Differentiating the aging of the mitral valve from human and canine myxomatous degeneration

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Abstract During the course of both canine and human aging, the mitral valve remodels in generally predictable ways. The connection between these aging changes and the morbidity and mortality that accompany pathologic conditions has not been made clear. By exploring work that has investigated the specific valvular changes in both age and disease, with respect to the cells and the extracellular matrix found within the mitral valve, heretofore unexplored connections between age and myxomatous valve disease can be found. This review addresses several studies that have been conducted to explore such age and disease related changes in extracellular matrix, valvular endothelial and interstitial cells, and valve innervation, and also reviews attempts to correlate aging and myxomatous disease. Such connections can highlight avenues for future research and help provide insight as to when an individual diverts from an aging pattern into a diseased pathway. Recognizing these patterns and opportunities could result in earlier intervention and the hope of reduced morbidity and mortality for patients.

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Introduction

Myxomatous mitral valve degeneration (MMVD) causes severe valvular regurgitation and ultimately death for millions of dogs around the world.1,2 In humans, this disease is the primary cause of mitral valvular regurgitation leading to the need for surgical valve repair.3 For humans afflicted with MMVD, it has been observed that the mean patient age is 60 years old. As opposed to calcific aortic valve disease, which has a clearly demonstrated association with aging,4 MMVD in humans appears distinct from age-related changes.5–7 This review will describe the age-related changes in mitral valve composition and material behavior and

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contrast these changes with those found in MMVD, with a particular emphasis on comparisons of human, porcine, and canine mitral valves.

**Gross and micro-anatomy of the mitral valve**

The canine mitral valve complex, similar to human and other mammalian species, consists of the annulus, two leaflets (anterior and posterior) of different shape and size, numerous chordae tendineae, and the papillary muscles (Fig. 1A). The function of the mitral valve is to direct the blood flow from the left atrium to the left ventricle during diastole and to prevent the backflow of blood into the left atrium during systole. In order to do so, the mitral valve complex requires all of its components, together with the adjacent atrial and ventricular muscle, to work in a synchronized fashion.

The mitral valve leaflets each consist of four layers that differ from each other in extracellular matrix (ECM) makeup and functionality. From top to bottom, the layers are the atrialis, spongiosa, fibrosa, and ventricularis. The fibrosa, the thickest layer of the valve, is comprised of dense, circumferentially aligned collagen, in contrast to the loose collagen found within the atrialis, spongiosa, and ventricularis. The dense collagen of the fibrosa provides the mitral valve leaflets with tensile strength, whereas the looser collagen and high glycosaminoglycan (GAG) content in the other layers provides the leaflets with compressive strength. The atrialis, the layer located on the inflow side of the mitral valve, has an abundance of the protein elastin, which allows the valve to undergo considerable stretch and then recoil back to its original undeformed shape during the cardiac cycle. The spongiosa contains a high concentration of the GAG hyaluronan (HA) and hydrated chondroitin/dermatan sulfate proteoglycans (PGs), which also provide compressive strength for the valve. The ventricularis layer of the mitral valve leaflets is very thin compared with other layers and consists of elastic fibers and regularly spaced, circumferentially oriented collagen fibers. The regions containing the highly aligned collagen are also rich in small leucine-rich proteoglycans (SLRPs) that connect to and provide mechanical support for the collagen fibrils.

In addition to the mitral valve leaflets, the chordae tendineae are a critical component to the functioning of the mitral valve apparatus. These chordae connect the underside of the valve
leaflet to the papillary muscles along the left ventricular wall. As the left ventricle contracts during systole, the papillary muscles contract as well and the chordae in turn provide mechanical support to the mitral valve leaflets, helping them maintain closure during the highly pressurized left ventricular systole. In order to withstand the high tensile stress applied to the chordae, they are comprised of highly organized collagen fibers that are oriented in the direction of load. As in the leaflet, these regions of highly aligned collagen are also rich in SLRPs, such as decorin and biglycan. The GAGs found within the chordae are predominantly those found on the SLRPs, and demonstrate an abundance of iduronate and 4-sulfated-N-acetylgalactosamine. The chordae also contain a small distribution of elastic fibers in their outer sheath.

The mitral valve leaflets also vary in makeup and function from the annulus to the free edge. For example, there is a distinct difference between the so-called "rough zone" and "clear zone" of the anterior leaflet. Historically, the "rough zone" has referred to those parts of the leaflet that contain chordal attachments, while the "clear zone" has referred to regions without chordal attachment. In our previous studies of mitral valve leaflets we have defined these regions as the mitral valve free edge (MVF) and the center of the anterior leaflet (MVAC), respectively (Fig. 1B). The MVAC maintains the tensile integrity of the valve compared with the MVF, the MVAC contains a thicker fibrosa layer and has a lower concentration of GAGs. The specific characteristics of the GAGs also vary between these two regions, e.g., the MVAC contains relatively less unsulfated, 6-sulfated, and 4-sulfated glucuronate than the MVF. In contrast, the MVF, which is more suited to bearing compressive loading, has a much higher concentration of HA than does the MVAC, mainly because the free edge has a much larger spongiosa component compared to the MVAC. The composition of the mitral valve posterior leaflet (also known as the mural leaflet), which experiences a compressive load during the cardiac cycle, closely resembles that of the other major compressive load-bearing region, the MVF.

Due to the lack of comprehensive studies on normal valvular aging in dogs, it becomes necessary to look to studies conducted on other species in order to provide some insight into aging and its relation to MMVD. However, it is important to note the anatomical and therefore potential histological differences between species when doing so. For example, it has been reported that larger animals have a distinct increase in the branching (first-order and second-order) of chordae when compared to the hearts of smaller animals. The need to support the larger valve cusp area of the larger animals seems to be the logical explanation for this structural arrangement. Variations in shapes of the anterior leaflet (also known as the aortic or septal leaflet) have also been reported when comparing human and canine mitral valves.

Human anterior leaflets have a clear line demarcating the boundary of the appositional area, the area of leaflet in contact during valve closure. In dogs, the absence of a clear line of demarcation, together with higher muscle content in both mitral cusps than found in human valves, has led to the speculation that canine mitral valves demonstrate a different closing mechanism. Humans and dogs also have different angles between the plane of the atrioventricular orifice in relation to the axis extending from the ventricular apex through the middle of the aortic valve. However, the significance of this finding is unknown and could be related to the 90° angle difference in anteroposterior axis orientation relative to the gravitational field between dogs (a quadruped mammal) and bipedal humans.

Aging and cell-mediated remodeling of the extracellular matrix

Studies of human and porcine mitral valves have shown the effect of aging on ECM and microstructural composition as well as the effects of these changes on valve function. With age, there is a reduction in cell density throughout the valve, across all layers. The valvular interstitial cells (VICS) that remain are increasingly associated with an "activated" remodeling phenotype of α-smooth muscle actin (α-SMA), co-localized with enzymes involved in collagen synthesis and degradation in porcine valves. Microstructurally, the layers become more delineated with advancing age, and the PGs decorin and biglycan show greater expression throughout the valve. These PGs colocalize with collagen and aid in collagen fibrillogenesis, lending further support that collagen remodeling occurs throughout aging. In contrast, with aging, the amount of elastin decreases throughout the valve, particularly for individuals over the age of 50 years. In addition, with advancing age, the incidence of lipid accumulation and calcification in valves increases throughout the human population (Fig. 2).

With respect to types of GAGs present in the mitral valve, with age there is a decrease in unsulfated and 6-sulfated glucuronate, as well as...
a decrease in the ratio of chondroitin sulfate to dermatan sulfate. Interestingly, the total amount of HA remains fairly constant with age in porcine mitral valves, although HA comes to represent an increasing percentage of the total GAGs in the MVF due to a reduction in 4-sulfated and 6-sulfated GAGs. In the MVAC, there is no age-related decrease in total GAGs or in HA, but there is an age-related increase in 6-sulfated glucuronate and 4-sulfated iduronate (Figs. 3 and 4).

Aging and mechanical behavior

With respect to mechanical behavior, the overall stiffness of the valve increases in both the radial and circumferential directions as a function of age. Accompanying these age-related changes are a reduction in extensibility and an increase in stress relaxation. Extensibility can be generally described as the magnitude that a tissue can stretch before the collagen fibers become fully uncrimped and bear load (Fig. 5). Stress relaxation is an indicator of the viscoelastic (time-dependent) behavior of the tissue; an increase in stress relaxation indicates that there is a substantial rearrangement of the extracellular matrix that progresses for some time after tissues are stretched. These altered material properties can be generally explained by thickening of the collagen-rich fibrosa layer as well as the increase in collagen production and remodeling that occurs throughout aging. It is also likely that there is collagenous reinforcement of elastic fibers with age, which would reduce extensibility. There is an increase in the radius of curvature of the transition region of the valve leaflet load–deformation curve with age, which we speculate is due to increased collagen alignment and crosslinking in the valve as well as greater abundance of PGs and GAGs in the collagenous fibrosa layer. Indeed, the SLRP decorin and biglycan, both of which become more abundant with age in valves, bind to and form bridges between collagen fibrils. Therefore, increased transmission of hemodynamic forces to collagen and the other underlying matrix proteins via interconnecting PGs could be responsible for the age-related changes in these mechanical properties.

Aging and innervation

Over the past several years, the recognition that mammalian mitral valve leaflets have well developed nerve plexuses, consisting of different transmitter-specific subpopulations of nerves, has challenged the conventional concept of mitral valve as a passive flap. The nerve plexus is most dense in the basal zone of the valve adjacent to the fibrous annulus ring, and becomes finer and more branched toward the free margin of the cusp. It has been shown in human mitral valves that the nerve density within the anterior leaflet is two fold greater than in the posterior leaflet and that the nerve fibers are closely associated with different types of VICs. Together with contractile cell types found within the valve (activated VICs and cardiomyocytes from the annulus), the nerve plexuses form a model for an active valve that is capable of contraction and relaxation, fine tuning its motion during each cardiac cycle. Giving support to this claim, experiments in dogs have shown that autonomic stimulation results in the valve better able to withstand pressure-induced displacement of the leaflets into the left atrium. With increasing age, there is a loss of innervation in the mitral valves of guinea pigs, mice and rats. In rats, both sensory and motor type of nerves were markedly diminished in older animals. There was no difference in mitral valve nerve density between young dogs aged 6–12 months and adult dogs aged 5 years, but there was a significant reduction in mitral valve nerve density in dogs above 10 years of age.
Aging and valvular cells

The valve leaflets are covered with a single layer of valvular endothelial cells (VECs) that is continuous with the endocardium of the heart. The VECs are believed to coordinate with the VICs located in the leaflet interior to maintain the ECM structure and contribute to the mechanical function of the valve. Although VICs and VECs demonstrate distinctly different phenotypes in the postnatal mitral valve, they share a common embryonic origin. During the earliest stage of valve development, the GAG-rich cardiac cushions arise within the atrioventricular canal and outflow tract of the looped heart tube. Certain endothelial cells on the surface of these cardiac cushions migrate into the interior of the cushion through a process referred to as endothelial—mesenchymal transformation (EMT) and ultimately become the VICs; it is believed that the underlying myocardium regulates the EMT. It does so through the regulation of growth factors including VEGF, TGF-β, NOTCH, and WNT, as well as extracellular matrix components, such as hyaluronan, which regulate pathways that direct endothelial cells toward mesenchymal transformation.

Once activated for mesenchymal transformation, the cells within these cushion regions then initiate ECM remodeling and later development of the valve leaflets. In contrast, the VECs do not undergo EMT during normal aging (Fig. 6). However, recent experiments indicate that some adult VECs retain the ability to differentiate in a manner similar to EMT while in a diseased state.

Although age-related changes in VECs have been investigated infrequently, scanning electron microscopy has been used to demonstrate denudation of the endothelium in some areas of the diseased valves of older dogs (9–15 years) with regional pleomorphism in adjacent areas of intact endothelium. In addition, it has been shown in human valves that fetal VECs (second and third...
trimesters) express significantly higher amounts ($p < 0.001$) of $\text{SMemb}$, matrix metalloproteinase-1 (MMP-1), MMP-13, and cell adhesion molecules ICAM-1 and VCAM-1 when compared to adult VECs, which had relatively negligible expression of these proteins.

Myxomatous changes alter valve motion, resulting in abnormal closure that, taken together with the changing in hemodynamic force due to regurgitation, may cause or contribute to endothelial damage. Damage to the VEC layer could influence the synthesis and release of vasoactive mediators that in turn interact with subendothelial matrix tissue. One such potent vasoconstrictor is endothelin, which also has the potential to stimulate proliferation of fibroblasts and increase collagen production. Indeed, an increase in endothelin receptor density has been found in the distal end of aged canine mitral leaflets (mean age 12 years) showing myxomatous changes.

VICs are the predominant cells found in all layers of mitral valve and are considered to be primarily responsible for maintaining valve structure and function. VICs are a heterogeneous and dynamic population of cells that have a range of distinct cellular phenotypes. This plasticity of VIC phenotype appears to be crucial in heart valve development, remodeling and repair, and progression of diseased states — all instances in which the myofibroblast cell type predominates. Liu et al. has proposed that there are five identifiable VIC phenotypes during different stages of mitral valve development, remodeling, and disease. These five are embryonic progenitor endothelial/mesenchymal cells (EMT stage), quiescent VICs (normal fibroblast like), activated VICs (diseased myofibroblast like), progenitor VICs (stem cell like) and osteoblastic VICs (Fig. 7).

Despite such characterizations, the current understanding of VIC biology remains very limited and information on the aging aspect of VIC biology is particularly rare. An early investigation of mitral valves from humans aged 5 days to 89 years reported a gradual age-related decrease in VIC number and nuclei size, notably after the fifth decade of life. High densities of VICs were found in fetal mitral valves from human, sheep and pig. In pigs, the highest VIC densities were
found in the first trimester, after which there was a decrease in cell number throughout fetal developmental and postnatal aging. Interestingly, distinctions between layers are evident as early as the first trimester, with the spongiosa layer having a lower cell density than the outer leaflet layers. During fetal valve development, most VICs stain positive for vimentin, α-SMA, and the collagenase MMP-13, and are thus considered to have an activated myofibroblast phenotype, the cell phenotype most involved in tissue remodelling. Throughout development and into adulthood, there is a significant reduction in the numbers of α-SMA positive VICs and a relatively greater number of the normal quiescent fibroblast phenotype in humans.

**Myxomatous mitral valve degeneration**

MMVD is the most common cardiac disease in the dog. The disease is also referred to as mitral valve degeneration (MVD). It is characterized by the degeneration of the mitral valve leaflets, which can lead to regurgitation of blood back into the left atrium. The exact cause of MMVD is unknown, but it is thought to be related to a combination of genetic, environmental, and immunological factors. The disease typically affects older dogs, particularly those between the ages of 10 and 15 years.

**Figure 5** Representative stress-strain curve seen in mechanical testing of valves in tension (solid line). The first linear portion of the bi-linear curve is called the toe region. It corresponds to the region where collagen is crimped and elastic fibers dominate the material properties of the valve. The second linear portion can be referred to as the collagen region because it corresponds to the region where collagen is fully uncrimped and is dominating the material properties. The slope of this region of the curve is defined as the Young’s modulus or stiffness of the material. One can determine a value for the extensibility by extrapolating the collagen region backwards using the Young’s modulus as the slope (dashed line) to where it crosses the x-axis. The extensibility corresponds to the degree of crimping of collagen in the valve. The curved area between these two regions is the transition region. It corresponds to the region in which collagen is actively uncrimping. The sharpness of this transition, defined by the radius of transition curvature, corresponds to the alignment and crosslinking present in the collagen in the tissue.

**Figure 6** Graphic overview of heart development and endothelial–mesenchymal transformation (EMT). As the heart tube develops it contains three layers, an inner lining of endothelial cells, a middle separating layer of ECM referred to as cardiac jelly, and an outer layer of myocardium. As valves form, a subset of the endothelial cells undergo EMT by delaminating, differentiating, and then migrating into the cardiac jelly. Then, in a process that is poorly understood, local swellings of cardiac jelly and mesenchymal cells (cardiac cushions) undergo remodeling and form heart valves. Reprinted from Armstrong, E. J., & Bischoff, J. (2004). Heart valve development: endothelial cell signaling and differentiation. Circ Res, 95(5), 459–470 with permission from Wolters Kluwer Heath.

**Figure 7** The functions of VICs can be organized into five major phenotypes: embryonic progenitor endothelial/mesenchymal cells, quiescent VICs (qVICs), activated VICs (aVICs), stem cell-derived progenitor VICs (pVICs), and osteoblastic VICs (obVICs). Through the process of EMT, embryonic progenitor endothelial/mesenchymal cells differentiate into aVICs and qVICs. aVICs are capable migration, proliferation, and ECM synthesis. qVICs remain quiescent until the heart valve is injured, at which point they transition to aVICs and aid in the repair and remodeling process. In addition, they can differentiate into obVICs in certain conditions. pVICs are derived from bone marrow. They are present in the bone marrow, in circulation, and in the heart valve. They are an additional source of aVICs in adulthood. obVICs respond to osteogenic and chondrogenic factors and promote valve calcification. Hatched arrows represent hypothesized transitions for which there is currently not solid evidence. Other abbreviations used: EPCs = Endothelial Progenitor Cells, DCs = Dendritic cells. Reprinted from Liu AC, Joag VR, Gotlieb AI. The emerging Role of Valve Interstitial Cell Phenotypes in regulating Heart Valve Pathobiology. Am J Pathol. 2007;171(5):1407–1418 with permission from Elsevier.
endocardiosis or chronic valvular disease, and affects many different species, including pigs\textsuperscript{47-49} and humans in addition to dogs. In dogs, unlike humans, the prevalence and severity of the disease is found to be closely age dependent.\textsuperscript{50} However, as described later, some breeds of dog are more predisposed to the disease than others, suggesting a likely inherited component to the disease.\textsuperscript{51,52} Various reports have estimated the occurrence of MMVD as ranging from 21% to 89% in dogs.\textsuperscript{46,53,54} This wide range might be attributed to the disparate clinical sample selection of the groups of dogs, which could be affected by factors such as sample pool size, age, and breed.

MMVD involves a complex connective tissue degenerative process with little or no inflammatory reaction. Histologically, MMVD is manifested by excessive destruction and derangement of the valve stroma with loss of collagen bundle organization and accumulation of PGs and GAGs in the leaflets and chordae.\textsuperscript{7,16,55,56} In addition, elastic fibers are increased in number, but appear to be disrupted and granular in nature.\textsuperscript{56} In dogs, the pathology is grossly recognized by the presence of greyish-white, smooth, glistening nodules or plaque-like elevations, also referred to as lesions, often situated in the area of apposition on the atrial surface of the leaflets.\textsuperscript{57} In humans, a much larger region of the tissue is affected by myxomatous degeneration, which may extend throughout the leaflet and chordae. Normal healthy mitral valve leaflets are thin, pliable, translucent and soft. In contrast, diseased valve leaflets and chordae become opaque and thickened; they are more extensible and less stiff than normal leaflets.\textsuperscript{58} The structural changes due to myxomatous degeneration in the leaflets are not always uniformly distributed, but have long been identified as the most important factors contributing to mitral valve dysfunction in dogs.\textsuperscript{46,54,57,59,60} The rough zone of the leaflet on the ventricular side, especially where the tendinous chords attach, is particularly prone to myxomatous degeneration.\textsuperscript{61} At later stages of this disease, gross distortion of the apposition line of the leaflets can be caused by the development of interchordal hoodings,\textsuperscript{62} also referred to as parachute change. The middle section of the leaflet, which withstands the strongest pressure loading during the cardiac cycle,\textsuperscript{63} is less susceptible to myxomatous degeneration in dogs. In humans with MMVD, the mid-section of the leaflets is frequently involved, especially the middle scallop of the posterior (mural) leaflet with changes observed in 88% of patients.\textsuperscript{6,64}

Both dogs and humans with MMVD demonstrate phenotypic changes in VICs from a quiescent fibroblast to an activated myofibroblast (α-SMA positive).\textsuperscript{42,59,65-67} Many of these cells also express stem/progenitor cell markers CD-34 and/or CD-117.\textsuperscript{42,68,69} In the early stages of MMVD in dogs, α-SMA positive cells are found to congregate toward the atrialis edge, then extend more deeply into the tissue.\textsuperscript{70} At later stages of this disease, α-SMA positive cells also accumulated in the distal ventricular side, presumably as a result of regurgitation-driven forces.\textsuperscript{65} Moreover, VICs staining positive for α-SMA were frequently co-localized with expression of TGF-β1 and TGF-β3. Interestingly, in the overtly myxomatous area of canine valves, α-SMA positive cells were uncommon, as was TGF-β3 expression, although TGF-β1 was strongly expressed.\textsuperscript{71} There was also a reduction in vimentin-positive cells in myxomatous areas of canine valves,\textsuperscript{65} compared with normal valve regions. Cell density in myxomatous areas was not significantly altered between different grades of MMVD or with canine age, downplaying a role for cell recruitment or cell proliferation in this condition in dogs.\textsuperscript{72} In humans, myxomatous mitral valves demonstrate an increase in certain catabolic enzymes and other phenotypic markers, including MMPs, cathepsins, SMemb, and interleukin-1β, that are associated with "activated myofibroblasts".\textsuperscript{42}

MMPs and tissue inhibitors of metalloproteinases (TIMPs) both play an important role in the remodeling of ECM, both in aging and in MMVD. For example, both MMP-2 and MMP-9 are increased in myxomatous mitral valves.\textsuperscript{73} In addition, it was shown that the expression of MMP-2 decreased and the expression of MMP-14 and of TIMP-2 and -3 increased with increased MMVD severity and age,\textsuperscript{74} with the decrease in MMP-2 attributed to increased abundance of its inhibitor TIMP-3.\textsuperscript{75,76}

In dogs, an increase in the severity of valvular damage in MMVD corresponds to a worsening of the clinical disability.\textsuperscript{1} MMVD leads to regurgitation of blood across the closed mitral valve during ventricular systole. The lesions of MMVD represent a gradual process and may cause no detectable clinical signs during early stage of structural change. Indeed, valve closure in the early stages may appear normal. However, progression of the disease will lead to insufficient coaptation of the leaflets, increasing regurgitation of blood back into the atria, and in the late stage of the disease, dilation of the left ventricle and mitral annulus occurs, as well as jet lesions and ruptured chordae in some cases. As a consequence, these lesions will lead to mitral systolic murmurs, and in severe cases, congestive heart failure. This disease progression is similar to MMVD in humans in terms of the
pathological change and the clinical outcome. In humans, however, displacement of mitral valve leaflets into the atrium can be less common, and the frequency of chordal rupture is very high.

Due to considerable variation in the size of the normal heart valve in different breeds of dogs, a visual scoring method is widely used to classify the diseased valve lesions into one of four grades (Table 1). The grading system is easy to apply and can provide useful information on the pathogenesis of the disease as each grade represents a stage in the development of the disease. However, this grading system requires a familiarization with all the manifestations of the disease and is somewhat subjective. Although MMVD in humans shares several of these changes, no similar classification system exists for human patients, partly because of the difficulty of identifying patients experiencing the early stages of the disease (Table 2).

Comparison of age-related changes and disease

Mitrval valve disease begins during the first third of a dog’s life, frequently producing mitral and occasionally tricuspid insufficiency during the middle years. The degree of valvular malformation caused by this disease is fairly well tolerated in some individuals up to 9 years of age or older. Although there is no comparable grading system in human medicine, the age distribution in dogs is comparable with the prevalence of mitral valve disease in human patients. The prevalence of another valvular disorder, mitral valve prolapse (MVP), is low in young children, but increases from childhood to adolescence, a change that is concomitant with the growth spurt of adolescence. Most reported human patients requiring mitral valve repair surgery or replacement tend to be middle-aged, with mean age at 56 years old. This age distribution suggests that while MMVD might not be a part of normal aging, aging and growth have a significant impact on disease progression.

A favored hypothesis at present is that MMVD is a response to repeated impact of the leaflet edges. Myxomatous degeneration then alters the apposition zone and valve motion resulting in abnormal closure. In addition, the changes in motion and apposition cause regurgitation that then impacts the distribution of hemodynamic forces and ultimately results in further tissue damage. An observation that links this hypothesis to the aging process is the fact that the heart rate, and thus the frequency of mitral leaflet coaptation impacts, increases with age in certain breeds. This greater frequency of impact could augment the progression of MMVD in older dogs. The location and the histology of the thickened foci on leaflet edges supports this hypothesis, but the sequence of events and the time course is still unknown.

In view of the high prevalence of MMVD in the aged canine population and that its clinical symptom, mitral regurgitation, represents the most important geriatric heart disease, some veterinarians and researchers have classified MMVD as a geriatric disease. Regardless of classification, increasing age has a marked effect on prevalence and severity in canine MMVD as well as human MVP. For this reason as well as the histological similarities between dogs and humans, canine MMVD has been proposed as a naturally occurring model of human MVP.

As described earlier, important ECM changes that occur during the aging process include remodeling of collagen, altered GAG and PG composition, and decreased elastin. Moreover,

![Table 1](https://example.com/table1.png)

<table>
<thead>
<tr>
<th>Lesion Grade</th>
<th>Nodules (Location)</th>
<th>Leaflets (Location)</th>
<th>Chordae</th>
<th>Valvular Competence</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Few, small oedematous (apposition)</td>
<td>Areas of diffuse opacity (basal)</td>
<td>Unchanged</td>
<td>Maintained</td>
</tr>
<tr>
<td>2</td>
<td>Multiple, grayish-white, coalesce (apposition)</td>
<td>Areas of diffuse opacity (throughout)</td>
<td>Unchanged</td>
<td>Maintained</td>
</tr>
<tr>
<td>3</td>
<td>Increased size, plaque-like (apposition)</td>
<td>Defined areas of opacity (basal)</td>
<td>Thickened at junction with leaflets</td>
<td>Some incompetent</td>
</tr>
<tr>
<td>4</td>
<td>Greyish-white, plaque-like elevations (apposition)</td>
<td>Grossly distorted; ballooning overgrowth (closure line)</td>
<td>Thickened, can be stretched/ruptured</td>
<td>Majority incompetent</td>
</tr>
</tbody>
</table>

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progressive thickening of the leaflet cusps was a notable change during the aging process.\textsuperscript{20,88}

Given the similarity between these histological and gross findings and those of MMVD in dogs it is possible that a link exists between MMVD and normal aging of dogs. MMVD might, in fact, be an extreme or abnormal amplification of a normal aging process; perhaps aged “normal” subjects in a human study would be classified as abnormal in a veterinary context. Furthermore, in human patients with MMVD, the mitral valve leaflets and chordae are thicker, and the ECM derangements are significantly more pronounced, compared with age-matched control human valves.\textsuperscript{7}

Using his lesion grading system, Whitney found the prevalence and severity of mitral valve lesions correlated with increase in age, based on a post-mortem study of 200 dogs.\textsuperscript{57} When combining all four grades, he found lesions in 37% at 0–4 years of age, 80% at 5–8 years of age, 93% at 9–12 years of age and 100% at 13–16 years of age. Type 1 lesions (15%) were found predominantly in dogs under 5 years old. Type 2 and 3 lesions (49%) were common in dogs 9–12 years old, whereas Type 3 and 4 lesions (88%) were predominately confined to dogs 13 years and over. The more advanced Type 3 and 4 lesions were found in 24% of the dogs under 9 years old, in comparison to 58% of dogs over 9 years old; the majority of dogs with Type 3 and 4 lesions showed evidence of congestive heart failure.\textsuperscript{57} A modification of Whitney’s lesion grading strategy was proposed by Kogure.\textsuperscript{54} Instead of 4 types of lesions, Kogure categorized the mitral lesions into 3 groups. Group I corresponds to Whitney’s types 1 and 2, whereas Group II is similar to Whitney’s type 3 and Group III

<table>
<thead>
<tr>
<th>Property</th>
<th>Age-Related Changes</th>
<th>Disease Related Changes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mechanical behavior and gross anatomy</td>
<td>Reduced extensibility and increased elastic modulus Thickening of leaflet cusps</td>
<td>Increased extensibility and reduced strength Both: Leaflets become opaque and thickened (especially along coaptation edge) Dogs: Greyish-white, smooth glistening nodules/plaques on atrial surface of leaflets and in rough zone Humans: Larger region affected throughout leaflets and chordae</td>
</tr>
<tr>
<td>Layers (microstructure)</td>
<td>Thickening of fibrosa layer Layers more delineated</td>
<td>Disruption of atrialis layer, expansion of spongiosa layer and destruction of fibrosa layer</td>
</tr>
<tr>
<td>Cells</td>
<td>VICs more activated (remodeling phenotype) Decrease in cell density</td>
<td>VICs and myofibroblasts activated phenotype Cell density (DNA concentration) unaltered</td>
</tr>
<tr>
<td>Collagen and MMPs</td>
<td>Increased production and remodeling Reduced crimp</td>
<td>Loss of fiber bundle organization but increase in immature collagen MMP-2 and -9 increased MMP-2 decreased MMP-14, TIMP-2 and -3 increased with severity and age</td>
</tr>
<tr>
<td>Elastin</td>
<td>Decreases Increase in collagen reinforcement of elastic fibers</td>
<td>Increase in number but appear more granular</td>
</tr>
<tr>
<td>PGs and GAGs</td>
<td>Increased abundance in fibrosa layer Decorin and biglycan greater expression Altered expression of certain sulfated GAGs Increase of chondroitin sulfate/dermatan sulfate ratio</td>
<td>Accumulate in leaflets and chordae</td>
</tr>
</tbody>
</table>

Table 2: Summary of changes seen in aging and myxomatous disease.

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resembled Whitney’s type 4. Among the 64 dogs with mitral valve lesion studied by Kogure, 30 (47%) dogs with a median age of 2 years old were classified into Group I; 22 (34%) dogs with a median age of 4 years old were in Group II, and 12 (19%) dogs with a median age of 10 years old were in Group III.54 The result was parallel to Whitney’s finding that the severity of lesions increased with age. Grading lesions as 1–4, corresponding to Whitney’s Type I–IV, is widely performed in the veterinary cardiology community. These two comprehensive studies have demonstrated the structural change in canine mitral valves is an age dependent process.

Effect of breed and genetic inheritance

Dogs of smaller breeds, including Cavalier King Charles Spaniels (CKCS), Chihuahuas, Cocker spaniels, Dachshunds and Miniature schnauzers, are particularly predisposed to MMVD,89 even though dogs of smaller breeds generally live much longer than dogs of larger breeds.90 It is well documented that certain highly predisposed breeds like the CKCS can show signs of MMVD as early as one to two years of age, with average initial diagnosis at 6.5 years old. This age is much lower than the average age of initial diagnosis of 12 years in other breeds.51,91 Because of this breed association, it has long been thought that the disease is inherited.52 Echocardiographic studies of mitral regurgitation in CKCS and Dachshunds have been conducted to confirm a parent to offspring inheritable relationship,92–94 while other studies lend support that MMVD is an autosomal dominant complex polygenic trait with variable penetrance.93 A breeding program aimed at reducing the prevalence of MMVD in the Swedish CKCS population started in 2000, but thus far it has been unsuccessful.5 An inherited form of the disease is seen in human Marfan syndrome patients who develop myxomatous mitral degeneration at an early age. However, Marfan syndrome is caused by the mutation a single gene encoding for the extracellular structural protein fibrillin.95–97 Some cases of mitral valve disease in humans, specifically familial cases of apparently autosomal dominant or X-linked inheritance of MVP or MMVD, have been traced to the expression of certain genes,98–101 although there does not appear to be a clear, universal genetic link between all cases. The specific loci identified thus far include chromosome 11 (11p15.4),102 13 (13q31–32),102 chromosome 16 (16p11–p12),100,102 and the X chromosome (Xq28).99 The underlying genetic defects are not yet known for chromosomes 11, 13, and 16, but there appears to be some linkages to defects in filamin A in certain families with MVP with X-linked inheritance.102

Conclusions and future directions

It has proven challenging to distinguish the effects of normal aging from myxomatous mitral valve disease for both human and canine mitral valves. Human heart valves, both normal and diseased, are becoming more challenging to study for many reasons, such as legislation concerning human subjects protections and the asymptomatic nature of the early stages of valve disease. Canine valves also present a challenge because of the high incidence of the disease. Regardless of these hurdles, it appears that there is a distinct relationship between age and MMVD, although this relationship differs between humans and dogs. This distinction may be due to the focus upon breed-specific studies of MMVD in dogs. Indeed, age does not appear to be the only factor in the development of MMVD; as shown in studies of breed-specific and familial MMVD, there is likely a role for genetic inheritance. In contrast with MMVD, it is unclear whether age is a factor in other diseases that affect dogs, such as the less common myxomatous lesions on the tricuspid and aortic valves103 or in the relationships between heartworm disease, pulmonary hypertension, and mitral valve lesions.53,104 It is evident that significant research remains in addressing the challenges of mitral valve disease in many mammalian species. One of the most promising directions appears to be focusing on the biology of valvular cells from myxomatous and unaffected regions of mitral valves105,106 from dogs of all ages. Other studies have addressed the role of mechanical stimulation on the production of GAGs and PGs by mitral VICs grown from either leaflets or chordae15,107,108 as well as the impact of growth factors on mitral VIC migration and wound healing,109,110 all of which are relevant to MMVD. These in-vitro approaches, together with studies that focus on certain proteins and their signaling pathways, are expected to provide insight into connections between aging and MMVD and to provide genetic linkages to disease mechanisms. Given the purported role of repeated mechanical impacts in initiating and driving myxomatous degeneration, it may be useful to employ in-vitro mechanical stimulation bioreactor approaches to investigate the mechanobiology of these cells and their ability to remodel the ECM. These efforts should reveal new information about the biology of this integral
component of the cardiovascular system and provide critical insight into the development of novel therapies for this devastating disease.

Conflict of interest

Authors have no conflict of interest.

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Aging of the mitral valve


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