Transcriptomic profiling of atrial fibrillation canine models at disease onset

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ABSTRACT (249 words)

Atrial fibrillation (AF), the most common arrhythmia with an overall prevalence of 0.51%, is associated with increased morbidity and mortality, as well as important healthcare costs. AF develops over many years and is often associated with substantial changes in the structural and electrophysiological properties of the atria. Because AF may lack subclinical symptoms at onset, it has been difficult to study the molecular and cellular events associated with earlier stages of this pathology in humans. Here, we characterized comprehensively the transcriptomic changes that occur in the atria of two robust canine AF models, electrically maintained AF without and with a controlled ventricular rate achieved by radiofrequency-ABLATION of the atrioventricular node/ventricular pacing, after one week of maintenance. Our RNA-sequencing experiments identified thousands of genes that are differentially expressed, including a majority that have never before been implicated in AF. Gene set enrichment analyses identified known (e.g. extracellular matrix structure organization) but also many novel pathways (e.g. muscle structure development, striated muscle cell differentiation) that may play a role in tissue remodeling and/or cellular trans-differentiation. Of interest, we found dysregulation of a cluster of non-coding ribonucleic acids (RNAs), including many microRNAs but also the MEG3 long non-coding RNA orthologue, located in the syntenic region of the imprinted human DLK1-DIO3 locus. Our results capture molecular events that occur at an early stage of disease development using well-characterized animal models, and may therefore inform future studies that aim to further dissect the causes of AF and factors involved in its progression in humans.
INTRODUCTION

Atrial fibrillation (AF) is the most common type of sustained arrhythmia, with an estimated lifetime risk of 22%-26% and association with increased morbidity and mortality\(^1\). Despite advances in antiarrhythmic therapies, their suboptimal efficacy and adverse effects have limited their use\(^2\). Therefore, there is a need to further characterize fundamental arrhythmia mechanisms in order to discover new therapeutic targets\(^2\). Although AF is known to be a final common endpoint of atrial remodeling resulting from a variety of heart diseases, it can also be, in turn, a cause of remodeling. This vicious cycle is called “AF begets AF”\(^3\) and explains the progressive nature of this arrhythmia and the complexity of its management.

Atrial remodeling is characterized by ion channel dysfunction, Ca\(^{2+}\) handling abnormalities, and structural changes, which result in AF induction and maintenance\(^4-5\). Heart disease, and even rapid atrial activity itself, cause the development of atrial fibrosis, which is a hallmark of structural remodeling. The degree of fibrosis is positively correlated with the persistence of AF\(^6\). Atrial cardiomyocytes subjected to rapid activation release factors that induce fibroblast-to-myofibroblast differentiation that leads to increased collagen synthesis\(^7\).

Any arrhythmia causing a rapid ventricular rate, including AF, is a well-recognized inducer of ventricular dysfunction, so-called “arrhythmia-induced cardiomyopathy”\(^8\). Heart failure enhances atrial stretch and sympathetic tone, making AF more resistant to rate- or rhythm-control treatments\(^9\). AF promotion results from the rapid atrial rate, but rapid ventricular rates due to inadequate rate-control also promote AF-related atrial remodeling with a different profile from the remodeling produced by rapid atrial rate alone\(^10\). Radiofrequency atrioventricular node ablation (AVB) with right ventricular pacing is a nonpharmacological strategy for rate control that can improve symptoms and outcomes\(^11\).
Our previous work in canine AF models showed that maintaining AF for one week by rapid atrial pacing activates fibroblasts, collagen gene expression and cardiomyocyte ion channel changes, without yet causing fibrosis\textsuperscript{12}. Continued electrical maintenance of AF for 3 weeks produces fibrosis, but electrically-maintained AF with ventricular rate control through AVB produces less profibrillatory remodeling than 3 weeks of AF alone\textsuperscript{10}. However, how AF with and without AVB impact atrial remodeling at the molecular level has not yet been assessed comprehensively. To answer this question, we took advantage of our well-characterized AF dog models and performed RNA-sequencing (RNA-seq) of cardiomyocyte-enriched atrial samples after one week to capture the molecular actors of atrial remodeling. In comparison with control (CTL) dogs, we found thousands of messenger-ribonucleic acids (mRNAs), long non-coding RNA (lncRNA) and microRNA (miRNA) that are differentially expressed in the atria of the canine AF models. Pathway analyses of the transcriptomic data highlighted known biological processes, but also potential novel modulators of arrhythmia initiation which may shed new light on our understanding of AF in humans.
METHODS

All results and R code are available at https://github.com/lebf3/DogAF.

Canine atrial fibrillation model

A total of 18 adult mongrel dogs of either sex, weighing 18-32 kg, were obtained from LAKA Inc and assigned to control (CTL) group (n=6) and two canine AF-models (n=6/group). Animals were handled in accordance with the “Guide for the Care and Use of Laboratory Animals” established by the National Institutes of Health as approved by the Montreal Heart Institute Ethics Committee (2016-47-01, 2019-47-03 for control dogs, 2015,47-01, 2018.47.12 for AF dogs).

To induce AF, animals were subjected to atrial tachypacing without (AF)\textsuperscript{13} and with (AF+AVB)\textsuperscript{14} atroventricular-node ablation under 0.07 mg/kg acepromazine (IM), 5.3 mg/kg ketamine (IV), and 0.25 mg/kg diazepam (IV), and 1.5% isoflurane anesthesia. In the AF group, a bipolar pacing lead with fluoroscopic guidance was placed in the right atrial appendage (RAA). In the AF+AVB group, pacing leads were inserted into the RAA and right ventricular apex. Pacing leads were connected to a subcutaneous pacemaker implanted in the neck (right side). In the AF+AVB group, radiofrequency catheter ablation was used to create AF+AVB. For this purpose, a quadripolar catheter with fluoroscopic guidance was placed across the tricuspid valve via the right femoral vein. Radiofrequency energy was then used to perform ablation when action potential at the His bundle was detected. Twenty-four to seventy-two hours after surgery, dogs in the AF group were subjected to AF-maintaining atrial tachypacing at 600 bpm for seven days. In the AF+AVB group, RA and right ventricle were paced at 600 and 80 bpm, respectively. In animals of the CTL group, no pacemaker was inserted.
Enrichment of dog atrial cardiomyocytes

Cardiomyocytes were enriched from the left atrium (LA) with enzymatic digestion through the coronary artery-perfused Langendorff system, as previously described\textsuperscript{15}. Briefly, dogs were anesthetized with 2 mg/kg morphine (IV) and 120 mg/kg alpha-chloralose and mechanically ventilated. Hearts were aseptically and quickly removed after intra-atrial injection of 10,000 U heparin and placed in Tyrode’s solution containing 136 mM NaCl, 5.4 mM KCl, 2 mM CaCl\textsubscript{2}, 1 mM MgCl\textsubscript{2}, 10 mM dextrose, 5 mM HEPES, 0.33 NaH\textsubscript{2}PO\textsubscript{4} (pH was adjusted to 7.3 with NaOH). The left coronary artery of the isolated heart was cannulated, and the LA was dissected free and perfused with 100% oxygenated Tyrode’s solution (37°C, 1.8 mM Ca\textsuperscript{2+}). The arterial branches were ligated to have a leak-free system, and LA tissues were perfused with Ca\textsuperscript{2+}-free Tyrode’s solution for ~10 minutes, followed by ~1-hour perfusion with ~0.45 mg/mL collagenase (CLSII, Worthington, Lakewood, NJ) and 0.1% bovine serum albumin (Sigma–Aldrich, Oakville, ON) in Ca\textsuperscript{2+}-free Tyrode’s solution for enzyme digestion. Digested tissue was removed from the cannula and cut into small pieces, and atrial cardiomyocytes were harvested.

RNA-seq/miRNA-seq

Library preparation and sequencing

mRNA and miRNA libraries were prepared at Genome Québec. mRNA libraries were made with the NEBNext_dual kit (rRNA-depleted stranded) and sequenced on NovaSeq 6000 S2 PE100 Illumina platform generating 32-123M Paired-end reads per sample. miRNA libraries were prepared with TruSeq smRNA and sequenced on the HiSeq 4000 SR50 Illumina platform generating 10-12M reads per sample.
**Bioinformatic processing and DE analysis**

The complete analysis can be found at https://github.com/lebf3/Dog_AF_transcriptomic. Briefly, mRNA reads were pseudomapped on reference transcriptome CanFam3.1.98 with Kallisto\(^\text{16}\) with the options quant -t 5 -b 100 and the rest as default. We aggregated transcripts by genes with tximport\(^\text{17}\) and quantified with DESeq2\(^\text{18}\). Genes with 0 reads in more than 12 samples were removed. Shrunken log\(_2\) transformed expression corrected for library size, with and without fibroblast fraction as a covariate (within DEseq2’s model) were then analyzed for DE with Wald test for all pairwise comparisons of CTL, AF and AF+AVB and likelihood ratio test for total assay DE. We plotted the PCA with fibroblast fraction as a covariate from log\(_2\) transformed expression values corrected with Limma’s removeBatchEffect() function\(^\text{19}\) for visualization of fibroblast effect on the top 1000 most variable genes. We then compared sets of GENEIDs found to be up (L2FC > 0 & \(p < 0.01\)) or down (L2FC < 0 & \(p < 0.01\)) in all possible contrasts.

For miRNAs, we trimmed reads using fastp\(^\text{20}\) with default settings and aligned them to CanFam3.1.98 genome with STAR v2.7.1a\(^\text{21}\) according to ENCODE protocol\(^\text{22}\). DEseq2 DE analysis was then conducted with the same parameters as described above for mRNAs.

**Deconvolution of RNA-seq data**

To account for potential tissue heterogeneity, we used a murine atrial gene signature matrix described in\(^\text{23}\) and our matrix of gene expression in Fragments Per Kilobase of exon model per Million reads mapped (FPKM) in CIBERSORTx online tool\(^\text{24}\). We then performed nonparametric Wilcoxon test on all possible comparisons for fibroblast fraction with a statistical significance threshold of \(p < 0.05\).
**Gene set enrichment analyses**

For each gene sets described above, we performed hypergeometric testing against the human Gene Ontology (GO) Biological Processes (BP) from Molecular Signatures Database v7.1 with the HypeR package.

**miRNA target prediction**

For DE miRNA present in the 5 most cited miRNA databases (DIANA, Miranda, PicTar, TargetScan, and miRDB), we defined genes as targets if they were: i-annotated with a human homolog in the ensemble database, ii-predicted targets by at least 3 out of 5 databases queried with the MiRNAtap package, iii-differentially expressed (mRNA FDR < 0.01), iv-inversely correlated (Pearson’s $r$<-0.5) log2 expression, corrected for the fibroblast fraction (expression values corrected with Limma’s removeBatchEffect() function). We then performed a GSEA with the remaining 82 predicted targets of the miRNA located on the syntenic region of the Dlk1-Dio3 locus (CanFam3.1 Chr8:68961744-69696779) as described above.

**RNA-seq and miRNA-seq DE genes comparison between human AF patients and canine AF models**

DE genes in our canine AF models with annotated human orthologues in the ENSEMBL database were compared to 2 human AF atrial tissue RNA-seq studies investigating both miRNA and mRNA\(^{25-26}\). miRNA precursors were retrieved with the R package miRBaseConverter and then compared across species. Because only 1 miRNA was found to overlap with human DE miRNA, only mRNA genes found to be DE in human and our canine AF models are represented as an Upset plot.
Proteomics

Dog cardiomyocytes were lysed by sonication, reduced and alkylated. Protein was precipitated, resuspended quantified and subjected to tryptic digest. Peptides (500 ng) were analyzed by reverse phase nano-HPLC coupled to a Bruker maXis II mass spectrometer (positive mode, mass range 150 - 2200 m/z, collision induced dissociation of top 20 precursors). LC-MS/MS data were analyzed for protein identification and label-free quantification using MaxQuant\textsuperscript{27} (1.6.1.0) against the public database UniProt with taxonomy Canis lupus familiaris and common contaminants (downloaded on 01.08.2019, 29809 sequences) with carbamidomethylation on Cys as fixed and oxidation on Met as variable modification with decoy database search included (mass tolerance 0.006 Da for precursor, 80 ppm for product ions; 1 % PSM and protein FDR, match between runs enabled, minimum of 2 ratio counts of quantified razor and unique peptides).

DE analysis and correlation

Proteins with > 3 missing values per treatment were removed. The remaining missing intensities were replaced with random values taken from the Gaussian distribution centered around a minimal value from the 10th quantile with the \textit{DEP} package’s Minprob function, to simulate a relative label-free quantification (LFQ) value for those low abundant proteins. Two-sample t-tests with subsequent multiple testing correction by FDR were used to identify DE proteins (\(p<0.01\)) with the fibroblast fraction as covariate using the Limma package. Because proteomic processing does not always converge to a single protein, only 752 genes out of the 1029 in the proteomic matrix were correlated to their corresponding RNA-seq data. We compared overlapping genes’ mean log\textsubscript{2} transformed expression in proteomic and RNA-
seq. The distribution of mean RNA-seq expression of the 752 overlapping genes was then compared to the full mean RNA-seq gene expression values.
RESULTS

RNA-sequencing of cardiomyocyte-enriched atrial samples from canine AF models

We analyzed data from three groups of six dogs. The first group (CTL) was the control group without atrioventricular ablation (AVB) nor pacemaker, in the second group (AF), right atrial-tachypacing at 600 beats per minute (b.p.m.) was used to maintain AF electrically for one week, and the third group (AF+AVB) included dogs with electrically-maintained AF for one week in the presence of AVB and ventricular pacing at 80 b.p.m. to control the ventricular rate. We reasoned that transcriptomic profiling of atria from these animals should allow us to discover the molecular changes that occur over the first week after the onset of AF, and play a role in the development of the tissue remodeling accompanying the transition from paroxysmal to persistent AF.

Initial analysis of bulk RNA-seq data hinted at some heterogeneity of cellular composition across samples. Therefore, we estimated the fraction of the major cell types in each sample using an in-silico deconvolution technique implemented in CIBERSORTx (Fig. 1A). Because of the induced tissue remodeling due to the AF treatments, we found that both AF and AF+AVB dogs had more fibroblasts in their atria than CTL animals (Fig. 1B). To emphasize the transcriptional differences between conditions that are not a result of variable cellular composition, we included the fibroblast fraction as a covariate in all subsequent differential expression (DE) analyses. Correction for this confounding variable reduced inter-group variability (Fig. 1C-D).

Proteomic analysis largely confirms the transcriptomic results

To validate our RNA-seq results, we took advantage of mass spectrometry (MS)-based protein quantification results from the same 18 dog atrial cardiomyocyte-enriched cell extractions that
were generated in a parallel study (detailed proteomic results will be presented elsewhere). After stringent quality control, we obtained relative quantification for 752 proteins. For these genes, the relative RNA and protein levels were strongly correlated (Pearson’s $r=0.49$, $P=1.57 \times 10^{-46}$) (Fig. 2A). Many of the genes that are well-correlated encode abundant cardiomyocyte proteins, such as titin ($TTN$), myosin light chain-4 ($MYL4$), desmin ($DES$), and tropomyosin-1 ($TPM1$). We found that RNA-seq could profile transcripts with a wider range of expression profiles, whereas MS-based proteomics preferentially captured proteins whose genes are expressed at high levels. (Fig. 2B).

**Transcriptomic changes in cardiomyocyte-enriched atrial samples**

Pairwise comparisons of gene expression levels between the three groups of dogs identified 434, 5971, and 7867 genes that are DE (false discovery rate (FDR) <0.01) in atrial cardiomyocyte-rich fractions in AFvsCTL, AF+AVBvsCTL, and AFvsAF+AVB, respectively (Fig. 3A-B). All differential gene expression level results are available in Table S1 and https://github.com/lebf3/DogAF). Many genes previously implicated in AF are dysregulated in both AF and AF+AVB dogs when compared to controls, thus validating the experimental design. This includes $FHL1$ involved in myofilament regulation$^{28}$, $SORBS2$ involved in intercalated disc gap junction regulation$^{29}$, and $KCNA5$, which regulates atrial action potential repolarization$^{30}$. However, our data also implicates genes not previously recognized to be involved in AF, such as leukocyte receptor cluster member-8 ($LENG8$), transcription elongation regulator-1 ($TCERG1$), ligand dependent nuclear receptor corepressor ($LCOR$), formin-binding protein-4 ($FNBP4$), and $ENSCAFG00000049959$ (orthologue of the long non-coding RNA (lncRNA) $MEG3$) (Fig. 3A, Table S2, and https://github.com/lebf3/DogAF).
To understand what pathways are modulated in the atria of these canine AF models, we performed gene set enrichment analyses (GSEAs) on the DE genes (Fig. 3C and Table S3). In AFvsCTL, we noted an up-regulation of genes associated with profibrotic pathways (e.g. extracellular structure organization, biological adhesion, response to wounding) and a down-regulation of genes implicated in angiogenesis, such as blood vessel morphogenesis. Genes implicated in muscle biology were up-regulated in the AF+AVBvsCTL analysis (e.g. muscle structure development, striated muscle cell differentiation) whereas the same comparison implicated down-regulated genes involved in ion transport and signaling pathways (e.g. sensory perception). We confirmed that this enrichment was not due to a smaller fraction of cardiac neurons found in the atria of AF+AVB dogs (Kruskal-Wallis’ $P=0.32$). Because of the large overlap in genes that are down-regulated in AF+AVBvsCTL and up-regulated in AFvsAF+AVB (Fig. 3B), we identified similar pathways in the GSEA for these two comparisons (in Fig. 3C, compare AF+AVBvsCTL and AFvsAF+AVB). Finally, genes that were down-regulated in the AFvsAF+AVB analysis implicated genes with more generic functions in gene expression and chromatin modifications, such as the histone-lysine N-methyltransferase SETD5 and the DNA methyltransferase TET2. Dysregulation of the expression of these chromatin-related genes and pathways is consistent with the extensive transcriptomic changes observed in the atria of AF+AVB dogs, in sheep models of AF (Fig. S1) as well as in AF patients$^{31-32}$.

**Dysregulation of miRNA expression**

Because miRNA play important roles in AF biology$^{33}$ but are not detected in standard RNA-seq protocols, we performed in parallel miRNA-seq on the same dog samples. We found 31, 19 and 21 miRNA that are DE (FDR <0.01) in AFvsCTL, AF+AVBvsCTL and AFvsAF+AVB, respectively (Fig. 4A, Table S1 and link to app). When comparing miRNA expression in the
two AF models, *MIR185* on the dog chromosome 26 was the most DE miRNA with strong up-regulation in the atria of AF animals. We also noted that 11 of the most strongly DE miRNA in the AF+AVBvsCTL and AFvsCTL analyses (*MIR136, MIR411, MIR370, MIR127, MIR493, MIR494, MIR485, ENSCAF00000025655 (96.20% identity to hsa-mir-379), MIR758, MIR543, MIR889*) mapped to the chr8:68,900,000-69,700,000 region in the dog reference genome CanFam3.1 (*Fig. 4B*). This region, highly conserved in mammals, is syntenic to the imprinted 14q32 region in humans (also known as the *DLK1-DIO3* locus)\(^34\). The IncRNA *MEG3*, which we described above as being over-expressed in the AF canine models is also located in the same *DLK1-DIO3* syntenic dog locus.

The dysregulation of the expression of IncRNA and miRNA at the same locus suggested that they might co-regulate the expression of genes implicated in the same biological pathway(s). To address this possibility, we used in silico predictions to infer the DE RNA that are possible direct targets of these DE miRNA located at the syntenic *DLK1-DIO3* locus. For this analysis, we focused on DE miRNA and DE RNA that are predicted to physically interact by at least three out of five databases and that have expression levels that are negatively correlated in the RNA-seq/miRNA-seq experiments (Pearson’s *r* < -0.5). Using these filters, we identified 82 potential target genes for the DE miRNAs at this locus, with most genes targeted by a single miRNA (*Fig. 4C*). GSEA on these 82 genes indicated a common role in synaptic signaling involving glutamate signaling (*Fig. 4D* and *Table S4*). Some of the key genes within these pathways are metabotropic glutamate receptor-1 and -8 (*GRM1, GRM8*), glutamate ionotropic receptor delta type subunit-1 (*GRID1*), glutamate ionotropic receptor AMPA type subunit-1 (*GRIA1*), and corticotropin-releasing factor-binding protein (*CRHBP*).
Partial transcriptomic overlap between chronic human and acute dog AF atrial samples

To emphasize the ability of our canine AF models to potentially capture early transcriptomic changes which would be missed by profiling the atria of human AF patients who have developed the disease over years, we compared the DE genes identified in AF dogs with results from two recent human atrial transcriptomic profiling experiments. We found a single miRNA, MIRLET7F2, that is DE in both dog and human AF samples (Table S5). Out of 9,151 DE genes in our canine AF models, we found 203 that were also DE in human AF atria in at least one study (Fig. 5). The sets of DE genes present in both dog and human atria included genes implicated in ion transport (KCNQ4), tissue remodeling and fibrosis (COL1A1, COL3A1) and TGF-beta signaling (INHA, BMP5), probably reflecting common pathophysiology.
**DISCUSSION**

In this study, we used a transcriptomic approach to comprehensively assessed the molecular remodeling of AF-induced remodeling with and without AVB from atria cardiomyocyte-enriched samples. We confirmed the occurrence of known AF factors like the reactivation of developmental pathways, but also found a strong and novel association with microRNAs and IncRNA from the *DLK1-DIO3* locus, including the *MEG3* canine orthologue. This finding is concordant with the many chromatin remodeling genes dysregulated in our models, which is an emerging phenotype of AF both in human and sheep models\(^{35}\).

**Molecular remodeling in AF with versus without AVB**

We did not expect to find a smaller number of DE genes in the AFvsCTL analysis, given our previous observation that AF treatment alone without AVB results in more important tissue remodeling\(^{10}\). One possible explanation is that our prior histological studies were done in AF animals treated for three weeks,\(^{10}\) whereas the results presented here reflect RNA changes after one week of AF. The transcriptomic changes in the AF+AVB group show that cells are under active chromatin modification, indicating ongoing adaptation to the stimulus. This is not observed in the AF group (lacking AVB), which may indicate that this adaptation has already occurred. This idea would be consistent with the down-regulation of chromatin-related genes recently noted in the atria of sheep AF models\(^{35}\), and may be a result of earlier establishment of profibrotic transdifferentiation in AF compared to AF+AVB canine models.

**Potential role of non-coding genes at the *DLK1-DIO3* locus in early AF**

The highly-conserved *DLK1-DIO3* locus hosts two differentially DNA-methylated regions (DMR) modulating the expression of its non-coding RNA clusters, where in humans the maternal allele is hypomethylated with concomitant expression on the hypermethylated
paternal allele of non-coding RNA and other protein-coding genes \((DLK1, RLT1, \text{and } DIO3)\)\(^{34}\). In both AF+AVBvsCTL and AFvsCTL, we found a large proportion (58% and 23%, respectively) of DE miRNA at this locus, underlying its importance in AF-related adaptation. We also found dysregulation of the \(MEG3\) lncRNA canine orthologue at this locus. \(MEG3\) is a highly expressed lncRNA that has been studied in various pathologies, including cancer\(^{36}\) and more recently cardiovascular diseases\(^{37}\). Non-coding RNAs at this locus have been shown to mediate various cardiac developmental programs\(^{34}\). More specifically, \(MEG3\) can contribute to the recruitment of the Polycomb repressive complex-2 (\(PRC2\))\(^{38}\), a key chromatin modulating factor. Of particular interest, Mondal et al. showed that through interaction with the H3-Lys-27 methyltransferase \(EZH2\), \(MEG3\) can repress TGF-beta target genes, which are known to promote a profibrotic response\(^{39}\). Data have been presented that suggest an important role of \(EZH2\) and/or \(EZH2\)-regulated genes in AF\(^{40}\).

**Glutamate receptor regulation by miRNAs from the \(DLK1-DIO3\) locus**

Our GSEA analysis-predicted gene targets of DE miRNA at the \(DLK1-DIO3\) locus suggest a role for glutamate signaling in AF. Immunostaining has confirmed the presence of glutamate receptors on cardiomyocytes\(^{41}\). Glutamate was also found to be significantly increased in AF patients left atrial appendages\(^{42}\). Glutamate signaling is important in vagal afferent neurons\(^{43}\), and remodeling of the glutamate system in AF may relate to the extensive previous evidence of autonomic dysfunction in AF patients\(^{44}\). Therefore, the \(DLK1-DIO3\) miRNA cluster may be an adaptative regulator of cardiomyocyte excitability or of neural cells in the presence of AF.

**Limitations**

We used cardiomyocyte-enriched samples in an attempt to obtain clearer results from the transcriptomic analysis by excluding extrinsic variability due to changes in cell composition.
However, while our samples are enriched in cardiomyocytes, they do not constitute a pure cardiomyocyte preparation. A disadvantage is that variability due to changes in cell composition is not eliminated. Our cardiomyocyte-enriched samples allow us to detect otherwise potential features of AF related to non-cardiomyocyte cells, such as autonomic dysregulation mediated by neural cells; however, we cannot unambiguously attribute DE genes to transcriptomic changes in a specific cell type. In part, we were able to control for fibroblast composition by adjustment through analysis for expression of fibroblast-related RNA-expression patterns. Nevertheless, features underlined here should be confirmed in pure cell cultures or single cell transcriptomic assays. A second limitation is the difficulty in extrapolating our findings to gene expression changes in humans. We found only modest overlap of DE genes in our model compared to reported DE gene patterns in human; several explanations could explain this (e.g. differences in biospecimen preparation, tissue heterogeneity, fundamental differences between dog and human AF pathology). It is also possible that different transcriptomic programs may be involved at the initiation of arrhythmia and tissue remodeling (AF and AF+AVB dog models) when compared to those dysregulated in the atria once the pathology has been present for years (as is typically the case for human patients from whom atrial samples are obtained).

**Conclusion**

Understanding the pathophysiology of chronic human diseases such as AF is challenging because they develop over many years and initially present with only unremarkable pre-clinical symptoms. In this study, we took advantage of two well-characterized canine AF models to chart the transcriptomic changes that occur at the onset of arrhythmia. Despite the inherent limitations in relating dog models to human AF, our results offer interesting new hypotheses for future testing, including in man. In particular, the up-regulation of miRNAs at the *DLK1-DIO3* locus after 1 week of tachypacing suggests that they may be early biomarkers of tissue...
remodeling and/or adaptation in the atria. If this miRNA signature is also detectable in the blood\textsuperscript{45}, it might be promising to test if it could predict development towards AF in individuals with no or only pre-clinical symptoms.
Acknowledgments

This work was funded by the Fonds de Recherche en Santé du Québec (FRQS), the Canada Research Chair Program, the Montreal Heart Institute Foundation (MHIF), Heart and Stroke Foundation of Canada (grant #18-0022032), Canadian Institutes of Health Research (CIHR) (grant # 148401), the Austrian Science Fund (FWF) (projects KL1645, W1226 and F73 to RBG). F.L. received scholarships from the CIHR, FRQS, MHIF and Université de Montréal.
References


22. ENCODE ENCODE_miRNA-seq_STAR_parameters


Figure 1. Deconvolution of canine atria cell composition using bulk RNA-sequencing.

(A) We inferred cell fractions with CIBERSORTx and an atrial-specific gene signature matrix obtained using orthologous murine genes. We present cell fractions for each dog sample that we analyzed in this study. CTL, control; AF, Atrial-tachypacing; AF+AVB, AF with Atrio-Ventricular Block. (B) When we group animals per treatment arm, we observed a significantly higher fraction of fibroblasts in the atrial fibrillation dog models (AF and AF+AVB) than in the control animals (AFvsCTL Wilcoxon's test $P=0.0087$ and AF+AVBvsCTL $P=0.015$). Principal component analysis of the top 1000 most variable genes expressed in canine atria before (C)
and after (D) correction for fibroblast fraction show treatment-dependent clustering after correction for cell composition.
Figure 2. Validation of highly expressed RNA by proteomics. (A) In 18 atrial samples, 752 genes found in both datasets are correlated at the protein (x-axis) and RNA (y-axis) levels (Pearson’s r=0.49, P=1.57x10⁻⁴⁶). For reference, we annotated 15 genes that are differentially expressed in the RNA-seq experiment and have high protein expression levels. N.S., not differentially expressed in the RNA-seq or proteomic experiment; RNA.sign, genes that are differentially expressed (DE) in the RNA-seq assay only; Both.sign, DE genes in both the RNA-seq and proteomic experiments. The grey area around the line corresponds to the 95% confidence interval. (B) Relative expression level of all transcripts measured in the RNA-seq experiment. The histogram shows that genes that are present in both the RNA-seq and
proteomic experiment are highly expressed (Common, dark grey) in comparison to the expression levels of all transcripts measured (all_RNA, light grey).
Figure 3. Analyses of differentially expressed atrial genes identify many biological pathways that are dysregulated in atrial fibrillation dog models. (A) Volcano plots of all transcripts that we analyzed in this study. Transcripts in red have a false discovery rate (FDR)<0.01. We found 434, 5971 and 7867 genes that were differentially expressed (DE) in the AFvsCTL, AF+AVBvsCTL, and AFvsAF+AVB analyses, respectively. The full DE results are available in Table S1. (B) Upset plot showing the intersection of up-and down-regulated DE genes (FDR<0.01) in each analysis. (C) The five most significant biological pathways identified using gene-set enrichment analyses (GSEA) for each set of DE genes (FDR <0.01). Full results are available in Table S3.
Figure 4. 11 differentially expressed microRNAs (miRNAs) map to a canine chromosome 8 region that is syntenic to human DLK1-DIO3. (A) Volcano plots of all miRNA that we measured in our experiments. We identified 31, 19 and 20 miRNA that are differentially expressed (DE) (false discovery rate (FDR) <0.01) in the AFvsCTL, AF+AVBvsCTL and AFvsAF+AVB analyses, respectively. (B) Miami plots of miRNA and their corresponding statistical significance (y-axis) for the AF+AVBvsCTL (top) and ATvsCTL (bottom) analyses. An arrow indicates the miRNA cluster located on the canine chromosome 8 region that is syntenic to human DLK1-DIO3. The odd and even chromosomes FDR values are in blue and red respectively. (C) Upset plot showing the DE miRNA targets located in the syntenic DLK1-
DIO3 locus and their corresponding number of potential target RNA. We identified potential targets with the MiRNAtap package (predicted by ≥ 3 databases) from DE miRNA (FDR < 0.01) and DE mRNA (FDR < 0.01). (D) Gene-set enrichment analyses (GSEA) with the potential gene targets (x-axis) of the DE miRNA located at the syntenic DLK1-DIO3 locus. We only present the top five pathways enriched in this analysis. A red square in the heatmap indicates membership of a given target gene to the biological pathways located on the left (empty columns were removed for clarity). GSEA FDR and AF + AVB vs CTL DE FDR are on the right and top of the heatmap, respectively.

**Figure 5.** Overlaps in genes differentially expressed in canine AF models and human AF patients. We compared differentially expressed (DE) genes in our canine AF models with annotated human orthologues that are DE in human AF atrial tissue25-26.