Gene network and canonical pathway analysis in canine myxomatous mitral valve disease: A microarray study

Keywords: Myxomatous mitral valve disease; Canine; Transcriptome; Pathway analysis; Microarray; Cavalier king Charles spaniel.

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Abstract: Myxomatous mitral valve disease (MMVD) is the single most common acquired heart disease of the dog and is particularly common in small pedigree breed dogs such as the cavalier king Charles spaniels (CKCS). There are limited data on the mitral valve transcriptome and the aim of this study was to use the microarray technology in conjunction with bioinformatics platforms to analyse transcript changes in MMVD in CKCS compared to normal dogs (non-CKCS). Differentially expressed genes (n = 5397) were identified using cut-off settings of fold change, false discovery rate (FDR) and P < 0.05. In total, 4002 genes were annotated to a specific transcript in the Affymetrix canine database, and after further filtering 591 annotated canine genes were identified: 322 (55%) were up-regulated and 269 (45%) were down-regulated. Canine microRNAs (cfa-miR; n = 59) were also identified.

Gene ontology and network analysis platforms identified between 6 and 10 significantly different biological function clusters from which the following were selected as relevant to MMVD: inflammation, cell movement, cardiovascular development, extracellular matrix organisation and epithelial-to-mesenchymal (EMT) transition. Ingenuity pathway analysis identified three canonical pathways relevant to MMVD: caveolar-mediated endocytosis, remodelling of epithelial adherens junctions, and endothelin-1 signalling. Considering the biological relevance to MMVD, the gene families of importance with significant difference between groups included collagens, ADAMTS peptidases, proteoglycans, matrix metalloproteinases (MMPs) and their inhibitors, basement membrane components, cathepsin S, integrins, tight junction cell adhesion proteins, cadherins, other matrix-associated proteins, and members of the serotonin (5-HT)/transforming growth factor-β signalling pathway.
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Abstract

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Introduction

Myxomatous mitral valve disease (MMVD) is the single most common acquired heart disease of the dog and is characterised by endothelial damage, stromal matrix degeneration, interstitial cell proliferation in the sub-endothelial zone and interstitial cell phenotypic changes (Buchanan, 1977; Beardow and Buchanan, 1993; Corcoran et al., 2004; Black et al., 2005; Disatian et al., 2008; Han et al., 2008; Lacerda et al., 2009; Borgarelli and Buchanan, 2012). The end-stage disease results in significant mitral regurgitation which can lead to left-sided congestive heart failure (Häggström et al., 2009). While much is known about the structural and cellular changes in canine MMVD, less is known about the molecular mechanisms and biochemical changes (Richards et al., 2012). Limited proteomic and transcriptomic data are available and provide interesting insights into disease pathogenesis, for example, the role of serotonin (5-hydroxytryptamine or 5-HT) in MMVD (Oyama and Chittur, 2006; Lacerda et al., 2009).

The only genomics study of canine MMVD to date used the Affymetrix Canine Gene 1.0 array covering approximately 23,000 gene transcripts and looked at a cohort of mixed breed dogs (Oyama and Chittur, 2006). Since that study, there have been significant advances in the quality of canine gene microarrays with much improved transcript annotation (27,681 genes), transcript sensitivity (590,097 probes, 24 probes/gene) and bioinformatics tools. The study by Oyama and Chittur (2006) identified 229 probe sets that were differentially expressed (at least two-fold change; 70% up-regulated and 30% down-regulated) of which 166 could be assigned to recognised genes. The main functional classes attributed to the affected genes, in descending order of importance, included cell signalling, metabolism, extracellular matrix (ECM), inflammation, cell defence, immunity, cell transport and cell structure (Oyama and Chittur, 2006). Of particular note was the ~four-fold increase in the 5-
HT₂B receptor gene, which fits in with the serotonin hypothesis of MMVD pathogenesis.

Limited numbers of gene expression studies have also been undertaken in human MMVD with a reported number of differentially expressed genes of around 400 (Hulin et al., 2012; Sainger et al., 2012). These studies demonstrated informative changes in metallothioneins and ADAMTS (A Disintegrin and Metalloproteinase with Thrombospondin Motifs), and transforming growth factor β-2 (TGFβ-2) and bone morphogenetic protein 4 (BMP-4 of the TGF-β signalling superfamily), proteoglycans, collagens, SOX-9 and CRTAC1 (cartilage acidic protein 1), genes that are implicated in ECM production and remodelling.

However, the number of reported differentially expressed genes in any transcriptomic study is affected by the calculated false discovery rates (FDRs), fold change thresholds (correction for signal intensity) and batch effects. Additionally, there have been major developments in bioinformatics, canine annotated databases and analysis platforms. The aim of this study, therefore, was to use more stringent threshold criteria and quality control, in line with current recommended protocols and with an affected group consisting of the same breed (Cavalier King Charles spaniel, CKCS) in order to improve the reported differential expression of genes in canine MMVD, and to identify novel signalling pathways that might contribute to the pathogenesis of MMVD.

Materials and methods

Tissue sample

Myxomatous mitral valve leaflets (anterior) (n = 10, Whitney grade ≥ 3) were collected from CKCS dogs presented to the Hospital for Small Animals, Royal (Dick) School of Veterinary Studies, the University of Edinburgh. All dogs previously had been clinically confirmed to have MMVD and were on a range of treatments for congestive heart failure.
Control normal anterior mitral valve leaflets \((n = 6)\) were collected from young adult dogs euthanased for reasons other than cardiac disease. All samples were collected within 10 min of death and Whitney classified by at least two investigators (CCL, MML, BMC). The valve leaflets were immediately placed in RNAlater (Invitrogen) and stored at -20 °C. All tissue samples were collected with full owner informed consent and the study conformed to national (UK) and institutional ethical guidelines for the use of animals in research.

RNA extraction and quality assessment

Tissue (100 mg) was minced, snap-frozen in liquid nitrogen and pulverised for 1 min at 2000 oscillations/min in a liquid nitrogen cooled dismembranator (Braun Mikro-Dismembrator Vessel, Braun Biotech International). Phenol/guanidine HCl reagents (1 mL; TriReagent, Sigma) was added to the powdered sample and 200 µL of chloroform was added to the microcentrifuge tubes prior to centrifugation at 12,000 g for 10 min. RNA extraction from the aqueous phase and DNA digestions were carried out using commercially available kits (RNeasy Mini Kit and RNase-Free DNase Set; Qiagen). RNA purity was analysed by spectrophotometer (Thermo Scientific NanoDrop 1000, 260/280 ratio \(\approx 2\)) and RNA quality and integrity and RNA integrity number (RIN \(\geq 7\)) determined by electropherogram (Agilent 2100 Bioanlyser). Four normal and four diseased samples reached the required quantity and quality of RNA and were stored at -80 °C prior to reverse transcription.

Microarray hybridisation

The Affymetrix GeneChip WT Terminal Labelling Kit and the Ambion WT Expression Kit were used to generate amplified and biotinylated sense-strand DNA targets for the Affymetrix Canine Gene 1.0 ST Array. Each ST (Sense Target) array selected probes were distributed along the entire length of each transcript. Briefly, total RNA was reverse-
transcribed to single-stranded cDNA, and then converted to double-stranded cDNA. The double-stranded cDNA then underwent in vitro transcription to generate unlabelled cRNA. The cRNA was cleaned for any antisense RNA, reverse-transcribed to the dUTP incorporated second single-stranded cDNA and fragmented with uracil DNA glycosylase and apurinic/apyrimidinic endonuclease 1. The fragmented DNA was labelled by terminal deoxynucleotidyl transferase (TdT) with the Affymetrix proprietary DNA Labeling Reagent, covalently linked to biotin, and then hybridised to the array. The arrays were washed, stained and scanned. The two cycle RNA-cDNA amplification, hybridisation, and chip scanning were performed by ARK Genomics (Roslin Institute, UK).

For differential gene expression analysis, all the raw Affymetrix CEL files, which contained probe-sets ID and calculated pixel intensity value for each array, were imported into Partek Genomic Suite 6.6 (PGS). Normalized signal intensity data for whole probe sets were also produced by Affymetrix Expression Console. Robust Multi-array Average (RMA) was used for data normalisation and final summarisation. QC metrics combined with principal component analysis (PCA) were generated for post-import quality assessment, and two potential outliers were identified and excluded. Analysis of variance (ANOVA) was used to identify differentially expressed genes between the two groups. Hierarchical clustering was based on the significant genes, and the original gene list was used to generate a Volcano plot (Fig. 1) for total unadjusted gene expression pattern. Finally, a list of genes of interest was generated according to the cut-off of fold-changes, FDR, and signal intensity.

Multi-platform in silico categorisation and network analysis

Gene categorisation and pathway analysis were conducted using a combination of ToppFun (gene list enrichment tool), DAVID 6.7 and the Ingenuity Pathways Analysis (IPA)
database by uploading the differentially expressed gene list to the respective online servers.

Functional analysis of a gene network identified the biological functions that were most attributable to the genes in that network (Chen et al., 2009; Huang et al., 2009).

Quantitative reverse-transcription polymerase chain reaction (RT-PCR)

Assays for 13 canine genes and one reference gene (Table 1) were designed using methods previously reported (Clements et al., 2006). Primers (MWG Biotech) and probes (Roche Diagnostics) were synthesised using locked nucleic acid with 5′-end labelled with a reporter fluorescein dye (FAM, 6-carboxy fluorescein) and 3′-end labelled with a dark quencher dye. In brief, quantitative RT-PCR assays were performed in triplicate in 96-well plates (LightCycler 480 system using Probes Masters; Roche Diagnostics) with one no-template control for each sample with total reaction volume of 5 μL per well. The amplification was performed according to a standard protocol. Quantitative RT-PCR data were analysed using LightCycler 480 Basic Software (Roche Diagnostics). The reference genes was used to normalise target gene relative expression level and to calculate the -ΔΔCt values.

Results

General description

RNA of sufficient quality and quantity for transcriptomic analysis was obtained from four dogs from each group (RIN > 7). There was one outlier in each group and when these were excluded 5397 differentially expressed genes were identified fulfilling the cut-off settings of fold change (> 1.5), FDR <0.05, and P < 0.05. In total, 4002 genes were annotated and 1395 probes were not assigned to a specific transcript in the Affymetrix canine database and their sequences were tracked and matched using the NCBI nucleotide BLAST and
miRBase BLAST tools. Only 139 probes were successfully identified as transcript homologs to other species and 59 of these were matched with canine microRNA family members (cfa-mir-RNA) (details to be included in a further report). The gene and the data sets were further filtered by individual signal intensity value and a final total of 635 differentially expressed probe sets, representing 591 annotated canine genes, were identified, 322 (55%) were up-regulated and 269 (45%) genes were down-regulated (Appendix; Supplemental file 1).

**Quantitative RT-PCR**

For data validation between different platforms, differential expression of 13 genes were examined and 12 (LAMA2, ENG, COL6A3, HTR2B, MYH11, MMP12, BMP6, ANGPT1, CHAD, ADAMTS19, ACTG2, and KERA) were found to be in agreement with the microarray data, while one (SOX-9) was not (Appendix; Supplemental file 2).

**Gene ontology, and network and canonical pathways**

Using ToppFun, 28 significant terms for gene ontology (GO) and biological processes (BP) were identified. Six functional subgroups were further identified based on the biological functions of each gene ontology term (GO: BP), and included inflammation and immune response, cellular adhesion and movement, cardiovascular development, ECM, osteogenesis, and epithelial cell proliferation. In each subgroup, GO: BP terms deemed biologically relevant to MMVD were selected and identified genes that are involved in response to wounding, biological adhesion, positive regulation of cell migration, cardiovascular system development, ECM organization, ossification, and regulation of epithelial cell proliferation (Appendix; Supplemental file 3).
DAVID identified 16 biological categories based on gene functions (Table 2).

Functional annotation clustering analysis after data enrichment identified 10 clusters based around similar biological function (Table 3) and the specific named genes are in Supplementary file 4 (Appendix). Analysis using IPA identified 11 significant biological functions and diseases relevant to the gene set, including endocrine system disorders, cellular movement, connective tissue disorders, inflammatory disease, immune cell trafficking, cancer, haematological system development, cardiovascular disease, humoral immune response, cardiovascular system development and cell death and survival. IPA also identified 33 canonical pathways with three functions applicable to MMVD, including caveolar-mediated endocytosis signalling, remodelling of epithelial adherens junctions, and endothelin-1 signalling, were identified (Appendix; Supplementary file 5). Gene networking analysis identified three significant and biologically relevant networks for MMVD, including cardiovascular system development and function, cellular movement and cell-to-cell signalling (Fig. 2). By overlaying these data with the disease-functions-canonical pathway analysis, specific genes of interest were identified (Table 4). Analysis by IPA also identified the top 10 upstream regulators: IL-1B, IL-13, TGFβ1, DYSF, IFN-γ, TNF, LDL, CSF2, NFκB, and NFE2L2. Of these, LDL activation appears to be the most directly relevant to MMVD.

By combining analysis from the three platforms, the following relevant functional and disease categories can be derived for the MMVD transcriptome: inflammatory response, cell movement, cardiovascular development, ECM organisation and epithelial-to-mesenchymal transition.

*Individual gene signal intensity*
Using the data from annotation clustering analysis, genes deemed biologically relevant to MMVD from the five selected categories were examined on the basis of their signal intensity and the details are shown in Table 5. The gene families of particular interest were: collagens, ADAMTS peptidases, proteoglycans, matrix metalloproteinases (MMPs) and their inhibitors (TIMPs), basement membrane components (nidogen1 and laminin2), cathepsin S, integrins, tight junction cell adhesion proteins (claudin, occludin), cadherins, and other matrix-associated proteins (fibrillin, fibronectin, periostin, fibulin, HAS2). Other genes that showed reasonably high expression, but were not necessarily significantly changed between the two groups, included: elastin, NOS, adhesion molecules (PECAM, VCAM), NOTCH, Snai1, β-catenin, members of the TGF-β superfamily (TGFβ-1, endoglin, BMP9, BMPR1B) and cytoskeleton proteins (actin, myosin, SM22).

Discussion

Transcriptional profiling for this study was carried out using the Affymetrix Canine Gene 1.0 ST Array. In total, 590,097 probes against 27,681 genes (both annotated and predicted canine genes) were included according to the genome information from canFam2. The design of 26 unique 25-mer probes for each transcript allowed for the highest coverage yet and provided a more accurate detection of transcribed genes and a higher resolution than 3’-biased microarrays. The only previous report of the canine mitral valve transcriptome used the first generation array Affymetrix GeneChip Canine Genome 1.0 (Oyama and Chittur, 2006). The current study confirmed the previous findings of up-regulation of 5-HTR2B, endoglin and BMP-6, but did not find differences in the expression of TGF-β or TGF-β receptors.
In human MMVD, the two published transcriptomic studies also reveal an orchestrated regulation of TGF-β and BMP-6, and in functional studies, control of ECM production and valve interstitial cell activation by BMP-4 (Hulin et al., 2012; Sainger et al., 2012). However, there are no reports of up-regulation of 5-HTR2B in any of the human valve studies. The findings of interest in the human mitral valve transcriptome are down-regulation of metallothioneins-1 and -2 (MT1/2) and members of the ADAMTS family of proteases, and increased expression of genes encoding for ECM components, including collagens, proteoglycans and MMPs. In contrast, in the dog, ECM genes were generally down-regulated or unchanged. While canine and human MMVD share many similar features, the diseases appear to be different at least at the transcriptomic level.

The possible involvement of the TGF-β signalling superfamily in MMVD is not surprising as it has important roles in cancer, fibrosis, and calcification (Geirsson et al., 2012). TGF-β signalling through SMAD pathways triggers myofibroblastic differentiation of valve interstitial cells and increases expression of SMAD-targeted genes associated with ECM such as COL1A, 3A1, 6A1, 6A3, elastin and TIMP1 (Verrecchia et al., 2001; Walker et al., 2004). However, there was no differential expression of the TGF-βs and their receptors or downstream target genes between the normal and affected dogs, although high signal intensity for various members indicated their role in valve matrix homeostasis. Increased TGFβ-1 and TGFβ-3, but not TGFβ-2, expression has been shown using immunohistochemistry in canine valves (Aupperle et al., 2008). These differences between gene and protein expression may be due to post-transcriptional modification. This contrasts with human myxomatous mitral valves where there is higher transcriptional level of all isoforms of TGF-β and the downstream genes COL1A1, COL3A1 and elastin, suggesting activation of fibrotic mechanisms (Geirsson et al., 2012).
The identification of LDL activation and its positive association with 17/23 downstream genes using IPA was an unexpected but interesting finding for MMVD. LDL signalling through low-density lipoprotein receptor-related protein-5 (Lrp5) has been shown to play a crucial role in calcific degeneration in human aortic and mitral valves (Neufeld et al. 2014). There is an association between LDL and aging in dogs which is coincidental with the natural history of MMVD, and LDL is worth considering as a potential contributing factor to MMVD in this species (Buchanan, 1977; Beardow and Buchanan, 1993; Osorio, 2009; Borgarelli and Buchanan, 2012).

Increased expression of inflammation-associated cytokine genes in the diseased mitral valve appears to be a consistent finding, but is not reported in the human mitral valve (Oyama and Chittur, 2006; Hulin et al., 2012; Sanger et al., 2012). IPA identified, central to the inflammation network, up-regulation of toll-like receptor 4 (TLR4) and interleukin 18 (IL-18), as well as IL-6, TLR1 and TLR8. Since there is no evidence of inflammatory cell contribution to MMVD pathogenesis, ECM degeneration and remodelling might be triggering TLR receptor signalling which could further contribute to ECM changes. Heat shock protein 70, fibronectin, hyaluronic acid, heparan sulfate, and hyaluronan, all important components of ECM remodelling in MMVD, can act as endogenous ligands for TLRs (Chao, 2009). Finally, IL-6 can trigger endothelial-mesenchymal transformation (EndoMT), an important mechanism during valve development, and potential contributor to the pathogenesis of MMVD (Mahler et al., 2013).

Matrix gene expression changes have been found in canine and human MMVD, and included genes encoding for ECM and basement membrane (BM) proteins. The BM is important for maintaining endothelial integrity, and endothelial damage, activation and cell
loss are features of canine MMVD (Corcoran et al., 2004; Han et al., 2013). Gene network analysis identified down-regulation of laminin beta 1 (LAMB1) and alpha 2 (LAMA2), nidogen-1 (NID1) and COL6A3, and increased expression of the protease cathepsin S (CTSS), all of which would reflect endothelial damage or BM dismantling as part of EndoMT (Lakatta and Levy, 2003; Li and Bertram, 2010). Breakdown of NID1, LAMB, COL and elastin by CTSS has been shown to impair BM integrity and stability (Sage et al., 2012; Turk et al., 2012). Increased expression of CTSK and CTSS has been reported in human MMVD, and cyclic strain increases CTSK expression in sheep mitral valve myofibroblasts (Rabkin et al., 2001; Aikawa et al., 2006; Lacerda et al., 2012).

Regarding ECM proteins, there were not surprisingly high intensity signals for many collagen genes and these tended to be lower in the disease group (COL1, COL2, and COL4), but only statistically different for COL6A3. There is a marginal and localised reduction in collagen expression in MMVD, at least in mild to moderately affected dogs, and this would be reflected by these gene expression changes (Hadian et al., 2010). The reduction in COL6A3 is important because its role in BM production and force-resistant collagen bundle formation suggests a clear contribution to the pathogenesis of MMVD (Klewer et al., 1998; Kruithof et al., 2007). Collagen maturation from procollagen relies on the ADAMTS family of metalloproteases, and ADAMTS2, ADAMTS9, and ADAMTSL4 were all significantly down-regulated in the diseased canine mitral valves. Lower expression of ADAMTS2 and ADAMTS9 suggests an inactive collagen turnover state exists, and this has been shown in ADAMST9-deficient mice, and in canine MMVD using X-ray diffraction and HPLC (Hadian et al., 2010; Kern et al., 2010). ADAMTS4 also has a regulatory role on fibrillin-1 and low expression will decrease elastin fibre formation in MMVD in a manner similar to that seen in Marfan syndrome (Chandra et al., 2012).
Gene expression for a limited number of MMPs and their tissue inhibitors (TIMPs) has been previously reported using a combination of microarray and PCR, and expression at the protein level has also been reported using immunohistochemistry. In the current study, 25 MMPs and 4 TIMPs were identified, but in general the data contradicted previous reports in particular for MMP1, which had a low intensity signal, but has been previously reported as increased using PCR and immunohistochemistry (Oyama and Chittur, 2006; Disatian et al., 2008; Aupperle et al., 2009, 2012). Remodelling activity at the time of sampling are likely to have effects on global MMPs and TIMP expression profiles and changes in expression need to be interpreted with caution (Rabkin et al., 2001; Rabkin-Aikawa et al., 2004).

Overall gene ontology analysis for all studies reported to date show similar biological functions in the differentially expressed gene sets, including genes involved in cell signalling, inflammation, extracellular matrix, immunity, cell defence, and metabolism. In the current study, additional functional categories of cellular movement and epithelial-to-mesenchymal transition were identified. Three canonical pathways were selected that would appear most relevant to the pathogenesis of MMVD: caveolae-mediated endocytosis, which controls endothelial cell growth, cell migration and can affect TGF-β signalling-induced fibroblast activation; endothelin signalling, which has a variety of functions including ECM remodelling; and remodelling of epithelial adherens junction, which with vascular endothelial (VE)-cadherin can affect cell proliferation and migration (Mow and Pedersen, 1999; Salanueva et al., 2007; Galdo and Lisanti, 2008; Sowa, 2012).

For all ECM products, proteoglycan (PG) genes had the highest signal intensity with lumican, versican, and biglycan being the three strongest, but only chondroadherin (CHAD)
and keratocan (*KERA*) were significantly down-regulated in the CKCSs. *KERA* is a small leucine rich PG and plays a pivotal role in ECM assembly in the cornea to maintain translucency, and CHAD is an anchor to the matrix by binding tightly to collagens I, II, and VI in cartilage (Liu et al., 2003; Hessle et al., 2013). Nevertheless, the role of CHAD and KERA in the mitral valve is unknown. In a canine MMVD proteomic study, decorin and biglycan were found to be up-regulated in the early-stage MMVD, but down-regulated in the late-stage MMVD (Lacerda et al., 2009). In contrast in human MMVD, biglycan (protein), decorin (both mRNA and protein), and versican (protein) were found more abundantly expressed compared with normal mitral valves (Radermecker et al., 2003; Gupta et al., 2009). In our study, unchanged expression of the major PGs (e.g. lumican, versican, decorin, and biglycan) between diseased and normal mitral valves suggests that cellular and structural changes in end-stage MMVD had no direct effect on PG gene expression. The variation of PG expression in different studies may suggest post-transcriptional and translational modification of PG mRNA and proteins.

There were no changes in the endothelium adhesion molecules *VCAM-1, ICAM1* or *PECAM1*, but there was over-expression of *E-selectin* and *TLRs*. Strong staining for *E-selectin* and *VCAM-1* in human myxomatous mitral valves, without morphological evidence of inflammation, has been previously reported. (Müller et al., 2000) The closure and opening of the cell-cell endothelial adherens junctions, cell motility, and maintaining vascular permeability and integrity is controlled by VE-cadherin (CDH5), and there was high signal intensity for a range of CDHs (2, 11 and 13) in the microarray (Dejana and Orsenigo, 2013). High expression of the cadherins promotes cell migration and proliferation through the ERK1/2 pathway (CDH13), and mesenchymal cell (CDH11) and myofibroblast differentiation (CDH2) (Ivanov et al., 2004). *CDH5* had the lowest signal intensity and was
significantly reduced in the CKCS group, and down-regulation of CDH5 through NOTCH-
Snai1 or TGF-β signalling permits endothelial migration and EndoMT (Armstrong and
Bischoff, 2004). Down-regulation of CDH5 was matched by up-regulation of Snai1 in
affected valves, but NOTCH was also down-regulated, suggesting the possible presence of a
NOTCH-independent Snai1 signalling pathway in canine MMVD. The pattern of cadherins
expression also suggests a proliferative and migratory phenotype, with minimal osteogenic
activity, and the contribution of these adhesion proteins to disease pathogenesis needs further
investigation.

Certain expression data suggested the presence of EndoMT, such as differential
expression of genes associated with BM components, mesenchymal differentiation and
NOTCH signalling pathways. There is evidence of transition into a mesenchymal phenotype
in MMVD, represented by increased expression of mesenchymal markers such as α-smooth
muscle actin (α-SMA, ACTA2), SM22 (TAGLN), and γ-SMA (ACTG2). This mesenchymal-
transition could be initiated by the up-regulation of the TGF-β superfamily members BMP6
and BMPR1, increased NOTCH-Snai1 signalling, down-regulating the expression of CDH5,
decreased expression of the BM components NID1 and LAMA2, and increased expression of
the BM lytic enzyme CTSS. Lastly, the increased expression of hyaluronic acid synthase 2
(Has2) in the CKCS could provide the hyaluronic acid rich subendothelial matrix necessary
for transition of endothelial cells (Bakkers et al., 2004; Camenisch et al., 2001; Lagendijk et
al., 2013). These changes in the valve transcriptome are reminiscent of the changes needed in
valve development to allow the valve to form from the endocardial cushion through the
migration of endothelial cells into the hyaluronic-rich embryonic stroma, where they
differentiate into valve interstitial cells which then generate the valve matrix (Camenisch et
The data from the current study suggest that EndoMT contributes to MMVD pathogenesis and this area needs further study.

The main limitations of this study were the small sample size, the lack of age- and breed-matched controls, the extraction of RNA from a mixed tissue type, sampling at a single disease end-point and differences in in silico analysis platforms. The difficulties in getting sufficient RNA of adequate quality from valves are illustrated by rejection of 6/10 CKCS samples, but the statistical analysis confirmed three vs. three was sufficient for credible comparison. MMVD research is hampered by the lack of suitable age- and breed-matched controls, because of the ubiquity of the disease in aged dogs and the limited sample pool when having to use family pets. This limitation cannot be overcome, but at least in this study the test group were all from the same breed reducing variability to some extent. In silico analysis show differences among different platforms and network clustering and gene-gene interaction analysis are based on several different tissue and cell types and may not necessarily be directly applicable to the mitral valve. However the use of three platforms resulted in somewhat consistent conclusions as to the major gene categories in MMVD and identified two novel categories not previously reported. Network analysis did allow for the identification of potential pathogenesis pathways that are worthy of further investigation. Furthermore, with continual updates of annotations and increased accuracy of canine genomics databases, data set reanalysis will be possible in the future.

Conclusions

The characterization of the MMVD transcriptome identified differentially expressed genes associated with inflammation, cell movement, development, and extracellular matrix organization and EndoMT. Signal intensity analysis identified genes important in ECM,
EndoMT and valve development, with patterns of gene expression suggesting decreased collagen turnover, ECM weakening, BM disruption, increased cell migration, active endothelial and myofibroblast differentiation in MMVD affected dogs.

Acknowledgements
This study was funded in part by the Cavalier Kings Charles Club of England. Chien Lu was in receipt of a Charles Darwin Scholarship from the University of Edinburgh. Meng-Meng Liu was in receipt of a Chinese Government State Scholarship.

Appendix .Supplementary material
Supplementary data associated with this article can be found in the online version, at doi

Conflict of interest statement
None of the authors of this paper has a financial or personal relationship with other people or organizations that could inappropriately influence or bias the content of the paper.

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Table 1. Primer Sequences for Microarray Data Validation by Q-PCR are shown.

| Gene Name | Forward Primer | Reverse Primer | Probe Number |
|-----------|----------------|----------------|--------------|
| **LAMA2** | 5’CCGCATTGAGCTGACAGT 3’ | 5’CAAACTGGTGGAGCCATCT 3’ | 38 |
| **ENG**   | 5’GGTGCTCAAGAAAGACCT CATC3’ | 5’GCAGGACAACTGGTCATCT C3’ | 30 |
| **COL6A3**| 5’AAGGAAGCTTCAGCACAAG AGA3’ | 5’TGAATAGAAGCCAACCTTG C3’ | 39 |
| **HTR2B** | 5’GGTCTGGAGATTACAAACAG AATCG3’ | 5’TCCCTGTTGCTCACCAGTCTC3’ | 6 |
| **MYH11** | 5’ATGCAGCTGGCCAAGAG AGAG3’ | 5’TCTGAGCAATTTCATCATCA G3’ | 81 |
| **MMP12** | 5’CCAATTGGATTTTGTGAGCTG TGT3’ | 5’CACTGTCTTTTGGACTCTC TG GA3’ | 22 |
| **BMP6**  | 5’TCTCCAGCCCTCAGATATTACCTGA3’ | 5’TGGAAGCTCATACGACTCTC A3’ | 4 |
| **ANGPT2**| 5’AGGAAACGAAAAAGCAGAC ACTACA3’ | 5’ATCAGCACCATGTAAGATCA GG3’ | 18 |
| **CHAD**  | 5’CCAGTCTTTTCGGCAGTAC C3’ | 5’ACATGTCTTTGCTGCGTCAC G3’ | 20 |
| **ADAMTS19** | 5’TCAACCCCTGCAATGAGAA GA3’ | 5’CGTATCACTGGCAGTACAC A3’ | 14 |
| **ACTG2** | 5’GGTCATCACCATTGAGCAA C3’ | 5’TGAATCCCGAGCAGACTCCAT C3’ | 11 |
| **Sox-9** | 5’CCACCGCCATCTTCAAGG3’ | 5’GGAGTGCACCTGTCATC3’ | 63 |
| **KERA**  | 5’GACTATGCACTTTTGACT GTCC3’ | 5’TTTCACAGTATAAAGACGAGTA GGGAAA3’ | 29 |
| **MRPS25**| 5’TCTTGGGGAAGAACAAGG AA3’ | 5’AGTGGGGCTGGTGAGAAAG3’ | 15 |
Table 2. Gene ontology chart showing the differentially expressed genes in canine mitral valves using the gene analysis platform DAVID 6.7.

| Gene Ontology Term                                | Gene Counts | Gene % | p-value         |
|---------------------------------------------------|-------------|--------|-----------------|
| Response to wounding                              | 43          | 8.5    | 2.10E-08        |
| Inflammatory response                             | 31          | 6.2    | 9.50E-08        |
| Immune response                                   | 42          | 8.3    | 4.70E-05        |
| Cell adhesion                                     | 41          | 8.1    | 1.40E-04        |
| Biological adhesion                               | 41          | 8.1    | 1.40E-04        |
| Defence response                                  | 37          | 7.3    | 1.80E-04        |
| Vasculature development                           | 20          | 4      | 3.10E-04        |
| Positive regulation of cytoskeleton organization   | 8           | 1.6    | 4.30E-04        |
| Blood vessel development                          | 19          | 3.8    | 6.30E-04        |
| Regulation of cell proliferation                   | 42          | 8.3    | 7.50E-04        |
| Blood vessel morphogenesis                        | 17          | 3.4    | 8.90E-04        |
| Regulation of cytoskeleton organization            | 13          | 2.6    | 1.10E-03        |
| Innate immune response                            | 13          | 2.6    | 1.20E-03        |
| Response to oxygen levels                         | 13          | 2.6    | 1.40E-03        |
| Collagen metabolic process                        | 6           | 1.2    | 1.50E-03        |
| Epithelial to mesenchymal transition              | 5           | 1      | 2.00E-03        |
Table 3.

Functional annotation clustering using DAVID 6.7, listing the ten clusters and the relative gene density.

| Annotation Clusters                              | Enrichment Score | Gene Count | p-value    |
|-------------------------------------------------|------------------|------------|------------|
| Vasculature development                         | 3.25             | 20         | 3.10E-04   |
| Collagen metabolic process                      | 2.57             | 6          | 1.50E-03   |
| Protein processing                              | 2.41             | 11         | 2.50E-03   |
| Vacuole                                         | 2.36             | 18         | 2.20E-03   |
| Positive regulation of cytoskeleton organization| 2.24             | 10         | 4.30E-04   |
| Regulation of cell motion                       | 1.91             | 14         | 7.10E-03   |
| Regulation of bone mineralization               | 1.71             | 5          | 1.30E-02   |
| Positive regulation of cell motion              | 1.71             | 9          | 1.10E-02   |
| Cysteine-type endopeptidase activity            | 1.61             | 9          | 5.40E-03   |
| Epithelial to mesenchymal transition            | 1.52             | 5          | 2.00E-03   |
Table 4.
Gene networking analysis using the gene analysis platform IPA.

A. Cardiovascular system development

| Diseases and functions                        | Genes                      |
|-----------------------------------------------|----------------------------|
| Abnormal morphology of mitral valve           | NFATC1                     |
| Abnormal morphology of cardiovascular system  | NFATC1, PPDX1, RBPJ, NOTCH1,ZMIZ1 |
| Abnormal morphology of cardiac valve          | NFATC1, ZMIZ1              |
| Morphogenesis of endocardium                  | RBPJ                       |
| Morphogenesis of atrioventricular valve       | NOTCH1                     |
| Morphogenesis of heart                        | NOTCH1, RBPJ, ZMIZ1        |
| Looping morphogenesis of heart                | NOTCH1, RBPJ               |
| Differentiation of endocardial cells          | NFATC1                     |
| Differentiation of endothelial cells          | NFATC1, RBPJ, NOTCH1       |
| Development of mesenchymal cells              | NOTCH1                     |

Canonical Pathway

| Genes                          |
|--------------------------------|
| Regulation of the epithelial-mesenchymal transition pathway |
| NOTCH1, CSL-HIF-1A            |
| NOTCH signalling              |
| Secretase γ, RBPJ, NFκB1, NOTCH1, CSL-HIF-1A |

B. Cellular movement and connective tissue development

| Diseases and functions                        | Gene                      |
|-----------------------------------------------|---------------------------|
| Cell movement of fibroblast                   | SKAP2, WASF2, ANGPT1, ENPP2 |
| Familial thoracic aortic aneurism             | MYH11                      |
| Endothelial cell development                  | ANGPTL1, ANGPT1           |
| Cell movement of endothelial cells            | ANGPT1, ENPP2             |
| Injury of endothelial cells                   | ANGPT1                     |

Canonical Pathway

| Gene                      |
|---------------------------|
| Integrin Signalling       |
| Arp2/3, Talin, calpain, Akt, integrin, JINK1/2, Lfa-1 |

C. Cell-to-cell signalling and tissue development

| Diseases and functions                        | Gene                      |
|-----------------------------------------------|---------------------------|
| Adhesion of fibroblast cell lines             | TNMD, TENC1               |
| Adhesion of vascular endothelial cells        | ITGB1, SELE               |
| Association of extracellular matrix           | ITGB1                     |
| Attachment of smooth muscle cells             | ITGB1                     |
| Binding of vascular endothelial cells         | ITGB1, SELE               |
| Morphogenesis of endothelial tube             | PDPN                      |
| Basement membrane disruption                  | NID1, Laminin             |
Table 5. Examples of single gene intensity changes are shown.

| Category                  | Genes                                                  |
|---------------------------|--------------------------------------------------------|
| Collagen                  | *COL6A3*↓(H)                                           |
| ADAMTS                    | *ADAMTS2*↓(M), *ADAMTS19*↓(M), *ADAMTS4*↓(M)          |
| PGs and GAGs              | *KERA*↓(H), *CHAD*↓(H)                                 |
| MMP and TIMP              | *MMP12*↑(M), *MMP14*↓(M), *MMP16*↓(M)                 |
| Basement membrane         | *NID1*↓(H), *LAMA2*↓(L), *CTSS*↑(H)                    |
| Cathepsin                 | *CTSC*↑(M), *CTSS*↑(H)                                 |
| Integrin                  | *ITFG1*↑(H), *ITGA*(M), *ITGA8*↑(M), *ITGB1*↑(H), *ITGB4*↓(M), *ITGBL1*↑(M) |
| Claudin and occludin      | *CLDN1*↑(M), *CLDN11*↓(H)                             |
| Cadherin                  | *CDH5*↓(H)                                            |
| Others                    | *CILP*↓(H), *Has2*↑(M), *HAPLN1*↓(H)                   |
| Caveolin, PECAM, ICAM, SELE, and VCAM | *SELE*↑(L)                                      |
| NOTCH and SNAI1           | *NOTCH1*↓(L), *RBPI*↑(H)                              |
| Catenin, VEGF, and NFATc  | *NFATc1*↓(M)                                          |
| TGF-β and superfamily     | *ENG*↓(H), *BMPRI*↑(L), *BMP6*↑(M)                    |
| Actin, myosin, and SM22   | *ACTA2*↑(H), *ACTC1*↓(H), *ACTG2*↑(H), *TAGLN*↑(H)    |

Statistically significantly different genes comparing cavalier king Charles spaniels (CKCSs) and normal dogs, selected on the basis of their likely biological relevance to myxomatous mitral valvular disease (MMVD), and using the five selected categories derived from annotation clustering analysis (ToppFunn, DAVID and IPA). Genes were also examined on
the basis of their signal intensity; H, high signal intensity; M, medium signal intensity; L, low signal intensity. ↑ up-regulated, ↓ down-regulated.
Figures Legends

Figure 1. Volcano plot demonstrating the overall gene expression pattern based on the X-axis (fold-change value) and Y-axis ($P$). Each dot represents one gene. Two vertical (fold-change value at 1.5 and -1.5) cut-off lines and one horizontal ($P = 0.05$) cut-off line are shown. In general, the plot is equally distributed. The significantly down-regulated genes in zone 3 were more diffusely distributed compared with the significantly up-regulated genes in zone 4. Unchanged genes in zone 1 ($P > 0.05$) and zone 2 ($P < 0.05$) had the highest plot intensity.

Figure 2. Gene networking analysis using Ingenuity pathway analysis (IPA) identified three significant and biologically relevant networks for myxomatous mitral valvular disease (MMVD), including network (1) cardiovascular system development and function, network (17) cellular movement and Network 23) cell-to-cell signalling. Green is down-regulated and red is up-regulated, with fold changes shown beneath each gene.
Volcano plot CKCS vs control

P-value CKCS vs control

Fold – change CKCS vs control
Figure 2A

Click here to download high resolution image
Figure 2C

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Supplemental file 1
Click here to download Optional e-only supplementary files: Supplemental File 1 Gene List.docx
Supplemental file2
Click here to download Optional e-only supplementary files: Supplemental File 2 Q-PCR validation.docx
Click here to download Optional e-only supplementary files: Supplementary File 5 IPA.docx
Manuscript Highlights

“Gene network and canonical pathway analysis in canine myxomatous mitral valve disease: a microarray study”

1. Most complete coverage of the mitral valve transcriptome, several orders of magnitude better than then only other report from 2006.

2. Application of several gene analysis platforms giving greater levels of data analysis and gene categorisation relevant to disease pathogenesis.

3. Evidence of contribution of endothelial-to-mesenchymal transition to disease pathogenesis; one of the more novel theories of disease to emerge in recent years.

4. Identification of ~600 genes significantly implicated in MMVD giving a picture not only of the complexity of the disease, but important information that will allow hypothesis driven research to be applied to understanding this disease.
Reviewer #1:
This is a well-written and interesting study of gene expression analysis in mitral valve tissue from dogs with myxomatous mitral valve disease. The aim of this study was to make an improve analysis on gene expression compared to previous studies in canine MMVD, in order to identify novel signaling pathways that might contribute to MMVD.

The study has a number of limitations, and most of them are addressed in the limitations. The major limitation is that finding of a difference between MMVD dogs and control dogs may be due to breed differences because the MMVD dogs only include tissue from Cavalier King Charles Spaniels. Previous studies have shown breed differences, for example with regards to platelet aggregation response and circulating biomarkers (Olsen et al. 2001; Moesgaard et al. 2007).

This a valid comment and we will make sure it is mentioned in the limitations. The intention here had been to limit variability by concentrating on this one breed. In fact we have evidence that the cellular changes seen in CKCSs are the same as for other affected dogs, so might argue the CKCS is a good
model of the disease in dogs and the propensity for this breed to be affected is primarily a function of the time to onset and not the pathogenesis (although that need one very large study to confirm).

In addition, it is limitation that histopathology is not performed to confirm the diagnosis. I think this is important also to address these two limitations.

Histopathological confirmation was undertaken. Have included a short comment in Materials and methods

In general, more references in relation to the gene analysis would be appropriate.

We would prefer to leave as is Chen and Huang refer to DAVID and TOPPFUN and were the developers, and IPA is a commercial product for which there is no specific development reference.

Line 95: Whitney grade for control dogs are missing

Grade is zero; have amended

Line 98: How was MMVD clinically confirmed?

All cases had been clinically confirmed by ourselves using echocardiography; have commented in MS.

Line 99: Information of therapy is missing

All dogs had been in the terminal stages of the disease and would have been on standard treatment for congestive heart failure. There might have been subtle differences in types of medication and dosage, but there is no way we could control for those variables; have commented in MS.

Line 142: Was model control for the AVOVA performed

Parktek 6.6 software only runs ANOVA and so we used that to compare 4 vs 4. We are not too sure what is meant by “model control” in this context.

Line 165: Description of the statistic in more detail is missing

See above; since these programmes and platform as have their own in-built statistical methodologies, which we presume have been properly modelled we are not sure of the benefit of providing that detail in the MS. For qRT-PCR Mann-Whitney U test was used with Graphpad prism.

Reviewer #2: Gene Network and Canonical Pathway Analysis in Canine Myxomatous Mitral Valve Disease: A Microarray Study

In this manuscript the authors used microarray data and several analysis tools to better understand the mechanism and affected genes/pathways in Cavalier King Charles Spaniels with Myxomatous Mitral Valve Disease. These data use the newer and better annotated canine Affymetrix array.

I am accepting this paper but have several comments the authors should address:

General comments:

Capitalize Cavalier King Charles Spaniel throughout the entire manuscript

This would be incorrect as cavalier and spaniel are not proper nouns, but King and Charles are, so they are capitalised. For the acronym we think capitalisation is the right form hence “CKCS”

Correct spelling errors throughout the manuscript

Done

Abstract

When mentioning the changed gene families please mention the general direction of change (up or down regulated)

Can we have editorial advice as doing this for each gene might make the abstract look “messy”. We have regrouped the genes as “all down-regulated”, “all up-regulated” or “up and down-regulated”.


See amended text and advise if acceptable. An alternative is ↑↓but presume this would not be acceptable.

Introduction

Include more background about MMVD. What is the incidence of this disease? What are theories about why CKCS are more susceptible to the disease?

We would normally do so, but are constrained by the word count for the MS. Also much of what is being asked to include would be speculative. Have included one sentence on endothelial damage, VC activation and aberrant remodelling.

What are the current theories about MMVD formation?

See previous response

Please rewrite the aim of the study in the last paragraph as it is unclear (lines 86-91)

Has been reworded.

Materials and Methods

How was MMVD diagnosed in the dogs?

Clinically and confirmed by echocardiography; MS amended

Correct the discrepancy (or clarify the difference) between n = 6 at line 100 and n = 8 on line 117. How many samples were there?

Cannot find n=8 in the MS. Numbers are correct.

What kind of QC parameters were used to accept the microarray data?

If this means pre-hybridisation it is the RIN number which was >7, which is in the MS. If it means post-hybridisation we used the PCA plots, which mentioned in new line 128-130.

Line 145 specify the cut-off values used

Now line 134 with values entered for fold-changes etc.

What was the RNA input for the Q-RTPCR reaction? Line161

Not sure what is being asked here; please clarify. We used the exact same RNA samples for the microarray and the PCR. Have included comment in new line 145.

Correct the “-” in front of delta delta Ct on line 164 or explain what the “-” means.

No meaning; has been removed.

Table 2

Clarify what "Gene %" means? Does this refer to the % of the pathway which is changed? Add what general direction the pathways are changed to the table.

Sentence included to clarify

Table 3

Include which direction genes were changed (up or down regulated)

Have included sentence cross-referencing to Figure 3 which shows the fold changes. If that is not sufficient we can include in Table 3, but this this would make it very complicated.

Figure 4

This figure is very hard to read and interpret; please make it more clear. Suggestions include removing molecules from the networks which do not strengthen the authors point, building a unique heatmap including functional nodes and only necessary/unchanged gene changes, or representing the data in a
different format.

This has been replaced with Table 4; effectively Figure 4 without the graphic for LDL. Original Table 4 is now Table 5

Supplemental Figure 4 ◊ change title to "Identity" not Identify

Done

Reviewer #3: In this study, the authors used up-to-date canine microarray technology to investigate the transcript changes in MMVD of CKCS. They found genes, biological function clusters and pathways related to MMVD. The study was well conducted and the manuscript was well written. The authors only need to correct some punctuations to make the article more accurate and less confusing. eg. in Line58, there should be comma in front of ‘less is’; there should be right parenthesis some where in Line110. Please check your MS carefully to correct these issues.

MS has been checked.
Editor and Reviewer Comments:

Note from the Scientific Editor to the Authors,

As mentioned below in their comments, both reviewers have recommended your submission to be accepted for publication in The Veterinary Journal. However, for unknown reasons, I am unable to open the manuscript file and to make final edits. Can you please consider the few comments from Reviewer #1 below and save your manuscript in a different version of Word, as this may solve the problem?

In addition, please use the other submission (751: culture of VEC and VIC) that we just send back to you as an example of the edits that will be necessary for you to make before we can accept the manuscript. For example:

Done

1. P needs to be capitalized and italicized.
2. "n" needs to be italicized.
3. Do no include ":." after the section titles
4. In your in-text references, use the following format: Corcoran et al., 2014
5. Use "min", "h" and "s"
6. Use Fig., not Figure
7. In your reference list, the volume numbers should not be italicized. And replace all "&" by "and"

All done

Reviewer #1:

I find the paper very fine. I have only two minor comments:

I still find it important to address that the findings may be due to breed differences between the control group and the diseased group. I recommend that the sentence in the discussion (line 386-387): "The main limitations of this study were the small sample size, the lack of age-matched controls..." is changed to "The main limitations of this study were the small sample size, the lack of age-and breed-matched controls..."

Comment included

Model control in relation to ANOVA is performed to evaluate if raw data fit the statistical model. Evaluation of variance homogeneity can be performed. Non parametric tests need to be used if data not fit the model and transformation of the data does not obtain variance homogeneity. I find it important to know if the raw data fit the statistical models, it is not clear to me whether it is performed

We are unable to determine whether if the raw data fits the statistical model or not due to the small sample size. In PARTEK suite microarray analysis, the data are transformed (log2) such that all values are between 0 and 16. The software then normalises the data irrespective of sample size and analyses using ANOVA regardless of the statistical model (Gaussian). Applying FDR to this bioinformatics platform further corrects for any potential false discovery. This is a routine microarray analysis procedure and widely accepted considering the sample size limitation typical of this type of research. We have consulted with Mick Watson, Director of Edinburgh Genomics (merger of Ark Genomics and TheGenePool) and he is confident the data analysis is robust.
Gene network and canonical pathway analysis in canine myxomatous mitral valve disease: A microarray study

The Veterinary Journal

Dear Brendan,

Your revised paper has been edited by your Handling Editor, Dr Eric Blomme, and another edited version has been uploaded to the journal website for your approval. Please check this edited manuscript carefully for accuracy and completeness.

I would be grateful if you could address the editorial comments and annotations that Eric has made to your manuscript in the references.

In addition, please attend to the following editorial requests:

- In the Corresponding author's details, please remove "(0)" from the telephone number and change "Brendan Corcoran" to "B.M. Corcoran". Authors' names can be written in full in the main list of authors if desired.
  
  Done

- Please change "cavalier king Charles spaniels" to "Cavalier King Charles spaniels".

  Not sure if Cavalier is a proper noun, but done as requested.

- When writing numbers such as "n = 59", please write "n" in italics.

  Done

- Please check punctuation and spacing throughout the manuscript.

  Done; according to my own limited grammatical skills!!!

- Numbers less than 10 that are not linked to a unit should be written in full unless used as the start of a sentence, e.g. change "6 and 10 significantly different biological function clusters" to "six and 10 significantly different biological function clusters".

  Done

- Please change "Royal (Dick) School of Veterinary Study" to "Royal (Dick) School of Veterinary Studies".

  Done

- Please write the heading "Conclusions" in bold.
- References should be formatted with a hanging indent.

- Please write the article titles for Geirsson et al. (2012), Lakatta and Levy (2003), Liu et al. (2003), Mow and Pedersen (1999) and Rabkin et al. (2001) in sentence case.

- In the tables, the headings should be written in sentence case, not emboldened and placed on the next line below the headings "Table 1", "Table 2" and "Table 3".

- In Table 1, please write sequences in upper case in the format 5'-CCGCATTGAGCTGACAGTAG-3'. Please also change "Probe No." to "Probe number".

- In Table 4, please place additional data as footnotes rather than with the heading.

- Please check all references carefully for accuracy. Please also check that all references are cited in the text and that all citations are listed in the references.

- As recommended by Eric, Fig. 2B should be converted to a Table.

Done, and other figures re-numbered as appropriate.

- Please remove the solid borders from around Figs. 3A, B and C.

Done; now re-labelled Fig 2A, B and C.
- As noted by Eric, Fig. 4 is missing.

**My mistake; this figure had been removed but the text not amended accordingly. The figure does not add to the text.**

Please ensure that your revised paper conforms fully to the requirements of our Guide for Authors.

**IMPORTANT:** Your article has been edited by the Editor handling your paper and the Word filename includes the word 'edited'. You MUST ONLY use this version in preparing your revised text. You should click on download submission files link from your author menu on EES and ensure you are making changes to the most up to date version of the manuscript.

You should submit your revision online by logging onto the Elsevier Editorial System for The Veterinary Journal:
http://ees.elsevier.com/ytvjl/

Your username is: ******

We look forward to receiving your revised manuscript.

With kind regards,

Adrian

Dr Adrian W. Philbey BVSc(Hon) PhD MANZCVSc(Pathology) MRCVS,
Senior Scientific Editor,
The Veterinary Journal

On behalf of Dr Andrew Higgins BVetMed MSc PhD FSB MRCVS,
Editor-in-Chief,
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__________________________________________________________

**EDITOR'S COMMENTS:**

Note from the Scientific Editor to the Authors,

Based on your revisions, I have made some final edits to prepare your submission for publication. Please use ONLY the edited version of the manuscript (which should be downloaded from the Journal's Web site) to carefully review the text and make sure that I did not alter the meaning of it. I also include a few comments in the reference list for you to address. Make all changes in a font of a different color.

Finally, I could not locate figure 4, and Figure 2B would be better suited as a table.
With kind regards,

Eric Blomme, DVM, PhD, Dipl. ACVP

Scientific Editor,
The Veterinary Journal
Dear Editor.

29/7/2014

Please find enclosed manuscript “Gene network and canonical pathway analysis in canine myxomatous mitral valve disease: a microarray study” for consideration for publication in TVJ.

This paper provides a large amount of data on the mitral valve transcriptome that is not currently available to the research community. It markedly improves on the only other study previously reported in 2006, not least because of major advances in transcriptome coverage and bioinformatics platforms.

These data will permit increased numbers of hypothesis driven projects to be generated by the research community interested in canine MMVD, but also provide data of interest to colleagues working on the analogous human disease.

Brendan Corcoran