeal arches. Expression of dHAND (Fig. 5, H and I) and neuropilin-1 (data not shown) was relatively normal in the pharyngeal arches of HH stage 18 experimental embryos compared with controls. This result suggests that NCC become specified, migrate, and differentiate into mesenchymal cells during early pharyngeal development of antisense Ufd11-infected embryos. Taken together, these data indicate that functional attenuation of Ufd11 in cardiac NCC leads to an increased incidence of conotruncal septal defects in chick embryos and these defects do not result from nonspecific cell death before NCC migration.

DISCUSSION

Cross-species analysis is sometimes useful in understanding potential causes of human disease. Mice homozygous for deletion of *Ufd11* are early embryonic lethal, whereas mice heterozygous for a *Ufd11* mutation are normal, precluding an examination of whether Ufd11 plays a role in cardiac NCC development (16). Antisense nucleotides designed to hybridize to specific mRNA sequences have been used to attenuate the

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function of numerous genes both *in vitro* and *in vivo*, including developmentally regulated genes in chick embryos (22–24). By using a replication-incompetent retrovirus that has proved beneficial in studying the avian NCC lineage, we were able to introduce antisense *Ufd11* transcripts into chick cardiac NCC over long time spans and examine the role of Ufd11 in cardiac NCC *in vivo*.

Although functional attenuation of Ufd11 in NCC lineage resulted in an increased number of conotruncal septal defects in chick embryos, there was phenotypic variation. It is possible that the phenotypic variation may reflect the gene dosage effect of Ufd11 resulting from a variable efficiency of retroviral infection in each chick embryo. If this is the case, then conotruncal septation requires Ufd11 in a dose-sensitive manner in this chick model, although haploinsufficiency of Ufd11 alone is not sufficient for the septal defects in the mouse model. Alternatively, it may reflect allelic variability, variable penetrance caused by environmental factors, or stochastic events during embryonic development similar to that proposed in patients with 22q11DS. We cannot distinguish between these possibilities as it is not possible to examine effectively the level of attenuation of Ufd11 protein in premigratory NCC of embryos studied later in development for defects in cardiogenesis.

Recent genomic manipulations in mice have provided insight into potential roles of candidate genes for 22q11DS. Mice harboring a hemizygous deletion of 18 genes, including Ufd11, that are deleted in patients with 22q11DS (Df1/+) were generated and exhibit aortic arch malformations in ~30% of cases (16). Further analyses determined that haploinsufficiency of Tbx1 was responsible for aortic arch malformations in Df1/+ mice (10). Mice homozygous for the *Tbx1* mutation displayed craniofacial and cardiovascular defects similar to 22q11DS (9-11), suggesting that *TBX1* may be a major determinant of the phenotype of 22q11DS. However, TBX1 mutations have not been identified in patients who have the 22q11 deletion phenotype and do not have del.22q11 (10, 13). In addition, Df1/+ mice have broader phenotypic features that have not been observed in mice heterozygous for a Tbx1 mutation, including conotruncal and ventricular septal defect, hypoplasia of thymus and parathyroid glands, and signs of behavioral and psychiatric disorder (16, 25, 26). These results indicate that haploinsufficiency of Tbx1 and additional genes in the Df1deletion region together contribute to the entire phenotype of Df1/+ mice. Our data in the present study suggest a role for Ufd11 in conotruncal development and thus support a possibility that Ufd11 may contribute to conotruncal and ventricular septal defects as a modifier gene in Df1/+ mice. Ufd11 might also contribute to the hypoplasia of the thymus/parathyroid glands and behavioral/psychiatric phenotype of Df1/+ mice, because it is expressed in the developing pharyngeal structures and brain. The association of a promoter polymorphism of UFD1L with schizophrenia susceptibility has recently been reported (27). Definitive evaluation of Ufd11's role in particular tissues awaits tissue-specific disruption of Ufd11 in mice, currently being performed in our laboratory.

Although we determined that Ufd11 plays a role in cardiac NCC, cellular functions of Ufd11 in avian or mammalian cells remain to be studied. We speculate that Ufd11 is required for

postmigratory differentiation of cardiac NCC in the conotruncal region because *1*) the infection of retrovirus containing antisense *Ufd11* did not affect NCC survival and migration in primary NCC; *2*) Ufd11 was expressed at high levels in the conotruncal region; and *3*) functional attenuation of Ufd11 resulted in conotruncal septation defects, although the phenotype was milder than that observed after ablation of premigratory NCC in chick embryos. Recent biochemical studies demonstrated multiple functions of Ufd1, including the activation of membrane-bound transcription factors and transportation of proteins from the endoplasmic reticulum into the cytosol, by regulating ubiquitin/proteosome-dependent processing (28– 31). Identification of pathways affected by Ufd11 in cardiac NCC will be necessary to understand the mechanism through which Ufd11 regulates NCC development.

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