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RESEARCH ARTICLE

Age-Dependent Changes in Geometry, Tissue Composition and Mechanical Properties of Fetal to Adult Cryopreserved Human Heart Valves

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Abstract

There is limited information about age-specific structural and functional properties of human heart valves, while this information is key to the development and evaluation of living valve replacements for pediatric and adolescent patients. Here, we present an extended data set of structure-function properties of cryopreserved human pulmonary and aortic heart valves, providing age-specific information for living valve replacements. Tissue composition, morphology, mechanical properties, and maturation of leaflets from 16 pairs of structurally unaffected aortic and pulmonary valves of human donors (fetal-53 years) were analyzed. Interestingly, no major differences were observed between the aortic and pulmonary valves. Valve annulus and leaflet dimensions increase throughout life. The typical three-layered leaflet structure is present before birth, but becomes more distinct with age. After birth, cell numbers decrease rapidly, while remaining cells obtain a quiescent phenotype and reside in the ventricularis and spongiosa. With age and maturation—more pronounced in aortic valves—the matrix shows an increasing amount of collagen and collagen cross-links and a reduction in glycosaminoglycans. These matrix changes correlate with increasing leaflet stiffness with age. Our data provide a new and comprehensive overview of the changes of structure-function properties of fetal to adult human semilunar heart valves that can be used to evaluate and optimize future therapies, such as tissue engineering of heart valves. Changing hemodynamic conditions with age can explain initial changes in matrix composition and consequent mechanical properties, but cannot explain the ongoing changes in valve dimensions and matrix composition at older age.
Introduction

End stage valvular disease is commonly treated with heart valve replacement to alleviate cardiac, pulmonary, or systemic problems due to the disease. Semilunar valves, in particular the aortic valve, are most often replaced. Although current heart valve alternatives enhance survival and quality-of-life of most patients, they have several limitations. The most important drawback is that they do not consist of living tissue and, therefore, do not grow, repair, and remodel after implantation. Especially for pediatric and adolescent patients, who require multiple valve replacements, this is a significant problem. In this regard replacement of the diseased aortic valve by the autologous pulmonary valve autograft (Ross procedure), common for pediatric and adolescent patients, is considered clinically effective [1]. Nevertheless, redo surgery for autograft failure in second postoperative decade is not uncommon and the replacement pulmonary valve is at risk for an additional operation or intervention as well.

Heart valve tissue engineering (HVTE) seeks to overcome the current limitations of valve prostheses, allografts and autografts, by creating a living heart valve replacement that can grow and adapt in response to changing functional demands. Key design parameters for HVTE are: valve geometry and morphology, cell type, extracellular matrix (ECM) composition and architecture, tissue mechanical properties, and growth and remodeling potential, which may all change with the target age group for valve replacement. As the properties of native human heart valves represent the ideal blueprint for HVTE [2], data from human valves of different ages should be used for optimizing HVTE for different target groups.

So far, valvulogenesis and tissue morphogenesis of heart valves have been mainly studied using valves of animal origin (e.g. porcine) [2–7]. Studies with human valves mostly concentrate on fetal or adult valves [8–14], while studies on human pediatric and adolescent valves are only sparsely available [10,15,16]. Hence, the development of structure-function properties from young to old age is largely unknown. In addition, there is no consensus on potential differences in structure-function properties between human pulmonary and aortic valves.

Here, we provide a comprehensive data set on aortic and pulmonary valve properties at different stages of growth and development (fetal, child, adolescent, and adult) that can be used for the development, application, and evaluation of living, age-specific heart valve replacements. The development of structure-function properties and valve remodeling was assessed from quantitative (dimensions, biochemical assays, mechanical testing) and qualitative (histology) measures of the geometry, morphology, composition, and mechanical properties of sixteen pairs of structurally unaffected human aortic and pulmonary heart valves.

Methods

Tissue preparation

Sixteen sets of cryopreserved healthy human aortic and pulmonary valves from the same donor (0–53 year, n = 32 valves) were obtained with written informed consent from Dutch postmortem donors. For the pediatric valves, the written informed consent was obtained from the next of kin or legal guardians. All postnatal valves were assessed to be unfit for implantation and were obtained from, and after approval for the current study by, the Heart Valve Bank Rotterdam (Erasmus University Medical Center, Rotterdam, The Netherlands). The fetal valves (1 set; 21 weeks of gestation) were obtained after elective abortion following individual written informed consent, after approval of the current study by the medical ethical committee of the Leiden University Medical Center (MEC-P08.087). The investigation of the fetal as well as the postnatal valves was conform the principles outlined in the Declaration of Helsinki. All valves were structurally and mechanically unaffected as assessed by macroscopic observations at time
of cryopreservation at the Heart Valve Bank and after thawing by the researchers. The cause of death of the donors was not related to valvular disease or conditions known to precede valvular disease. Earlier studies demonstrated that the applied cryopreservation and thawing protocol applied to these valves did not affect the structural integrity of collagen and elastin [17] and mechanical properties [18]. The fetal valves were only studied with histology due to the size of these valves.

The cryopreserved valves were stored at -80°C and thawed just prior to ECM analysis and mechanical testing. The thawing protocol was similar for all studies heart valves. The thawed valves were cut according to a cutting scheme in preparation for the pre-defined analyses (Fig 1 and S1 Table). Samples for histology and biochemical assays were respectively fixed in formalin or snap-frozen within 24 hours, while mechanical testing was performed within 48 hours after thawing.

Valve dimensions and morphology

Valve dimensions were quantified by defining the annulus diameter, leaflet size (both radial and circumferential), and leaflet thickness. The annulus diameter and gross morphological appearance (e.g., presence of fibrosis or artheroma) of the valve leaflets were assessed upon explantation by the Heart Valve Bank Rotterdam. The size of the right (coronary) leaflet was measured after excision in circumferential and radial direction to assess the dimensions of the leaflets (S1B Fig). The thickness of the leaflets was measured as part of the indentation tests (see below).
Cell phenotype, tissue composition and maturation

Pieces of the leaflets (leaflet, hinge region and part of the wall; Fig 1D) were fixed overnight in formalin, processed and subsequently embedded in paraffin. They were sectioned at 10 μm thickness to qualitatively study phenotype of the valvular interstitial cells and matrix composition using histology (H&E: tissue morphology, Masson Trichrome: collagen, Verhoeff-van Gieson: collagen and elastin, Safranin-O: proteoglycans) and immunofluorescent stainings (elastin, collagen type I and III, and alpha smooth muscle actin (αSMA)). The leaflet parts for biochemical assays (2–4 tests per valve) were lyophilized and digested in papain solution (100 mM phosphate buffer [pH = 6.5], 5 mM L-cystein, 5 mM EDTA, and 125–140 μg papain/ml) to determine total cell number (DNA content, μg/mg dry weight), matrix composition (sulfated glycosaminoglycans (sGAG) and hydroxyproline content, μg/mg dry weight), and collagen maturation (# cross-links per collagen triple helix). For details on the qualitative and quantitative measures, see S1 Appendix.

Indentation tests

To characterize local tissue mechanical properties, indentation tests were performed on all sets (n = 30 valves) in the belly region (~7 indentations per leaflet) and near the commissures (~4 indentations per leaflet) of hydrated right (coronary) leaflets (Fig 1C). For indentation tests, the fibrosa-side of the leaflet was placed face-down on a glass (coverslip) and the tests were performed at room temperature as described previously [19]. Briefly, a spherical sapphire indenter (diameter 2 mm) was used to compress the tissue with a constant indentation speed of 0.01 mm/s. At the indented locations, one preconditioning cycle followed by two indentation tests were performed to determine leaflet stiffness (E-modulus), which was calculated from the force-indentation curve up to 20% indentation. When the indenter touches the leaflet, a drop in the force signal was observed. The height of the indenter with respect to the coverslip at that moment was taken as the thickness of the indented sample.

Biaxial tensile tests

To further analyze mechanical properties of the valves, biaxial tensile tests were performed on 1 to 3 sets of valve leaflets per age group, except for the fetal valves (n = 12 valves in total, see S1 Table) as previously described [20]. In brief, rectangular samples were cut (see Fig 1), kept hydrated and placed on aluminum foil to mount the samples in a BioTester 5000 device (CellScale, Waterloo, Canada) using a 5N load cell and a BioRakes mounting system with 0.7 mm space between the pins. The samples were then tested while submerged in PBS at room temperature. The samples were biaxially stretched to peak values (failure) in circumferential (ε_c) and radial (ε_r) leaflet direction at a strain rate of the initial sample length per minute and following a step-wise protocol. A first series of strains was applied, according to ε_c: ε_r (% strain) = 0:60; 11.5:55; 23: 11.5: 25:0 to capture the non-linear mechanical behavior of the leaflets. The samples were unloaded and left to recover for 1 minute in between steps. Next, the strains were sequentially increased during up to 5 additional series in steps of 5% or 10% until failure. Before each series, the samples were preconditioned for 10 cycles first to the max ε_c and then to the max ε_r of the group.

The final stress-strain curves, where stresses were normalized for tissue thickness (Cauchy stress), were averaged per tested valve (max. 6 samples per valve, see Fig 1) and plotted for both radial and circumferential direction. The average curves stop when the first sample failed.

In order to achieve easily interpretable figures for mechanical properties and to account for inter-subject variation in non-linear stress-strain behavior, we defined stiffness values (E-modulus) from the tangent of the fitted slope of the linear strain-hardening region of the stress-
strain curves and tissue extensibility from the x-intercept of the slope of the strain-hardening part of the stress-strain curves. These values were averaged per tested valve using the individual stress-strain curves that reached the strain-hardening phase.

Data analyses and mapping hemodynamic function
Data are presented as mean ± the standard error of the mean. Postnatal valves were subdivided into three age groups based on the age of the available donor valves (S1 Table): child (2 months to 11 years; n = 5), adolescent (18–22 years; n = 4), and adult (38–53 years; n = 6). Fetal valves were not included in statistical analyses. Relationships between valve properties or between valve properties and age were identified with correlation analysis (with r the correlation coefficient). Statistical differences (p<0.05) between age groups and the aortic and pulmonary valves were tested with two-way ANOVA, followed by Bonferroni post-hoc testing. GraphPad Prism software (GraphPad Software, Inc, USA) was used for the analyses.

In an attempt to correlate valve geometry and tissue composition with changing postnatal hemodynamic conditions, we collected hemodynamic data from several databases, including The National Heart Lung and Blood Institute, MedScape and literature [21–23].

Results
Dimensions of human heart valve leaflets continue to increase with age
The annulus diameter of the heart valves increased with age (p<0.001; Fig 2A), with the pulmonary annulus slightly larger than the aortic annulus. As expected, annulus size increased rapidly in the pediatric group and slowly thereafter, with continued, but slow increase up to old age. Concomitantly, leaflet size increased in circumferential and radial direction with age, and this increase was similar for both valves (p<0.001; Fig 2C and 2D).

The thickness of the aortic valve leaflets was heterogeneous (Fig 2B). Overall, the aortic valve leaflets were thicker than the pulmonary valve leaflets. This difference was predominant in the belly of adolescent and adult valves (p<0.001). The commissural regions of the aortic leaflets were thinner than the belly region, but only significant in the adolescent and adult valves (p<0.001).

In the pediatric valves, fibrosis was not observed at all. In the valves from the adolescent group mild fibrosis was seen, though not in all valves. However, adult valves all showed some degree of fibrosis. Fibrosis was mostly found in the central belly region, and incidentally at the fixed edge of the leaflet. Some adult valves showed atheroma spots at the fixed edge of the leaflet.

Three-layered structure is present before birth, but becomes more distinct with age
The typical three-layered leaflet structure was clearly visible in all studied valves, even in the fetal valves (Fig 3), but became more pronounced with age. Collagen was predominantly found in the fibrosa, as demonstrated by the Masson Trichrome (not shown) and Verhoeff-Van Gieson staining (Fig 2A, 2C–2F and 2H–2J). Here, collagen was mainly characterized as type I (Fig 3K–3N), whereas collagen type III was present throughout the leaflet (data not shown). Elastin was observed in the ventricularis (Fig 3A–3J). In the fetal valve, elastin was clearly observed with the elastin immunofluorescent staining, but not with the Verhoeff-Van Gieson staining (Figs 3B versus 3A and 2G vs. 2F). Proteoglycans were observed in the spongiosa of the leaflets and in the hinge region of the valves (Fig 3O–3R). Since the differences between the aortic and...
Fig 2. Evolution of valve dimensions. (A) Increase in annulus diameter in the aortic (grey) and pulmonary (black) valve. The diameter increases rapidly early in life and slowly, but continuously, thereafter. The pulmonary valve is slightly larger compared to the aortic valve. (B) The thickness of the leaflets is heterogeneous and larger in the belly compared to the commissures, which is similar for all age groups. Aortic valves are slightly thicker than pulmonary.
pulmonary valves were minimal, the aortic valve was chosen to represent the observed findings (S1 Fig for the histology of the pulmonary valve).

**Cellular content and phenotype change rapidly after birth**

Histological analyses (Fig 3) suggest that cell content decreases with age. Total cell number (DNA content) was indeed higher in the pediatric aortic leaflets compared to the adolescent and adult aortic leaflets (p < 0.01; Fig 4A). Especially, the leaflets from the youngest donors contained more DNA than the older leaflets. No differences in cell number were found between the aortic and pulmonary valve. Interestingly, in the fetal and pediatric leaflets, cells were observed throughout all layers of the leaflet, while the cells in the adolescent and adult leaflets were mainly located in the spongiosa and ventricularis.

αSMA-positive cells were frequently observed in the leaflets of the fetal and 8-month old donor, but were only sparsely observed in the older valve leaflets (≥ 4 years), indicating a more quiescent cell phenotype (Fig 3S–3W). When present, αSMA-positive cells were observed in the hinge region and the arterial wall for the fetal and pediatric valves, and predominantly in the arterial wall for the adult valves.

**Matrix composition shifts toward decreased sGAG, increased collagen content and maturity with age**

The sGAG content decreased with age in both the aortic (p < 0.05) and pulmonary valves (p < 0.01; Fig 4B). The decrease in sGAG with age was most obvious for the aortic valve, especially from pediatric to adult aortic valves (p < 0.01). In addition to a decrease in sGAGs, the hydroxyproline content, as a measure of valvular collagen, increased with age in the aortic valve (p < 0.01; Fig 4C). The hydroxyproline-to-sGAG ratio increased with age for both valves (p < 0.05; data not shown). Interestingly, pediatric aortic valves contained more sGAG relative to hydroxyproline, while in the adult aortic valves more hydroxyproline compared to sGAG was present.

The collagen matrix of both aortic and pulmonary valves matured with age, as indicated by an increase in cross-link density (Fig 5). In the aortic valve, HP collagen cross-link density increased slightly throughout life (p < 0.01), whereas in the pulmonary valve HP cross-link density only increased from adolescent to adult age (p < 0.05; Fig 5A). LP cross-link density increased with age in both aortic and pulmonary valves (p < 0.001; Fig 5B). HP cross-link density was always higher than LP cross-link density, but the HP-to-LP ratio decreased with age in both valves (p < 0.05), predominantly from childhood to adolescence (p < 0.05; Fig 5C).

**Leaflet stiffness and extensibility change with age**

In the aortic valve leaflets, the low-strain stiffness of the belly (p < 0.01), as well as the commissure (p < 0.05) increased with age (Fig 6A and 6B). This increase was striking from adolescence (belly: 3.5 kPa; commissure: 7.0 kPa) to adulthood (belly: 10.5 kPa; commissure: 20.8 kPa). Biaxial tensile testing showed that aortic leaflet stiffness increased both in circumferential and radial direction (Fig 7A and 7B; p < 0.05). In the pulmonary valve leaflets, the low-strain stiffness increased with age only in the commissures (Fig 6B; p < 0.01), although the belly showed a 2-fold increase (p < 0.05) in the low-strain stiffness from adolescence (4.5 kPa) to adulthood.
| Fetal | Child | Adolescent | Adult |
|-------|-------|------------|-------|
| ![Image](image1) | ![Image](image2) | ![Image](image3) | ![Image](image4) |
| ![Image](image5) | ![Image](image6) | ![Image](image7) | ![Image](image8) |

**Fig 3.** Representative histological and immunofluorescent stainings on aortic valves. Figures A-E represent the whole aortic valve (leaflet, hinge region, and wall, while in E-W a representative part of the leaflet is shown. (A, C-F, H-J) Verhoeff-Van Gieson staining for collagen (red) and elastin (black). (B, G) Elastin was observed in the fetal valve using immunofluorescence (red). (K-N) Collagen type I (red) was predominant in the fibrosa. (O-R) Safranin-O
staining showed proteoglycan presence (red/orange) mainly in the spongiosa and the hinge region. (S-W) αSMA (immunofluorescence; green) with cell nuclei (in blue). In the leaflets of the fetal (S) and 8-month old donor (T) αSMA-positive cells were observed, while in the older leaflets almost no αSMA-positive cells were observed (U-W). Scale bar: 500 μm. l: leaflet; w: wall; f: fibrosa; s: spongiosa; v: ventricularis.

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(10.3 kPa). Here, no age-related differences in circumferential and radial stiffness were observed. In both valves and at all ages, the stiffness in circumferential direction was higher than in radial direction, indicating anisotropic behavior.

The leaflets of the pediatric (11 year) and adolescent (20 year) valves were more extensible than the adult leaflets (Fig 7C and 7D) and also more extensible on the pulmonary side than on the aortic side. On the other hand, leaflet extensibility of the 8-month old valve was relatively low and comparable to the extensibility of the adult leaflets for both the aortic and pulmonary valve. Although this may reflect the natural variability amongst the donor valves, the 8-month valve could also be considered as an outlier in the absence of more valves of comparable age. Linear regression analysis excluding the data from the 8-month old valve revealed that leaflet extensibility in circumferential direction decreased significantly with age for both the aortic and pulmonary valves (p < 0.05, lines in 7E). For the pulmonary valve, also the extensibility in radial direction decreased significantly with age (p < 0.01). The E-moduli shows no correlation with age when excluding the data from the 8-month old valve (lines in 7F).

Leaflet mechanical properties are related to matrix composition and maturation

In the aortic valve, the belly stiffness positively correlated with collagen content (p<0.05; r = 0.62). Thus, tissue stiffness increases with increasing collagen content. Furthermore, the stiffness in circumferential direction positively correlated with HP cross-link density (p<0.05; r = 0.9).

In the pulmonary valve, a negative correlation was observed between sGAG content and commissure stiffness (p<0.05; r = -0.56), meaning that the stiffness is decreasing with increasing sGAG content. In addition, positive correlations between HP and LP cross-link densities and belly stiffness were observed for this valve (p<0.01; r = 0.71 and p<0.05; r = 0.58 for the HP and LP cross-links, respectively).

Reported valvular hemodynamics do not correlate with observed valve geometry and tissue properties in adults

In an attempt to correlate valve composition, geometry, and mechanical properties to valve hemodynamic loading, age-specific results were compared with average data on valvular hemodynamics reported in literature (heart rate, pressures). While changes in matrix composition and valve dimensions observed in the young age groups seem to coincide with reported changes in hemodynamic conditions, ongoing changes in dimensions and matrix composition later in life are difficult to relate to the more stable average hemodynamic conditions and may therefore also reflect the consequences of ageing. For instance, gradual increases in collagen content and stiffness, as well as a decrease in sGAG content, are observed for both the aortic and pulmonary valves during adult age, while hemodynamic loading conditions can be considered more stable (Fig 8).

Discussion

To determine structure-function properties, valve remodeling and differences thereof between pulmonary and aortic heart valves with age, we obtained data on valve dimensions,
Changes in DNA-, sGAG-, and hydroxyproline (HYP) content.

Changes in DNA- (A), sGAG- (B), and hydroxyproline (HYP) content (C) in μg/mg dry weight (left side: all data points used for correlation analysis with age; right side: grouped data). DNA content is higher in children, especially in the first years of life (0–4 years), compared to the adolescents and adults. sGAG content decreases with age. Hydroxyproline content increases with age in the aortic valve. Differences between groups (p<0.05) are indicated by paired symbols. AV: aortic valve, PV: pulmonary valve.

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Fig 5. Changes in collagen HP, LP cross-links, and the HP-to-LP ratio. Changes in collagen HP (A), LP cross-links (B), and the HP-to-LP ratio (C) in μg/ mg dry weight (left side: all data points used for correlation analysis with age; right side: grouped data). HP cross-links density increases with age in the aortic valve. In the pulmonary valve, the HP cross-link density increases from adolescent to adult age. LP cross-link density increases with age in both valves. Only
composition, maturation and mechanical properties of pairs of human aortic and pulmonary cryopreserved (donor) valve leaflets across a broad range of age. All studied leaflets, including the fetal leaflets, demonstrated a three-layered structure, with mainly collagen type I in the fibrosa, sGAG in the spongiosa, and elastin in the ventricularis. Aikawa et al. described that the three-layered structure, with elastin in the ventricularis, becomes apparent at 36 weeks of gestation [16], whereas our data demonstrated the presence of elastin already by week 21. This

Fig 6. Changes in stiffness at low strains as measured with indentation tests. The stiffness was computed in the belly (A) and commissures (B) (left side: all data points used for correlation analysis with age; at the right side). In the aortic valve, the stiffness in the belly and commissures increase with age, whereas in the pulmonary valve only the commissural stiffness increases with age. In both valves, E-moduli show a steep increase from adolescence to adulthood. Significant differences between groups (p<0.05) are indicated by paired symbols. AV: aortic valve, PV: pulmonary valve.

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Fig 7. Averaged stress-strain curves, E-modulus at high strains, and extensibility obtained from biaxial tensile tests. Averaged stress-strain curves (A, B), E-modulus at high strains (C), and extensibility (D) obtained from biaxial tensile tests. The E-moduli (E) in both circumferential and radial direction increase with age in the aortic valve (p<0.05). In the pulmonary valve the E-modulus does not change with age. The leaflets of the 11 and 20-year-old donors.
discrepancy might be caused by differences in staining techniques, as Votteler et al. demonstrated tropoelastin/elastin expression in the semilunar valve leaflets as early as week 7 of gestation and concluded that Verhoeff’s containing ECM-visualizing stains fail to identify developing, immature elastic fibers [24]. This corresponds with our findings that elastin was hardly observed in the fetal valves with the Verhoeff-Van Gieson staining, but clearly observed with the immunofluorescent staining.

With respect to valve dimensions, we found that the annulus increases rapidly during the first years of life, and slowly thereafter, even at older age. This can be due to growth or to a decrease in compliance with older age, causing dilatation of the annulus, as suggested by Merryman [25]. Annulus dilation may hinder movement and/or proper coaptation of the leaflets, thereby promoting regurgitation [26]. Interestingly, however, we also observed an increase in leaflet size with age. This may indicate an active remodeling or growth process to ensure proper valve closure. The thickness of the leaflets did not change with age. However, in the adolescent and adult age groups, the belly was thicker than the commissures. While this can be explained from the required load bearing capacity of the belly region [27,28], it might also be partly related to the presence of fibrotic spots, which were observed in the fibrosa of all adult and some adolescent leaflets.

From fetal to adolescent and adult age, the number of valvular interstitial cells decreased, which might be attributed to an increased cell proliferation-to-apoptosis ratio in young valves [16]. At older age, the number of cells remained constant and the cells mainly homed in the ventricularis and spongiosa. After birth, the cellular phenotype is assumed to be regulated by environmental cues [26], such as local tissue strains due to hemodynamic loading. Similar to the findings of Aikawa et al., the cells in the leaflets of the fetal and 2- and 8-month old donor were αSMA-positive, suggesting an activated myofibroblast phenotype relevant for fast matrix synthesis and remodeling [29,30]. Interestingly, no differences in αSMA-positive cells between the aortic and pulmonary valve were observed, suggesting that both valves have comparable remodeling capacity at young age. In the older aortic leaflets only quiescent cells (no αSMA) were observed, but mainly in the spongiosa and ventricularis and not in the fibrosa. This is surprising, since the fibrosa of all older valves showed evidence of fibrotic areas, suggesting ongoing matrix remodeling. Possibly this remodeling is so slow or passive due to ongoing collagen cross-linking, that it is not reflected by VIC presence or phenotype. Alternatively, valvular endothelial cells on the fibrosa side may contribute to the onset of fibrosis via shear stress-induced increase in inflammatory receptors [31] and decrease in fibrosis inhibitors [32]. In the absence of VICs, these negative effects cannot be suppressed [33] and will continue to contribute to valve degeneration.

Our mechanical characterization indicated anisotropic mechanical behavior in all studied age groups, similar to other studies using porcine or human valves from single age groups [6,13,15,34–37]. In addition to this, the adult valve leaflets were less extensible than the pediatric and adolescent leaflets. This was previously reported for porcine [6] and human [15] aortic leaflets and our study adds the pulmonary valve leaflets to that list. Leaflet stiffness was shown to increase with both increasing collagen content and decreasing sGAG content. Merryman hypothesized that ECM stiffness increases with age [25], which might be explained from our observation of the shift towards more collagen compared to sGAG in the ageing heart valve.
Fig 8. Valve hemodynamics during postnatal life in the aortic and pulmonary valve. Trend lines for reported data on valve hemodynamics (black) and our measured data on annulus diameter (grey), matrix composition, and stiffness of the belly (red) during postnatal life in the aortic (A) and pulmonary valve (B). Left-sided pressures increase during childhood, while the right-sided pressures decrease to adult values rapidly after birth. Hemodynamic data are collected from several databases, including The National Heart Lung and Blood Institute, MedScape and literature. Trend lines for our experimental data were obtained using the mean values of the data sets depicted in Figs 2, 4 and 6.

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leaflets. The increase in stiffness with age can also be explained from the observed increase in collagen cross-links with age, which corresponds to previous work from our group [36].

The valvular hemodynamics are unchanging during adult life, while the geometry, composition, and mechanical properties of the valve do change. The change in annulus diameter and leaflet dimensions during adult age, being either the consequence of growth or dilation, seems to correspond with the decrease in extensibility of the leaflets. It might be argued that a loss of tissue extensibility is compensated by growth of the leaflets to ensure valve closure with age.

Remarkably, we hardly found differences in studied parameters between the aortic and pulmonary leaflets, except for the thickness of the leaflet belly. We even found comparable mechanical properties between the aortic and pulmonary leaflets, similar to others [14,35]. It is peculiar that the evolution of mechanical properties of both valve leaflets is similar, although the transvalvular pressures on both valves remain distinct after birth (Fig 8).

Our data can be used as input for designing TE heart valves or to assess the outcomes of (other) living prostheses with known properties. In our group, Mol et al. [38] and Kortsmit et al. [39,40] investigated matrix composition and mechanical properties of TE heart valves created from human myofibroblasts seeded into rapidly degrading scaffolds. Although it is difficult to compare values of native and engineered heart valves due to possible contributions of scaffold remnants in the TE heart valves, the DNA and sGAG content of engineered heart valves were similar, while the collagen content is much lower, compared to native valves. In the engineered valves there is more sGAG compared to collagen, while in the adult native heart valves there is more collagen compared to sGAG. Since the ECM composition of the heart valve is related to maturation and mechanical properties, it might be suggested that TE protocols should be optimized to improve the hydroxyproline content and therewith the (mechanical) functionality of these heart valves. TE valves are stiffer when compared to the mechanical properties found in the present study for native valves. This suggests that not only the matrix composition and maturation are important for defining the mechanical functionality, but especially the matrix architecture seems to be important for the mechanical functionality [41,42]. Thus, to increase the long-term in vivo functionality of the TE heart valves, not only the matrix composition and maturation should be optimized, also the collagen architecture that defines the anisotropic properties of the leaflets should be studied in native valve leaflets to improve HVTE.

In conclusion, we present here for the first time an complete overview of tissue mechanical properties, matrix composition and maturation of pairs of human aortic and pulmonary valve leaflets of different age groups (fetal, child, adolescent, adult). Interestingly, the only differences between the aortic and pulmonary valve were found to be in thickness and hemodynamic properties, while the differences in tissue composition and mechanical properties were minimal for all age groups. This suggests that future therapies can focus on one living replacement for both the pulmonary and the aortic valve. Further, this study proposes that future living valve replacements for children should have a remarkably different ECM composition (lower collagen-to-sGAG ratio) and architecture to result in a lower stiffness as compared to the adult replacements.

**Study Limitations**

This study presents data on cryopreserved valves across a broad range of ages and includes sets of pulmonary and aortic valves from the same donors. As such it adds significantly to exiting studies on human aortic valves and porcine aortic valves. Still, numbers are relatively low and described trends and correlations are based on the sparsely available donor valves obtained during >5 year of research. Ideally, a large and normally distributed data set would be ideal.
and allow for regression analyses with age as a continuous variable to provide for sound conclusions on age-related effects, but this is unrealistic for donor valves. Therefore, we provided correlation graphs (Figs 2 and 4–7), as well as a more pragmatic distribution of age groups, based on the available but limited data set. As such, the child group included all available valves in children’s ages (2 months–11 years), the adolescent group the valves of donor 18–22 years of age and the adult group the other available valves (only above the age of 35). To strengthen our data we encourage other researchers to add to our data, which is available upon request.

Despite reported similarities in tissue microstructure and mechanical properties of cryopreserved valves [17, 19], it is difficult to assume the complete absence of cryopreservation effects on tissue biochemical composition and function. For instance, it has been reported that valvular GAG content is influenced by cryopreservation, but is better preserved when the valves are frozen in open leaflet position [43], as was the case in our study. Ongoing studies in our labs therefore compare fresh versus cryopreserved valves with respect to mechanical behavior and biochemical composition. Nevertheless, assuming comparable impact of cryopreservation on tissues of different age groups, all valves in the present study underwent similar freeze-thawing protocols, suggesting that the observed trends are likely to be attributed to age or age-related changes and not to cryopreservation.

The mechanical properties of the valve leaflets in the present study were obtained from experimental tensile and indentation studies only and presented as leaflet extensibility and (compressive and tensile) moduli. One should note that indentation and tension lead to different material behavior and therefore the compressive and tensile moduli cannot be directly related. For a complete description and prediction of mechanical behavior of the leaflets, however, a numerical-experimental approach comprising a constitutive model of the non-linear visco-elastic behavior is required. While this has been applied in recent studies by our group for both tensile [20] and indentation test [44] of the human native heart valve sets, here we chose a more comprehensive representation of leaflet properties, which is quite common in papers addressing in-vivo outcomes of engineered heart valves and heart valve pathology.

Supporting Information

S1 Appendix. Extended description of the cell phenotype, tissue composition and maturation analysis.

S1 Fig. Representative histological and immunofluorescent stainings on pulmonary valves. Figures A–E represent the whole pulmonary valve (leaflet, hinge region, and wall, while in E–W a representative part of the leaflet is shown. (A, C–F, H–J) Verhoeff-Van Gieson staining for collagen (red) and elastin (black). (B, G) Elastin was observed in the fetal valve using immunofluorescence (red). (K–N) Collagen type I (red) was predominant in the fibrosa. (O–R) Safranin-O staining showed proteoglycan presence (red/orange) mainly in the spongiosa and the hinge region. (S–W) αSMA (immunofluorescence; green) with cell nuclei (in blue). Scale bar: 500 μm. l: leaflet; w: wall; f: fibrosa; s: spongiosa; v: ventricularis.

S1 Table. Age groups composition and experiments. Overview of the composition of the age groups and the experiments performed on the sets of pulmonary and aortic heart valve leaflets (indicated with an ‘X’).

Supporting Information
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Author Contributions

Conceived and designed the experiments: DvG ALFS ADM FPTB CVCB. Performed the experiments: DvG ALFS PJAO MWJTJvdB. Analyzed the data: DvG ALFS PJAO ADM MWJTJvdB FPTB CVCB. Contributed reagents/materials/analysis tools: PJAO AJvdB AJJCB MJTHG. Wrote the paper: DvG ALFS PJAO ADM MWJTJvdB AJvdB AJJCB MJTHG FPTB CVCB.

References

1. Ross D, Jackson M, Davies J. The pulmonary autograft—a permanent aortic valve. Eur J cardio-thoracic Surg. 1992; 6: 113–116.

2. Schoen FJ. Heart valve tissue engineering: quo vadis? Curr Opin Biotechnol. 2011; 22: 698–705. doi:10.1016/j.copbio.2011.01.004 PMID: 21315575

3. Moretti A, Whitehouse MW. Changes in the mucopolysaccharide composition of bovine heart valves with age. Biochem J. 1963; 87: 396–402. PMID: 16749009

4. Stephens EH, Grande-Allen KJ. Age-related changes in collagen synthesis and turnover in porcine heart valves. J Heart Valve Dis. 2007; 16: 672–682. PMID: 18095519

5. Stephens EH, Chu C-K, Grande-Allen KJ. Valve proteoglycan content and glycosaminoglycan fine structure are unique to microstructure, mechanical load and age: Relevance to an age-specific tissue-engineered heart valve. Acta Biomater. 2008; 4: 1148–1160. doi:10.1016/j.actbio.2008.03.014 PMID: 18448399

6. Stephens EH, Jonge N de, McNeill MP, Durst CA, Grande-Allen KJ. Age-related changes in material behavior of porcine mitral and aortic valves and correlation to matrix composition. Tissue Eng Part A. 2010; 16: 867–878. doi: 10.1089/ten.TEA.2009.0288 PMID: 19814589

7. Kasyanov V, Moreno-Rodriguez Ra, Kalejs M, Ozolanta I, Stradins P, Wen X, et al. Age-related analysis of structural, biochemical and mechanical properties of the porcine mitral heart valve leaflets. Connect Tissue Res. 2013; 8207: 1–9.

8. Gross L, Kugel MA. Topographic Anatomy and Histology of the Valves in the Human Heart. Am J Pathol. 1931; 7:445–474.7. PMID: 19969978

9. Bashey RI, Torii S, Angrist A. Age-related collagen and elastin content of human heart valves. J Gerontol. 1967; 22: 203–208. PMID: 6022112

10. Maron BJ, Hutchins GM. The development of the semilunar valves in the human heart. Am J Pathol. 1974; 74: 331–344. PMID: 4817158

11. Vessely I, Casarotto DC, Gerosa G. Mechanics of cryopreserved aortic and pulmonary homografts. J Heart Valve Dis. 2000; 9: 27–37. PMID: 10678373

12. McDonald PC, Wilson JE, McNeill S, Gao M, Spinelli JJ, Rosenberg F, et al. The challenge of defining normality for human mitral and aortic valves: geometrical and compositional analysis. Cardiovasc Pathol. 2002; 11: 193–209. PMID: 12140123

13. Stradins P, Lacis R, Ozolanta I, Purina B, Ose V, Feldmane L, et al. Comparison of biomechanical and structural properties between human aortic and pulmonary valve. Eur J cardio-thoracic Surg. 2004; 26: 634–639.

14. Hinton RB, Yutzey KE. Heart valve structure and function in development and disease. Annu Rev Physiol. 2011; 73: 29–46. doi: 10.1146/annurev-physiol-012110-142145 PMID: 20809794

15. Christie GW, Barratt-Boyes BG. Age-dependent changes in the radial stretch of human aortic valve leaflets determined by biaxial testing. Ann Thorac Surg. 1995; 60: S156–B. PMID: 7646149

16. Aikawa E, Whitaker P, Farber M, Mendelson K, Padera RF, Aikawa M, et al. Human semilunar cardiac valve remodeling by activated cells from fetus to adult: implications for postnatal adaptation, pathology, and tissue engineering. Circulation. 2006; 113: 1344–1352. PMID: 16534030

17. Gerson CJ, Goldstein S, Heacox AE. Retained structural integrity of collagen and elastin within cryopreserved human heart valve tissue as detected by two-photon laser scanning confocal microscopy. Cryobiology. 2009; 59: 171–179. doi: 10.1016/j.cryobiol.2009.06.012 PMID: 19591817
18. Virues Delgadillo JO. Effect of freezing on the passive mechanical properties of arterial samples. J Biomed Sci Eng. 2010; 03: 645–652.
19. Cox MAJ, Driessen NJB, Boerboom RA, Bouten CVC, Baaijens FPT. Mechanical characterization of anisotropic planar biological soft tissues using finite indentation: experimental feasibility. J Biomech. 2008; 41:422–429. PMID: 17897653
20. Soares ALF, van Geemen D, van den Bogaerd AJ, Oomens CWJ, Bouten CVC, Baaijens FPT. Mechanics of the pulmonary valve in the aortic position. J Mech Behav Biomed Mat. 2014; 29:557–567.
21. Merryman WD, Youn I, Lukoff HD, Krueger PM, Guilak F, Hopkins R, et al. Correlation between heart valve interstitial cell stiffness and transvalvular pressure: implications for collagen biosynthesis. Am J Physiol Heart Circ Physiol. 2006; 290: H224–31. PMID: 16126816
22. Kriemler S, Jansen C, Linka A, Kessel-Schaefer A, Zehnder M, Schürmann T, et al. Higher pulmonary artery pressure in children than in adults upon fast ascent to high altitude. Eur Respir J. 2008; 32: 664–669. doi: 10.1183/09031936.00166407 PMID: 18417505
23. Gao Y, Raj JU. Regulation of the Pulmonary Circulation in the Fetus and Newborn. Physiol Rev. 2010; 90: 1291–1335. doi: 10.1152/physrev.00032.2009 PMID: 20959617
24. Votteler M, Carvajal Berrio D a, Horke A, Sabatier L, Reinhardt DP, Naïr A, et al. Elastogenesis at the onset of human cardiac valve development. Development. 2013; 2353: 2345–2353.
25. Merryman WD. Mechano-potential etiologies of aortic valve disease. J Biomech. 2010; 43: 87–92. doi: 10.1016/j.jbiomech.2009.09.013 PMID: 19811785
26. Schoen FJ. Mechanisms of function and disease of natural and replacement heart valves. Annu Rev Pathol. 2012; 7: 161–183. doi: 10.1146/annurev-pathol-011110-130257 PMID: 21942526
27. Driessen NJB, Bouten CVC, Baaijens FPT. Improved prediction of the collagen fiber architecture in the aortic heart valve. J Biomech Eng. 2005; 127: 329–336. PMID: 15971711
28. Sacks MS, David Merryman W, Schmidt DE. On the biomechanics of heart valve function. J Biomech. 2009; 42: 1804–1824. doi: 10.1016/j.jbiomech.2009.05.015 PMID: 19540499
29. Rabkin E, Hoerstrup SP, Akaiw M, Mayer JE, Schoen FJ. Evolution of cell phenotype and extracellular matrix in tissue-engineered heart valves during in-vitro maturation and in-vivo remodeling. J Heart Valve Dis. 2002; 11: 308–314. PMID: 12056720
30. Rabkin-Akaiw E, Farber M, Akaiw M, Schoen FJ. Dynamic and reversible changes of interstitial cell phenotype during remodeling of cardiac valves. J Heart Valve Dis. 2004; 13:841–847. PMID: 15473488
31. Simmons CA, Grant GR, Manduchi E, Davies PF. Spatial heterogeneity of endothelial phenotypes correlates with side-specific vulnerability to calcification in normal porcine aortic valves. Circ Res. 2005; 96:792–799. PMID: 15761200
32. Sucosky P, Balachandran K, Elhammali A, Jo H, Yoganathan AP. Altered shear stress stimuli upregulates endothelial vcam-1 and icam-1 in a bmp-4 and tgf-beta1-dependent pathway. Artherosclerosis Tromb Vasc Biol. 2009; 29:254–260.
33. Hjortnaes J, Shapero K, Goettsch C, Hutcheson JD, Keegan J, Kluin J, et al. Valvular interstitial cells suppress calcification by valvular endothelial cells. Atherosclerosis. 2015; 242:251–260. doi: 10.1016/j.atherosclerosis.2015.07.008 PMID: 26232165
34. Sauren AA, Hout MC van, Steenhoven AA van, Veldpaus FE, Janssen JD. The mechanical properties of porcine aortic valve tissues. J Biomech. 1983; 16: 327–337. PMID: 6885834
35. Leeson-Dietrich J, Boughner D, Vesely I. Porcine pulmonary and aortic valves: a comparison of their tensile viscoelastic properties at physiological strain rates. J Heart Valve Dis. 1995; 4; 88–94. PMID: 7742995
36. Balguid A, Rubbens MP, Mol A, Bank RA, Bogers AJJC, Kats JP van, et al. The role of collagen crosslinks in biomechanical behavior of human aortic heart valve leaflets—relevance for tissue engineering. Tissue Eng. 2007; 13: 1501–1511. PMID: 17518750
37. Hasen A, Rabaret K, Swieszowski W, Selimović S, Paul A, Camci-Unal G, et al. Biomechanical properties of native and tissue engineered heart valve constructs. J Biomech. 2014; 13: 1949–1963.
38. Mol A, Rutten MCM, Driessen NJB, Bouten CVC, Zünd G, Baaijens FPT, et al. Autologous human tissue-engineered heart valves: prospects for systemic application. Circulation. 2006; 114: 1152–8. PMID: 16820565
39. Kortsmit J, Rutten MCM, Wijaars MW, Baaijens FPT. Deformation-controlled load application in heart valve tissue engineering. Tissue Eng Part C Methods. 2009; 15: 707–716. doi: 10.1089/teng.TEC.2008.0658 PMID: 19275473
40. Kortsmit J, Driessen NJB, Rutten MCM, Baaijens FPT. Nondestructive and noninvasive assessment of mechanical properties in heart valve tissue engineering. Tissue Eng Part A. 2009; 15: 797–806. doi: 10.1089/ten.tea.2008.0197 PMID: 19105603

41. Sacks MS, Schoen FJ. Collagen fiber disruption occurs independent of calcification in clinically explanted bioprosthetic heart valves. J Biomed Mater Res. 2002; 62: 359–371. PMID: 12209921

42. Lindeman JHN, Ashcroft BA, Beenakker J-WM, Es M van, Koekkoek NBR, Prins FA, et al. Distinct defects in collagen microarchitecture underlie vessel-wall failure in advanced abdominal aneurysms and aneurysms in Marfan syndrome. Proc Natl Acad Sci U S A. 2010; 107: 862–865. doi: 10.1073/pnas.0910312107 PMID: 20080766

43. Dainese L, Barili F, Topkara VK, Cheema FH, Formato M, Aljaber E, et al. Effect of Cryopreservation Techniques on Aortic Valve Glycosaminoglycans. Artificial Organs. 2006; 30(4): 259–264. PMID: 16643384

44. Oomen PJA, Loerakker S, van Geemen D, Neggers J, Goumans M-JTH., van den Bogaerd AJ, et al. Age-dependent changes of stress and strain in the human heart valve and their relation with collagen remodeling. Acta Biomater. 2016; 29: 161–169. doi: 10.1016/j.actbio.2015.10.044 PMID: 26537200