UDP-glucose Dehydrogenase Polymorphisms from Patients with Congenital Heart Valve Defects Disrupt Enzyme Stability and Quaternary Assembly*

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Background: UDP-glucose dehydrogenase (UGDH) polymorphisms were identified in a screen of candidate genes for heart valve defects.

Results: Two individual mutants fail to rescue cardiac valve defects in UGDH-deleted zebrafish and have reduced stability in vitro.

Conclusion: UGDH loss of function mutations result in a subset of human congenital cardiac valve defects caused by reduced enzyme activity during morphogenesis.

Significance: Screening these alleles could predict valve defects.

Cardiac valve defects are a common congenital heart malformation and a significant clinical problem. Defining molecular factors in cardiac valve development has facilitated identification of underlying causes of valve malformation. Gene disruption in zebrafish revealed a critical role for UDP-glucose dehydrogenase (UGDH) in valve development, so this gene was screened for polymorphisms in a patient population suffering from cardiac valve defects. Two genetic substitutions were identified and predicted to encode missense mutations of arginine 141 to cysteine and glutamate 416 to aspartate, respectively. Using a zebrafish model of defective heart valve formation caused by morpholino oligonucleotide knockdown of UGDH, transcripts encoding the UGDH R141C or E416D mutant enzymes were unable to restore cardiac valve formation and could only partially rescue cardiac edema. Characterization of the mutant recombinant enzymes purified from Escherichia coli revealed modest alterations in the enzymatic activity of the mutants and a significant reduction in the half-life of enzyme activity at 37 °C. This reduction in activity could be propagated to the wild-type enzyme in a 1:1 mixed reaction. Furthermore, the quaternary structure of both mutants, normally hexameric, was destabilized to favor the dimeric species, and the intrinsic thermal stability of the R141C mutant was highly compromised. The results are consistent with the reduced function of both missense mutations significantly reducing the ability of UGDH to provide precursors for cardiac cushion formation, which is essential to subsequent valve formation. The identification of these polymorphisms in patient populations will help identify families genetically at risk for valve defects.

Congenital heart defects, approximately one-fourth of which result from improper valve or septal formation (1), are a prevalent health concern (2). In many cases, the defects may be mild, and patients are not diagnosed until later in life, or the early defect is managed surgically but continues to require vigilant health awareness. Several genetic mutations have been correlated with these conditions (3, 4), but only a few of the molecular factors critical for normal heart development in humans have been well characterized.

Heart morphogenesis is a complex multistep process involving changes in expression of hundreds of different genes, as well as input from physical stimuli during development (3, 4). Model organisms such as zebrafish have been invaluable for delineation of important signaling molecules and receptors that modulate the necessary changes (5–8). In addition to specific signals, cell surface and extracellular adhesive molecules such as proteoglycans and glycosaminoglycans are essential for presentation of small molecules, local magnification of their potency, and generation of the hydrodynamic cushions ultimately needed for heart chamber and valve differentiation (3, 9, 10). The roles of genes involved in synthesis of these diverse anionic carbohydrate polymers have been defined by elegant genetic studies in which pre-cardial tissue from lethal disruptions is explanted and developmentally rescued with transduction of the targeted gene (9, 11).

UDP-glucose dehydrogenase (UGDH)3 is one candidate gene essential for development in frogs (12), fruit flies (13),...
nematodes (14), and mice (9, 15), shown in zebrafish to be critical specifically for cardiac valve formation (11). This enzyme produces UDP-glucuronate, key sugar nucleotide precursor for incorporation into the extracellular matrix glycosaminoglycan hyaluronan (16), modification of lipophilic hormones for enhanced solubilization and elimination (17), polymerization of heparan sulfate and chondroitin sulfate chains, as well as conversion to UDP-xylose to initiate proteoglycan production (10). Studies in bacterial (18, 19), plant (20, 21), bovine (22), and human (23) UGDH homologues show that all of them catalyze two successive oxidations of the 6′ carbon to orchestrate the same conversion of UDP-glucose to UDP-glucuronate (18–23). Notably, the enzyme is active in bacteria as a dimer (24), although in mammals and higher eukaryotes such as plant, nematode, fruit fly, and zebrafish, the enzyme purifies and functions as a hexamer (23, 25). This structure is a trimeric assembly of dimeric units (26), which are homologous to the bacterial dimer, organized in a toroidal formation that has been visualized by electron microscopy (25) and x-ray crystallography (see Fig. 1).

The structure has been solved for the human UGDH T131A apo form (Protein Data Bank code 3ITK), as a ternary complex with NAD⁺ cofactor and UDP-glucuronate product (Protein Data Bank code 2QG4), as a ternary complex with NADH cofactor and UDP-glucose substrate (Protein Data Bank code 2Q3E) (27), and for UGDH E161Q as a thiohemiacetal between Cys-276 and the singly oxidized UDP-glucose, captured following the addition of NAD⁺ and UDP-glucose (Protein Data Bank code 3KHU) (28). In a prior study, we used the human UGDH model to identify residues likely to be critical for catalytic activity and/or binding of the substrate and cofactor, which we subsequently confirmed using site-directed mutagenesis and enzymology (23, 29).

In a previously described large scale sequencing experiment using candidate genes essential for atrioventricular (AV) canal, AV septum, and AV valve formation, three patients with mitral valvar prolapse and mitral regurgitation were identified with novel missense mutations in the UGDH gene from among 192 patients with clinically diagnosed AV septal defects (30). Two of these individuals were heterozygous for a single nucleotide substitution that converted Arg-141 of UGDH to cysteine (R141C). A third individual was heterozygous for a mutation that converted Glu-416 to aspartate (E416D). Neither of these mutations on steady state enzyme activity, quaternary structure, and intrinsic stability of the protein. This comprehensive approach provided insights into the underlying molecular causes of valve defects in patients with UGDH polymorphisms and revealed novel aspects of UGDH enzymatic function.

**EXPERIMENTAL PROCEDURES**

**In Situ Hybridization**—Embryos were fixed in 4% formaldehyde, embedded in paraffin, and sectioned for *in situ* hybridization (10–14 μm). Labeled RNA probes were used to detect cardiac tropomyosin I or UGDH mRNA expression by *in situ* hybridization according to a previously described method (31). Representative images were digitally captured to illustrate message expression relative to AV canal and outflow tract.

**Cardiac Valve Morphogenesis and in Vivo Rescue of Cardiac Edema in Zebrafish**—Wild-type and transgenic (Tg(Tie2: EGFP) by zebrafish were maintained under standard conditions (32). Two independent antisense morpholino oligonucleotides (MOs) were used to knock down Ugdh expression: ugdh-ATG MO, which was previously described (11), and ugdh-Splice MO (supplemental Fig. S1, 5′-AACAGCTCACA-CAAAACACACCTGTA-3′). MOs were injected into embryos at the one cell stage (4 ng of ugdh-ATG MO or 2 ng of ugdh-Splice MO). Valve morphology was visualized by fluorescence microscopy on a Zeiss dissecting microscope. Representative digital images were captured at 48 h post-fertilization (hpf). For
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cardiac phenotype rescue experiments, in vitro transcribed mRNA encoding UGDH wild type or R141C or E416D mutants was injected (10 μg/ml, 1 nl/embryo) concurrently with Ugdh MOs (4 ng/embryo) at the one-cell stage. Cardiac edema was evaluated and quantified at 55 hpf.

Hyaluronan Detection—To confirm functional Ugdh knockdown in embryos, we stained for hyaluronan on paraffin sections as described previously (33). Serum-free medium of HEK293 cells expressing a neurocan-alkaline phosphatase fusion protein was a generous gift from Uwe Rauch (University of Lund).

Generation and Purification of UGDH Point Mutants—Point mutants of human UGDH were generated from the wild-type constructs, UGDH-pET28a (for protein purification) or UGDH-pCS2 (for RNA injections in zebrafish), using the QuikChange site-directed mutagenesis kit (Stratagene, La Jolla, CA) according to the manufacturer’s protocol. The sequences were verified by the Genomics Core at the University of Nebraska-Lincoln. Point mutants were expressed in Escherichia coli strain Rosetta2(DE3)pLysS (EMD Biosciences, Inc., San Diego, CA) and purified as N-terminal His6 fusions as previously described (23). Following nickel affinity chromatography, the proteins were dialyzed against 0.1 m sodium phosphate, pH 7.4, and 1 mM DTT. All protein was stored at a concentration of 10 mg/ml.

Enzymatic Activity Measurement and Kinetic Characterizations—Activity of the point mutants was assayed by monitoring the change in absorbance at 340 nm that accompanies reduction of NADH as reported previously (23). The R141C and E416D mutants exhibited saturable kinetics. Michaelis constants \( K_m \) and \( V_{max} \) for UDP-glucose and NAD was determined independently by holding [NAD \(^+\)] constant and varying [UDP-glucose] from 0 to 2 mM. Similarly, [NAD \(^+\)] measurements were made by holding [UDP-glucose] constant and varying [NAD \(^+\)] from 0 to 2 mM. Specific activities were calculated from absorbance values using the molar extinction coefficient for NADH of 6220 M\(^{-1}\) cm\(^{-1}\). Triplicate determinations for each concentration increment were plotted with PRISM (GraphPad Software, Inc., San Diego, CA). \( K_m \) and \( V_{max} \) were calculated by fitting the data to the Michaelis-Menten equation and assuming a single binding site/subunit for substrate and cofactor. Activity loss of wild-type and each mutant enzyme species was measured in aliquots removed at regular time points until no further activity was detectable. Each aliquot was assayed for 1 min with saturating cofactor and substrate. Specific activity was fitted to a semi-log plot following an initial lag period to calculate enzyme half-life. Statistical values were determined from these data by Student’s unpaired two-tailed \( t \) test.

Molecular Mass Determination—Oligomeric state of the apo enzyme was determined by size exclusion chromatography as previously described (23). Each sample was precleared by centrifugation prior to loading. To determine the effect of substrate and cofactor inclusion on oligomeric state, purified recombinant UGDH wild type and point mutants were individually loaded on a Superdex 200 HR 10/300 column (GE Healthcare) using a 250-μl loop and separated by FPLC in 1× PBS in the presence of 1 mM UDP-glucose and 5 mM NAD \(^+\) at a flow rate of 0.5 ml/min. Substrate and cofactor were included in each 250-μl load sample at the same final concentrations. Because of the interference from cofactor in UV absorbance readings from the column fractions of enzyme complexes, the protein content of these samples was measured at 595 nm upon the addition of Bradford reagent. The standards used were: thyroglobulin, 699 kDa; ferritin, 440 kDa; albumin, 67 kDa; \( \beta \) lactoglobulin, 35 kDa; and RNase A, 13.7 kDa. Resolution was sufficient to calculate molecular masses unambiguously in multiples of 57 kDa (the calculated molecular mass of monomeric His-tagged UGDH).

Protein Stability Determination—Purified recombinant UGDH wild type and point mutants were analyzed using a modified version of the Thermofluor stability assay as described by Ericsson et al. (34). Solutions of the wild-type and mutant apo enzymes (=5 μg of protein in all cases, except I331N, which required 45 μg) were prepared in 1× PBS containing Sypro Orange dye (Invitrogen; 1:500 dilution). Samples containing substrate and cofactor were prepared in the same manner with UDP-glucose, and NAD \(^+\) was added to a final concentration of 1 mM each. All samples were handled on ice throughout the preparation and transferred to an iCycler MyiQ Thermocycler (Bio-Rad) for analysis. The assay protocol was executed with the block initially cooled to 4 °C. After the addition of samples, the temperature was raised over a period of 76 min from 4 to 79 °C in 0.5 °C increments. The change in fluorescence was monitored for each sample at excitation and emission levels of 490 and 575 nm, respectively. Transitions representing the melting temperature, \( T_m \), were obtained from the derivative of fluorescence with respect to temperature for at least 8–12 replicates and plotted as the means ± S.D.

RESULTS

Embryonic Localization of UGDH in the Developing Mouse Heart—It was previously shown that ugdh is required for heart valve formation in zebrafish embryogenesis (11) and is critical for gastrulation in mice, which arrest at embryonic day 9.5 when its function is disrupted (15). We first examined Ugdh distribution in mouse embryos, specifically to address its localization in cardiac development. At embryonic day 11.5, strong expression of Ugdh was observed in the endocardial cushions located within the AV canal (Fig. 2A). Both the endothelial cells covering the cushions and the mesenchymal cells within the cushions stained positive for Ugdh expression. At embryonic day 12.5, Ugdh mRNA was present in the endocardial cushions of the AV canal, in the dorsal mesenchymal protrusion and in the outflow tract cushions (Fig. 2, B and C). At E14.5, specific expression of Lgdh was observed in the valve leaflets located in the AV canal (Fig. 2D), which eventually become the leaflets of the mitral and tricuspid valves. Ugdh expression was not detectable in myocardium, which was visualized by staining for cardiac troponin I (Tnni3) (Fig. 2, E–H).

Functional Examination of UGDH R141C and E416D Variants in Zebrafish—To examine the effects of the single nucleotide polymorphisms identified in the candidate screen of congenital heart disease sufferers, we used a transgenic zebrafish model in which the endogenous expression of Ugdh was knocked down by antisense MO injection. In addition to a pre-
vously published ugdh-ATG MO, we designed and characterized a new ugdh-Splice MO that efficiently affected splicing of the endogenous ugdh mRNA (supplemental Fig. S1). To assess the functional consequence of ugdh MO injection on Ugdh activity in the embryo, we stained the embryos for hyaluronan, which is a key molecular fate for the UDP-glucuronate product of UGDH. We observed reduced hyaluronan accumulation in multiple tissues, including the heart, in embryos injected with the ugdh MOs relative to uninjected embryos (Fig. 3, A and B).

Next, we used a transgenic zebrafish line in which a GFP reporter is expressed under an endothelial cell-specific promoter (Tg(Tie2:eGFP)). Use of this model facilitates the visualization of cardiac valve defects because GFP-positive endothelial cells normally cluster in a ring at the AV boundary beginning at 43 hpf when valve formation initiates. As expected from previously reported data (11), injection of MOs targeting ugdh in zebrafish embryos at the single cell stage eliminated endocardial ring formation (Fig. 3, C and D). We were able to restore endocardial ring formation by coinjecting the ugdh MOs with mRNA encoding wild-type human UGDH (Fig. 3E).

However, coinjection of UGDH R141C (Fig. 3F) or UGDH E416D (not shown) transcripts with the MOs failed to rescue the loss of endothelial clustering efficiently, suggesting that both polymorphisms conferred mutations to the UGDH enzyme that severely compromised its activity in vivo.

Embryos were further examined for cardiac morphogenesis as reflected in the presence of edema when the absence of endocardial ring formation failed to constrict the heart tube sufficiently for directional fluid flow. Cardiac edema was observed in >65% of embryos that had been injected with ugdh antisense MOs (Fig. 3G). Nearly all edema was eliminated by coinjection of wild-type human UGDH mRNA. However, consistent with the notion that functional defects are conferred by the UGDH R141C and E416D polymorphisms, coinjection of mRNA encoding either of these variants failed to rescue cardiac edema completely.

In Vitro Characterization of Purified UGDH R141C, E416D, and I331N (Jekyll)—To characterize the molecular defects in UGDH R141C and E416D variants more thoroughly and compare them with a previously reported defect, UGDH I331N, that prevented valve development in zebrafish, we generated a new ugdh-ATG MO, we designed and characterized a new ugdh-Splice MO that efficiently affected splicing of the endogenous ugdh mRNA (supplemental Fig. S1). To assess the functional consequence of ugdh MO injection on Ugdh activity in the embryo, we stained the embryos for hyaluronan, which is a key molecular fate for the UDP-glucuronate product of UGDH. We observed reduced hyaluronan accumulation in multiple tissues, including the heart, in embryos injected with the ugdh MOs relative to uninjected embryos (Fig. 3A and B).
The constructs encoding UGDH R141C and E416D mutants expressed abundant soluble recombinant protein in E. coli, and each enzyme form was purified to homogeneity by nickel affinity chromatography. Typical yields were similar for wild type and both of these mutants, suggesting that neither mutation intrinsically reduced gene expression or solubility. However, the UGDH I331N construct yielded significantly reduced quantities of purified protein that was highly unstable even at 4 °C. Substrate dependence of the enzyme reaction converting UDP-glucose to UDP-glucuronate was not saturable for this mutant, which had minimal detectable activity. However, kinetics for the other two mutants were saturable, and these measurements were used to calculate Michaelis-Menten constants (summarized in Table 1). The R141C mutant $V_{\text{max}}$ was only $\approx 40\%$ of the maximum value for wild-type UGDH, which may partially explain its human phenotype but exhibited slightly lower $K_m$ for both cofactor and substrate. In contrast, E416D had $\approx 40\%$ higher $V_{\text{max}}$ and moderately increased $K_m$ values relative to the wild-type enzyme.

**Half-life of Wild-type and Mutant UGDH**—The differences in activity of the recombinant mutant enzymes were relatively nominal considering the magnitude of the phenotype conferred *in vivo* by a heterozygous mixture of wild-type and mutant UGDH enzyme species. Thus, we postulated that the effects on intrinsic activity could be mediated either by different conditions than those in which we assayed (i.e., intracellular) and/or that the mutations influenced the stability of the functional enzyme. To test the latter point, we incubated UGDH wild type, R141C, and E416D individually at 37 °C and measured activity over time. The heterozygous condition was simulated in this experiment by pre-equilibrating a 1:1 mixture of wild-type and mutant protein as indicated in the legend were equilibrated overnight at 4 °C. The samples were then placed at 37 °C. At the indicated time points, activity was assayed by the addition of the enzyme to phosphate buffer containing saturation concentrations of NAD$^+$ and UDP-glucose. The values plotted are the means ± S.D. for triplicate determinations. For several points, the symbols obscure the error bars. Absolute turnover was measured as the change in NADH absorbance at 340 nm in a 1-min assay for comparisons of wild type with R141C (A) or E416D (B). Absorbance values were normalized to the value at time 0 to illustrate the rate of activity loss.

**FIGURE 4.** *Time-dependent activity of wild-type UGDH and point mutants.* Individual purified proteins or a 1:1 mixture of wild-type and mutant protein as indicated in the legend were equilibrated overnight at 4 °C. The samples were then placed at 37 °C. At the indicated time points, activity was assayed by the addition of the enzyme to phosphate buffer containing saturation concentrations of NAD$^+$ and UDP-glucose. The values plotted are the means ± S.D. for triplicate determinations. For several points, the symbols obscure the error bars. Absolute turnover was measured as the change in NADH absorbance at 340 nm in a 1-min assay for comparisons of wild type with R141C (A) or E416D (B). Absorbance values were normalized to the value at time 0 to illustrate the rate of activity loss.

**Structural Integrity and Intrinsic Stability of Mutant UGDH**—Our lab and others have previously shown that mammalian UGDH is active as a hexamer (23, 25, 35), and several of our previously characterized active site mutants show dissociation to an exclusively dimeric form with dramatic reduction in $V_{\text{max}}$ or increased $K_m$. Therefore, we compared the quaternary structure of the R141C, E416D, and I331N mutants with that of wild-type UGDH by gel filtration chromatography. The wild-type protein eluted with peaks at molecular masses of $\approx 360$ and $\approx 120$ kDa, corresponding to a hexamer and a dimer, respectively, of the $\approx 57$-kDa UGDH monomers (Fig. 5A, *solid black trace*), in which the hexamer was clearly the dominant peak. The elution profiles of both the E416D (Fig. 5B) and R141C (Fig. 5C) mutant enzymes were split in a hexamer-dimer equilibrium. In the case of E416D, the dimeric species accounted for $\approx 40\%$ of the absorbance ($\approx 60\%$ dimer), whereas R141C appeared to be $\approx 80\%$ dimeric. The I331N mutant was almost exclusively dimeric, with $\approx 10\%$ of the absorbance attributable...
to a peak potentially corresponding to monomeric enzyme. Consistent with its previously noted instability, the total magnitude of the eluted peaks for I331N UGDH was reduced 10-fold relative to the wild type and other mutants (Fig. 5D).

Interestingly, including saturating levels of NAD⁺ and UDP-glucose in the loaded protein and the column running buffer induced a complete shift of the wild-type UGDH elution profile to that of the hexamer (Fig. 5A, dotted gray trace). The same phenomenon was observed for the E416D mutant (Fig. 5B) and, to a lesser extent, the R141C mutant (Fig. 5C). Such a shift did not occur appreciably upon substrate and cofactor addition to I331N. Thus, it appeared that the formation of a holoenzyme ternary complex stabilized the hexameric quaternary assembly of the enzyme, and the extent of hexameric association was somewhat related to the level of stability of the wild type and each mutant as observed in the activity assays (Fig. 4).

To examine the intrinsic stability of UGDH and its point mutants in the presence and absence of its substrate and cofactor, we used a thermally sensitive fluorescence assay. Consistent with a previous report using this method (34), the observed melting temperature \( T_m \) of the wild-type apoenzyme form of UGDH was \( \sim 50 \) °C for the apoenzyme form and increased to \( \sim 65 \) °C when NAD⁺ and UDP-glucose were added to the reaction mixture (Table 2 and Fig. 6). Similar results were obtained for the E416D mutant. Although an increase in \( T_m \) was observed for both of the other mutants as well, the apoenzyme form of the R141C mutant was denatured with a \( T_m \) of \( \sim 8 \) °C lower than that of wild type, and the increase in \( T_m \) for I331N was only \( \sim 46 \) °C for the ternary complex. These results are consistent with greater overall thermal stability of the hexameric form of UGDH and reflect the differences observed in the stability of the respective mutants.

### DISCUSSION

UGDH is an enzyme that has been cloned and characterized from many diverse species and implicated in zebrafish heart development for over 10 years. To our knowledge, this is the first report of a characterization of mutations to UGDH occurring in a population of human patients with congenital heart valve defects. In this study, we localized expression of UGDH to a critical region of the developing mouse heart and followed its timed increase and subsequent decrease over the normal time frame for cardiac valve formation. We further demonstrated that the patient mutations R141C and E416D cannot rescue valve development or cardiac edema resulting from UGDH deletion in the zebrafish model. A detailed characterization of the structural and functional features of the mutant enzymes revealed modest alterations in the intrinsic activity of the enzyme that may translate to reduced function at cellular levels of substrate and cofactor. However, the most dramatic impact of the mutations on enzyme function appeared to derive from the striking destabilization of intrinsic thermal stability that leads to accelerated loss of activity in physiological conditions.

**FIGURE 5.** Hexameric assembly of wild-type and mutant UGDH is sensitive to the presence of substrate and cofactor. Purified wild-type and mutant UGDH was fractionated by gel filtration FPLC in the absence or presence of NAD⁺ (5 mM) and UDP-glucose (1 mM). Absorbance was monitored at 280 and 595 nm, respectively. Size determination of each peak was made by comparison with molecular weight standards. Representative traces are plotted for wild-type UGDH (A), E416D (B), R141C (C), and I331N (D). On each plot, the apo-enzyme trace (in black) is superimposed on its holo-enzyme complement (in gray) for comparison of quaternary structure.

| Enzyme     | Apo       | Ternary  |
|------------|-----------|----------|
| Wild-type  | 49.5 ± 0.3| 64.8 ± 0.2|
| E416D      | 50.7 ± 0.2| 63.5 ± 0.2|
| I331N      | 40.8 ± 2.7a| 46.0 ± 1.4ab|
| R141C      | 46.2 ± 1.0| 55.0 ± 1.0b|

a \( p < 0.001 \) relative to wild-type apoenzyme.
b \( p < 0.001 \) relative to wild-type holoenzyme.
Importantly, we demonstrated that the presence of the mutant forms in a 1:1 ratio with wild-type UGDH, representative of the heterozygous genotype found in all patients with the polymorphisms, promotes a loss of wild-type UGDH activity as well.

UGDH was first identified as a cardiac valve initiation factor in zebrafish by isolating the gene disruption that was present in the zebrafish valve mutant jekyll (11). The lesion was found to be a single nucleotide substitution from A to T that generated an ATC to AAC codon change. The amino acid change, published as isoleucine to aspartate, is actually an isoleucine to asparagine mutation at a position that is highly conserved across numerous species including humans. The substitution had not been previously characterized in the context of the enzyme structure. We found the I331N mutant to be highly unstable and poorly expressed, with further impairment in quaternary structure assembly and virtually undetectable residual activity. These properties are a direct consequence of perturbations to the local environment of the Ile-331 residue by the substitution (Fig. 7A). This residue is located in a strongly hydrophobic core motif and is well packed among several other nonpolar amino acid side chains. Significant disruption to this stable, well conserved motif would be expected by the introduction of a hydrophilic amino acid side chain in any one of these positions, so this accounts for the strong effect of the asparagine substitution on UGDH stability.

Examination of Glu-416 in the context of the UGDH ternary complex structure (Fig. 7B) illustrates its role in indirect positioning of the UDP-glucose substrate via two water molecules in collaboration with the side chain of lysine 339. The backbone of the loop containing Lys-339 is stabilized by hydrogen bonding to Glu-416. Thus, effects of the E416D substitution were predicted to include decreased binding of UDP-glucose. In fact, the $K_m$ for UDP-glucose is increased by $\approx 50\%$ (Table 1). This relatively modest reduction in apparent affinity for substrate in the context of enzymatic turnover is also accompanied by a significant acceleration in the loss of activity at 37 °C (Fig. 4) and destabilization of the hexameric quaternary assembly in the absence of substrate and cofactor (Fig. 5), although the intrinsic thermal stability is not compromised (Fig. 6). We previously found that the K339A substitution increased $K_m$ for UDP-glucose by $\approx 150\fold$, and yielded an exclusively dimeric apoenzyme. The more direct role in UDP-glucose coordination through interaction with both phosphate groups accounts for the significant difference in $K_m$ between the K339A and E416D mutants. The observed effect on the hexamer-dimer equilibrium is consistent with a link between active site integrity, substrate and cofactor binding capacity, and hexameric assembly, which ultimately determines enzyme stability. Reduced stabil-
ity of the E416D hexamer results in propagation of instability to the wild-type enzyme, as observed in a simulated heterozygous condition (Fig. 4), so this is a likely explanation for the effects of this mutation on cardiac valve morphogenesis, as discussed below.

Comparison of results from the R141C mutant is consistent with this explanation. Arg-141 is located at the interface between dimeric units of the hexamer (Fig. 1) and maintains a hydrogen bond network that stabilizes a helix-loop-β motif critical to the dimer-dimer association (Fig. 7C). Substitution of this residue with a cysteine eliminates these contacts and releases the helix from its tight association, which would be expected to disrupt the quaternary structure to an exclusively dimeric form as was shown in Fig. 5. The reduced stability of the dimer is associated with a reduction in enzymatic activity (Table 1), destabilization of the mutant in both a homozygous and a heterozygous reaction mixture (Fig. 4), and significantly reduced thermal stability in the absence or presence of ternary complex components (Fig. 6). The reductions in activity and stability are probable underlying causes of the phenotype associated with this missense mutation in human patients.

The relatively modest differences in enzymatic activity of the R141C and E416D mutants is not entirely surprising because bacterial homologues are only found as dimers (24). However, the inability to form normal complexes may nonetheless have dramatic impact on the stability of wild-type hexahexamers with either R141C or E416D and/or on the potential for allosteric regulation within a cellular context. This is particularly intriguing because the addition of substrate and cofactor at saturating levels can fully restore the hexameric structure. Although hexameric structure is not critical for achieving wild-type maximum reaction velocity if optimal conditions are provided (29), studies of the nucleotide sugar dehydrogenase family member GDP-mannose dehydrogenase (36) suggest that hexameric structure provides the capacity for allosteric regulation and cooperativity with respect to substrate use. Thus, the effects on quaternary assembly, particularly the ability to restore and maintain a stable ternary complex within a physiological range of available substrate and cofactor, could translate to significant differences in the net enzymatic activity in developmentally critical phases.

UGDH is the only enzyme that provides UDP-glucuronate, which is an essential precursor for hyaluronan synthesis, initiation of all proteoglycan synthesis, and elongation of the glycosaminoglycan chains that comprise proteoglycans. In mice, UGDH gene disruption is embryonically lethal (15), and in zebrafish Ugdh inactivation results in cardiac valve defects (11). UGDH and HAS2 gene disruption have identical phenotypes, and cardiac tissue explant studies, in which AV septum formation can be restored by exogenous hyaluronan, the product of the HAS2 gene, demonstrate that UGDH must function at a high level during heart formation to supply precursors for hyaluronan production (9). It has been further shown in the Xenopus model system that UGDH expression directly dictates the production capacity for hyaluronan during development (12). Our results show that UGDH is expressed at a very high level specifically in the region of the developing valves throughout valve development (Fig. 2). The introduction of transcripts encoding either the R141C or E416D mutant form of UGDH was insufficient to support valve formation or prevent cardiac edema (Fig. 3), which clearly illustrates a significant level of impairment resulting from these missense mutations in vivo.

Based on the infrequency with which UGDH mutations have been reported, it is likely that most UGDH polymorphisms do not yield viable embryos. In support of the extreme phenotypic distress conferred by such lesions, only the heterozygous and not the homozygous expression of R141C and E416D is observed in the population study (30). The characterizations we described above point toward a significant loss of enzyme stability and resultant reduction in UDP-glucuronate production below the levels that are required to sustain hyaluronan synthesis while the cardiac cushions are forming. Cardiac cushions are largely the result of rapid hyaluronan extrusion and accumulation between the layers of the myocardium and endocardium beginning at embryonic day 9.5. Their formation produces tissue protrusions that are invaded by endocardial cells, which delaminate because of epithelial to mesenchymal transition. Their absence prevents epithelial to mesenchymal transition for cell migration into the AV septum, which is needed for subsequent valve differentiation. Because this process necessitates rapid deposition of large quantities of hyaluronan, UGDH function is critical at this time, and even modest reductions in its function are clearly disruptive. Screening families for these alleles may assist in the prediction of cardiac valve defects and offer therapeutic targets.

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