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Research article

Age-dependent myocardial transcriptomic changes in the rat. Novel insights into atrial and ventricular arrhythmias pathogenesis

Modificări transcriptomice dependente de vârstă în miocardul de șobolan. Noi aspecte referitoare la patogeneza aritmiilor atriale și ventriculare

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Abstract

Background: Aging is associated with significantly increased prevalence of cardiac arrhythmias, but transcriptional events that underlie this process remain to be established. To gain deeper insight into molecular mechanisms of aging-related cardiac arrhythmias, we performed mRNA expression analysis comparing atrial and ventricular myocardium from Wistar-Kyoto (WKY) rats of different ages. Methods: Atrial and ventricular sampling was performed in 3 groups (n=4 each) of young (14-week-old), adult (25-week-old), and aging (47-week-old) WKY rats. mRNA expressions of 89 genes involved in cardiac arrhythmogenicity were investigated using TaqMan Low Density Array analysis. **Results:** Of the 89 studied genes, 40 and 64 genes presented steady atrial and ventricular expressions, respectively. All genes differentially expressed within the atria of WKY rats were up-regulated with advancing age, mainly the genes encoding for various K^+ , Ca^{2+} , Na^+ channels, and type 6 collagen. Atrial expression levels of 19 genes were positively correlated with age. Ventricular transcriptomic analysis revealed a balance between up-regulated and down-regulated genes encoding for the same ion channels. **Conclusion:** Our results indicate the induction of an up-regulation transcriptional response in atrial but not ventricular myocytes with advancing age, suggesting that the two chambers undergo different molecular remodeling programs. Aging atria displayed a transcriptomic profile consistent with higher propensity to arrhythmias, including up-regulation of genes encoding for $I_{\rm p} I_{Ca-P} I_{Ca-P} I_{Na^+}$ outward K^+ currents, and collagen, while ventricular transcriptome did not

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seem to be significantly altered by aging. These observations could explain the higher propensity to atrial than ventricular arrhythmias in the elderly.

Keywords: experimental model; aging; transcriptome; atrial fibrillation; ventricular arrhythmias

Rezumat

Introducere: Deși înaintarea în vârstă este asociată cu creșterea prevalenței aritmiilor cardiace, evenimentele transcriptomice care stau la baza acestui proces rămân neelucidate. Pentru a identifica aceste mecanisme am realizat analiza expresiei ARNm la nivelul miocardului atrial și ventricular la șobolani Wistar-Kyoto (WKY) de diferite vârste. Metodă: Expresiile ARNm a 89 de gene au fost analizate prin TLDA utilizând eșantioane atriale și ventriculare de la 3 grupuri (n=4 fiecare) de șobolani WKY tineri, adulți și vârstnici. **Rezultate:** Din cele 89 de gene, 40 și respectiv 64 de gene au prezentat expresii stabile la nivel atrial și respectiv ventricular. Toate genele exprimate diferit în atriile șobolanilor WKY au prezentat o up-regulare cu înaintarea în vârstă, în special genele care codifică canale de K^+ , Ca^{2+} , Na^+ și colagenul de tip 6. Expresia atrială a 19 gene a fost corelată pozitiv cu vârsta. Analiza transcriptomică ventriculară a evidențiat un echilibru între genele up-regulate și down-regulate codificând aceleași canale ionice. Concluzii: Rezultatele noastre indică un răspuns transcripțional atrial, dar nu și ventricular, caracterizat prin up-regulare odată cu înaintarea în vârstă. Aceste rezultate sugerează că cele două cavități sunt supuse unor programe de remodelare moleculară diferite. Atriile îmbătrânite au prezentat un profil transcriptomic sugestiv pentru o predispoziție crescută pentru aritmii, precum up-regularea genelor care codifică I_{f} I_{Ca-L} I_{Ca-T} I_{Na} I_{K} și colagenul, în timp ce transcriptomul ventricular nu pare să fie sever alterat de înaintarea În vârstă. Aceste observații ar putea explica predispoziția mai mare pentru aritmii atriale decât ventriculare la vârstnici.

Cuvinte cheie: model experimental; înaintare în vârstă; transcriptom; fibrilație atrială; aritmii ventriculare *Received:* 5th *December 2013; Accepted:* 17th *January 2014; Published:* 14th *February 2014.*

Introduction

Aging is associated with significantly increased prevalence of both atrial and ventricular arrhythmias (1,2), likely due to age-associated changes in heart structure and function (3). Factors such as DNA and/or protein damage due to increased oxidative stress or late-acting mutations could account for the adverse effects of aging on myocyte structure and function (4-8). However, molecular bases underlying the increased propensity to arrhythmogenesis in the elderly remain incompletely understood. Cardiac arrhythmias have been associated with altered expression of genes encoding for proteins involved in Ca2+-handling, ion channels, extracellular matrix, or cell-cell coupling (9), but expression profiles of these genes have never been studied in relation with aging.

Transcriptome analyses seem well suited for the identification of cardiac molecular abnormalities related to aging, and may provide a starting point for our understanding of pathophysiological mechanisms linking aging to cardiac arrhythmias. However, studies on aging are rather difficult in humans due to increased incidence of cardiovascular diseases that limit survival and complicate genetic studies, making it difficult to determine to what extent mRNA abnormalities are due to aging and not to other sources of biological variability (10). On the other hand, Wistar-Kyoto (WKY) rats are free from cardiovascular diseases (11-13); this makes WKY rats a good surrogate for human aging, allowing identification of age-associated changes in gene expression without interference from concomitant disorders.

In order to gain deeper insights into the molecular pathophysiology of aging-related cardiac arrhythmogenesis and to identify potential molecular markers and targets for novel antiarrhythmic therapies, we performed mRNA expression analysis using TaqMan Low Density Array (TLDA) analysis, comparing atrial and ventricular myocardium from WKY rats of different ages. Three stages of the aging process, 14 weeks of age, 25 weeks of age, and 47 weeks of age, corresponding to young, adult, and aging animals, respectively, were chosen in order to investigate and compare the transcriptome during

Material and methods

Animals

aging in WKY rats.

Three groups of male WKY rats 14 weeks (further mentioned as young; n = 4), 25 weeks (further mentioned as adult; n = 4), and 47 weeks (further mentioned as aging; n = 4) of age were purchased from Elevage Janvier (Le Genest Saint Isle, France).

All animals were housed in groups of two to three rats per cage in a climate-controlled room (21° to 24°C), with a 12-h/12-h light/dark cycle, in an accredited animal facility, and had free access to standard rat pellets and tap water.

All experiments were performed in compliance with the French Ministry of Agriculture guidelines for animal experimentation and were approved by the local Animal Ethics Committee.

Heart sampling

All animals were euthanized with an intraperitoneal injection of a terminal dose of sodium pentobarbital (>100 mg/kg). The hearts were excised and weighted; the left atrium and ventricle were isolated, placed in sterile dry tubes, snap frozen in liquid nitrogen, and stored at -80°C until processing.

Gene expression analysis

RNA extraction. Total RNA was isolated from frozen biopsies using TRIzol[®] Reagent (Life Technologies; Villebon sur Yvette, France) following manufacturer's instructions. RNA concentration was measured using Nanodrop ND1000 (Thermo Scientific; Illkirch, France) and quality was verified using the Agilent 2100 BioAnalyser (Agilent Technologies; Massy, France).

TaqMan Low Density Array analysis. To identify altered gene expression, we used a TLDA to examine the expression of 89 genes encoding for ion channels, proteins of the extracellular matrix, and proteins involved in cell-cell coupling. The choice of the 89 genes selected for mRNA expression analysis was based on their conclusive role in cardiac arrhythmogenicity (9). A detailed list of the 89 examined genes is provided in *Table I*. For the present study, the TLDA card was configured into four identical 89 genes set. For each set of genes, we used 6 endogenous control genes: *Ppia*, β -actin, Gapdh, Hprt1, Tbp, and 18s RNA.

Briefly, one μ g of total RNA was reverse-transcribed to complementary DNA (cDNA) by using High-Capacity cDNA Reverse Transcription Kits (Applied Biosystems) following manufacturer's instructions. The cDNA (2 μ L) was mixed with 48 μ L of H₂O and 50 μ L of 2X TaqMan Universal PCR Mix (Applied Biosystems). Each sample (100 μ L) was loaded into a port of the micro-fluid card and run on an ABI 7900HT System (Applied Biosystems) for 2 min at 50°C, 10 min at 94.5°C, followed by 40 cycles of 30 s at 97°C and 1 min at 59.7°C.

The TLDA data were analyzed using SDS 2.3 software (Applied Biosystems). Threshold cycle data for all target and control genes were transformed into molecule numbers (attomoles) considering polymerase chain reaction efficiency of 100%. Gene expression levels were normalized using TBP.

	samples of WKY rats of different ages.
Symbol	Name
Cacnalc	calcium channel, voltage-dependent, L type, alpha 1C subunit
Cacnald	calcium channel, voltage-dependent, L type, alpha 1D subunit
Cacnalg	calcium channel, voltage-dependent, T type, alpha 1G subunit
Cacnalh	calcium channel, voltage-dependent, T type, alpha 1H subunit
Cacna2d1	calcium channel, voltage-dependent, alpha 2/delta subunit 1
Cacna2d2	calcium channel, voltage-dependent, alpha 2/delta subunit 2
Cacnb1	calcium channel, voltage-dependent, beta 1 subunit
Cacnb2	calcium channel, voltage-dependent, beta 2 subunit
Cacnb3	calcium channel, voltage-dependent, beta 3 subunit
Cacnb4	calcium channel, voltage-dependent, beta 4 subunit
Cacng4	calcium channel, voltage-dependent, gamma subunit 4
Cacng6	calcium channel, voltage-dependent, gamma subunit 6
Casq1	calsequestrin 1 (fast-twitch, skeletal muscle)
Casq2	calsequestrin 2 (cardiac muscle)
Clcn2	chloride channel 2
Clcn3	chloride channel 3
Col6a1	collagen, type VI, alpha 1
Dsc2	desmocollin 2
Dsp	desmoplakin
Gjal	gap junction protein, alpha 1
Gja5	gap junction protein, alpha 5
Gja7	gap junction membrane channel protein alpha 7
Hcn1	hyperpolarization-activated cyclic nucleotide-gated potassium channel 1
Hcn2	hyperpolarization activated cyclic nucleotide-gated potassium channel 2
Hcn3	hyperpolarization-activated cyclic nucleotide-gated potassium channel 3
Hcn4	hyperpolarization activated cyclic nucleotide-gated potassium channel 4
Kcnal	potassium voltage-gated channel, shaker-related subfamily, member 1
Kcna2	potassium voltage-gated channel, shaker-related subfamily, member 2
Kcna3	potassium voltage-gated channel, shaker-related subfamily, member 3
Kcna4	potassium voltage-gated channel, shaker-related subfamily, member 4
Kcna5	potassium voltage-gated channel, shaker-related subfamily, member 5
Kcna6	potassium voltage gated channel, shaker related subfamily, member 6
Kcnab1	potassium voltage-gated channel, shaker-related subfamily, beta member 1
Kcnab2	potassium voltage-gated channel, shaker-related subfamily, beta member 2
Kcnab3	potassium voltage-gated channel, shaker-related subfamily, beta member 3
Kcnb1	potassium voltage gated channel, Shab-related subfamily, member 1
Kcnc3	potassium voltage gated channel, Shaw-related subfamily, member 3
Kcnc4	potassium voltage gated channel, Shaw-related subfamily, member 4
Kcnd1	potassium voltage-gated channel, Shal-related subfamily, member 1
Kcnd2	potassium voltage-gated channel, Shal-related subfamily, member 2
Kcnd3	potassium voltage-gated channel, Shal-related subfamily, member 3
Kcnel	potassium voltage-gated channel, Isk-related family, member 1
Kcnell	KCNE1-like
Kcne2	potassium voltage-gated channel, Isk-related family, member 2

 Table I. List of the 89 genes for which mRNA expressions were studied in left atrial and ventricular samples of WKY rats of different ages.

Table I. (continued)

Symbol	Name			
Kcne3	potassium voltage-gated channel, Isk-related family, member 3			
Kcne4	potassium voltage-gated channel, Isk-related family, member 4			
Kcnh2	potassium voltage-gated channel, subfamily H (eag-related), member 2			
Kcnh7	potassium voltage-gated channel, subfamily H (eag-related), member 7			
Kcnip2	Kv channel-interacting protein 2			
Kcnj11	potassium inwardly rectifying channel, subfamily J, member 11			
Kcnj12	potassium inwardly-rectifying channel, subfamily J, member 12			
Kcnj2	potassium inwardly-rectifying channel, subfamily J, member 2			
Kcnj3	potassium inwardly-rectifying channel, subfamily J, member 3			
Kcnj4	potassium inwardly-rectifying channel, subfamily J, member 4			
Kcnj5	potassium inwardly-rectifying channel, subfamily J, member 5			
Kcnj8	potassium inwardly-rectifying channel, subfamily J, member 8			
Kcnk1	potassium channel, subfamily K, member 1			
Kcnk3	potassium channel, subfamily K, member 3			
Kcnk5	potassium channel, subfamily K, member 5			
Kcnn2	potassium intermediate/small conductance calcium-activated channel, subfamily N, member 2			
Kcnq1	potassium voltage-gated channel, KQT-like subfamily, member 1			
Npr3	natriuretic peptide receptor C/guanylate cyclase C (atrionatriuretic peptide receptor C)			
Pak1	p21 protein (Cdc42/Rac)-activated kinase 1			
Myh7	myosin, heavy chain 7, cardiac muscle, beta			
Pkp2	plakophilin 2			
Ryr2	ryanodine receptor 2, cardiac			
Scn1a	sodium channel, voltage-gated, type I, alpha			
Scn1b	sodium channel, voltage-gated, type I, beta			
Scn2b	sodium channel, voltage-gated, type II, beta			
Scn3a	sodium channel, voltage-gated, type III, alpha			
Scn3b	sodium channel, voltage-gated, type III, beta			
Scn4a	sodium channel, voltage-gated, type IV, alpha subunit			
Scn5a	sodium channel, voltage-gated, type V, alpha subunit			
Scn7a	sodium channel, voltage-gated, type VII, alpha			
Scn9a	sodium channel, voltage-gated, type IX, alpha			
Smad6	SMAD family member 6			
Smad9	SMAD family member 9			
Smyd2	SET and MYND domain containing 2			
Trpc1	transient receptor potential cation channel, subfamily C, member 1			
Trpc3	transient receptor potential cation channel, subfamily C, member 3			
Trpc4	transient receptor potential cation channel, subfamily C, member 4			
Trpc5	transient receptor potential cation channel, subfamily C, member 5			
Тгрсб	transient receptor potential cation channel, subfamily C, member 6			
Trpm4	transient receptor potential cation channel, subfamily M, member 4			
Trpm5	transient receptor potential cation channel, subfamily M, member 5			
Trpv1	transient receptor potential cation channel, subfamily V, member 1			
Тгрνб	transient receptor potential cation channel, subfamily V, member 6			
Tgfb1	transforming growth factor, beta 1			
Camk2d	calcium/calmodulin-dependent protein kinase II delta			

Statistics

Data are expressed as means \pm SE. Nonparametric ANOVA (Kruskal-Wallis test) was used for multiple comparisons. Between-group comparisons were performed using the Mann-Whitney U test. Correlations were ascertained with Spearman's rank correlation method. A P value of less than 0.05 was considered statistically significant. Statistical analyses were undertaken using GraphPad Prism[®] software (GraphPad Software; San Diego, CA).

Results

Cardiac morphometry

Compared to young and adult WKY rats, aging rats had markedly increased body weights (Table II), but there was no significant difference in heart weights between WKY rats of different ages (all P > 0.05). Consequently, heart weightto-body weight ratios were significantly lower in aging WKY rats compared to young WKY rats (P = 0.04), while there was no significant difference between adult and young WKY rats, nor between aging and adult WKY rats (both P > 0.05). Among WKY rats, body weights were significantly positively correlated with age (Spearman r = 0.96, P < 0.001), but there was no significant correlation between heart weights and age (P =0.80), whilst heart weight-to-body weight ratios were significantly negatively correlated with age (Spearman r = -0.82, P = 0.01).

Gene expression profile

Left atrial and ventricular gene expression levels were compared between WKY rats of different ages. Significantly altered transcripts were selected by applying a fold change-cutoff of 1.5 and a *P*-value cutoff of 0.05.

Genes with steady expression profiles

Forty genes out of the 89 studied genes presented similar expression levels in atrial tissue samples from young, adult, and aging WKY rats (all P > 0.05). These genes included genes encoding for various subunits of the L-type Ca²⁺ current ($I_{C_{2}}$), for the Ca²⁺-binding protein calsequestrin 1, for various subunits of the delayed rectifier K^+ current (I_{K}), of the transient outward K^+ current (I_{to}), of the inward rectifier K^+ current (I_{K1}) , of the ATP-sensitive K⁺ current (I_{K-ATP}) , and of the acetylcholine-activated K^+ current (I_{K-Ach}), for various subunits of the voltage-dependent Na^+ current (I_{Na}), for non-selective cation currents, connexins, for the serine/threonine-specific protein kinase, for myosin heavy chain 7, for the N-lysine methyltransferase, and for the transforming growth factor *beta* (TGF β).

In ventricular tissue samples, 64 genes out of the 89 studied genes presented similar expression levels in young, adult, and aging WKY rats (all P > 0.05). These genes included genes encoding for I_{Ca-L} , I_{Na} , various subunits of I_K , I_{to} , and I_{K1} , for the hyperpolarization-activated inward current (I_p), various Cl⁻ currents, non-selective cation currents, for the atrionatriuretic pep-

Table II. Worphometric characteristics of WKT Tats.					
	Young WKY rats	Adult WKY rats	Aging WKY rats		
	(n = 4)	(n = 4)	(n = 4)		
Body weight (g)	398.2 ± 1.3	$509.7\pm4.8^{\dagger}$	$580.0 \pm 11.7^{*\ddagger}$		
Heart weight (mg)	1450.5 ± 131.6	1547.2 ± 32.5	1554.7 ± 69.8		
Heart weight / body weight (10-3)	3.6 ± 0.3	3.03 ± 0.1	$2.7 \pm 0.1^{*}$		

Table II. Morphometric characteristics of WKY rats.

Data are expressed as means \pm SE. * $P \le 0.04$ for aging versus young WKY rats; $^{\ddagger}P = 0.02$ for aging versus adult WKY rats; $^{\ddagger}P = 0.02$ for adult versus young WKY rats. P values refer to comparisons between groups based on the Mann-Whitney U test.

tide receptor C, for the serine/threonine-specific protein kinase, for various proteins involved in TGF β signaling, as well as for the N-lysine methyltransferase. None of the studied genes encoding for I_{K1}, I_{K-ATP}, I_{K-Ach}, Ca²⁺-binding proteins, or proteins involved in intercellular connectivity (connexins 40, 45, and 43, desmocollin 2, and desmoplakin) presented significantly different expression levels between the three groups.

Twenty-eight of the studied genes presented steady profiles in both atrial and ventricular tissue samples. These genes included most of the genes encoding for I_{Na} , for various outward K⁺ currents, Ca²⁺-currents, for the Ca²⁺-binding protein calsequestrin 1, for connexin-45, for the serine/threonine-specific protein kinase, and for the N-lysine methyltransferase.

Age-specific transcripts

Differentially expressed genes between young and adult WKY rats. Sixteen genes were differentially expressed with at least a 1.5-fold change in atrial tissue samples from young WKY rats compared to adult WKY rats. All these 16 genes were significantly up-regulated in adult compared to young rats. The most significant up-regulation was observed in *Hcn2*, encoding for I_{r} , which was 3.8 times more expressed in the atria of adult compared to young rats. Genes encoding for proteins involved in Ca²⁺-handling (*Cacnb1*, *Kcnn2*, and *Ryr2*), for I_{to} (*Kcna4*), and for a protein involved in TGF β signaling (Smad6) were also significantly up-regulated in adult compared to young WKY rats.

In ventricular tissue samples, 4 genes were differentially expressed between young and adult WKY rats, 3 of which were up-regulated, while 1 was down-regulated in adult compared to young WKY rats. The most significant up-regulation was observed in *Myh7*, encoding for myosin heavy chain 7, which was 3 times more expressed in adult than in young WKY rats. *Cacng4* and *Smad9*, encoding for the *gamma* subunit 4 of

the voltage-gated Ca²⁺-current and for a protein involved in TGF β signaling, respectively, presented both 1.5-fold higher expressions in adult compared to young WKY rats. Contrarily, *Col6a1*, encoding for non-fibrillar type VI collagen, was 1.5-fold less expressed in adult compared to young WKY rats. None of the genes with significantly different expressions within the atrial tissue presented a similar change within the ventricular tissue.

Differentially expressed genes between adult and aging WKY rats. Thirteen genes were differentially expressed in atrial tissue samples from adult compared to aging WKY rats. All 13 genes were significantly up-regulated in aging compared to adult WKY rats. The most significant up-regulation was observed in *Kcnb1* and *Kcna5*, both encoding for the ultrarapid component of I_{K} (I_{Kur}), which were 6-times and 4-times, respectively, more expressed in the atria of aging compared to adult WKY rats. *Col6a1*, encoding for type VI collagen, and *Npr3*, encoding for the atrionatriuretic peptide receptor C, were also significantly up-regulated in aging compared to adult WKY rats.

In ventricular tissue samples, 8 genes were differentially expressed between adult and aging WKY rats, 4 of which were significantly up-regulated, while 4 were significantly down-regulated in aging compared to adult WKY rats. The most significant up-regulation was observed in *Kcnal*, encoding for I_{K} , which presented a 15fold higher expression level in aging compared to adult WKY rats, while the most significant down-regulation was observed in Kcna4, encoding for I_{to}, which was 11-times less expressed in aging WKY rats than in adult WKY rats. The other up-regulated genes were Kcna6, encoding for I_{κ} , which had a 10-fold higher expression in aging than in adult WKY rats, Kcna3, encoding for I₁₀, which had a 5-fold higher expression in aging than in adult WKY rats, and Scn2b, encoding for a *beta* subunit of I_{N_a} , which was 3-times

more expressed in aging than in adult WKY rats. The other down-regulated genes were *Kcna2*, encoding for I_{Kur} , which had a 7-fold lower expression in aging than in adult WKY rats, *Myh7*, encoding for myosin heavy chain 7, which had a 3-fold lower expression in aging than in adult WKY rats, and *Scn4a*, encoding for I_{Na} , which was 1.7-times less expressed in aging than in adult WKY rats. Again, none of the genes with significantly different expression levels within the atrial tissue presented a similar change with-in the ventricular tissue.

Differentially expressed genes between young and aging WKY rats. Fourteen genes were differentially expressed in atrial tissue samples from young compared to aging WKY rats. All 14 genes were significantly up-regulated in aging compared to young rats. The most significant up-regulation was observed in Npr3, encoding for the atrionatriuretic peptide receptor C, which was 7-times more expressed in the atria of aging compared to young WKY rats. Hcn4, encoding for I_p Kcna5, Kcnj11, Kcnq1, encoding for various outward K⁺ currents, and Col6a1, encoding for type VI collagen, were also significantly up-regulated in aging compared to young WKY rats.

In ventricular tissue samples, 7 genes were differentially expressed between young and aging WKY rats, 4 of which were significantly up-regulated and 3 were significantly down-regulated in aging compared to young WKY rats. The most significant up-regulation was observed in *Kcnab2* and *Kcna1*, both encoding for I_{μ} , which were 22-times and 21-times, respectively, more expressed in aging than in young WKY rats, while the most significant down-regulation was observed in Kcna4, encoding for I,, which was 13-times less expressed in aging than in young WKY rats. The other up-regulated genes were Kcna6, encoding for I_{κ} , which had a 14fold higher expression in aging than in young WKY rats, and Kcna3, encoding for I_{to}, which had a 3.5-fold higher expression in aging than in young WKY rats. The other down-regulated genes were *Kcna2*, encoding for I_{Kur} , which had a 5-fold lower expression in aging than in young, and *Kcnab1*, encoding for I_{K} , which was 2.4-times less expressed in aging than in young WKY rats. Again, none of the genes with significantly different expressions within the atrial tissue presented a similar change within the ventricular tissue.

Gene expression levels and advancing age

Atrial transcriptomic analysis revealed 22 genes significantly differentially expressed between all three groups. One of these 22 genes presented a peak at adult age (*Hcn2*), two presented their lowest expression levels at adult age (Kcna3 and Kcnb1), while the remaining presented a continuous increase with advancing age (*Figure 1*). For these 19 remaining genes, atrial expression levels were significantly positively correlated with age. These 19 genes included genes encoding for proteins involved in Ca²⁺-handling, Cl⁻ currents, outward K⁺ and non-selective cationic currents, voltage-gated Na⁺ currents, the pacemaker current, collagen, proteins involved in intercellular connectivity, and atrionatriuretic peptide receptors.

At the ventricular level, 10 genes were significantly differentially expressed between all three groups. Four of these genes (*Cacnb2*, *Cacng4*, *Kcna2*, and *Myh7*) presented a peak at adult age, two (*Col6a1* and *Kcna3*) presented their lowest expression levels at adult age, while two (*Kcna1* and *Kcna6*) presented a continuous increase, and two (*Kcna4* and *Kcnab1*) presented a continuous decrease with advancing age (*Figure 2*). For 2 genes (*Kcna1* and *Kcna6*), ventricular expression levels were strongly positively correlated with age, whilst for other 2 genes (*Kcna4* and *Kcnab1*) ventricular expression levels were strongly negatively correlated with age.



Figure 1. Left atrial mRNA expression for genes significantly differentially expressed between young, adult, and aging WKY rats (n = 4 each). Data are expressed as means. Results are shown in log scale.



Figure 2. Left ventricular mRNA expression for genes significantly differentially expressed between young, adult, and aging WKY rats (n = 4 each). Data are expressed as means. Results are shown in log scale.

Transcripts with putative roles in aging-related cardiac arrhythmogenesis

At the atrial level, expression levels of several genes that have been previously related to increased arrhythmogenicity presented significant age-dependent changes.

Expression levels of *Hcn4*, encoding for I_P, were significantly higher in aging WKY rats compared to both young and adult WKY rats (160.8 ± 2.9 in aging *versus* 77.4 ± 0.3 in adult *versus* 39.4 ± 0.2 in young WKY rats, P = 0.01), and were significantly positively correlated with age (Spearman r = 0.93, P < 0.001). Similarly, expression levels of *Cacna1c* and *Cacna1g*, encoding for I_{Ca-L}, and for the T-type Ca²⁺ current (I_{Ca-T}), respectively, were also significantly higher in aging WKY rats compared to both young and adult WKY rats (268.6 ± 6.5 in aging *versus* 90.7 ± 1.1 in adult *versus* 49.8 ± 0.7 in young WKY rats, P = 0.01, and 478.7 ± 11.2 in aging *versus* 228.9 ± 0.3 in adult *versus* 93.1 ± 0.9 in young WKY rats, P = 0.03, respectively), and were significantly positively correlated with age (Spearman r = 0.88, P < 0.01, and Spearman r = 0.86, P < 0.01, respectively).

Expression levels of four genes encoding for outward K⁺ currents (*Kcnd3*, *Kcnip2*, *Kcnj11*, and *Kcnk3*) were also significantly higher in aging WKY rats compared to both young and adult WKY rats (*Figure 3*) and significantly positively correlated with age (Spearman r = 0.79, P =0.02; Spearman r = 0.77, P = 0.01; Spearman r = 0.81, P = 0.02; and Spearman r = 0.96, P <0.001, respectively). Up-regulated expression of genes encoding for I_{Na} (*Scn5a* and *Scn7a*) was also observed in aging WKY rats compared to both young and adult WKY rats (*Figure 3*) and were significantly positively correlated with age (Spearman r = 0.88, P < 0.01 for both genes).

Expression levels of *Col6a1*, encoding for type VI collagen, were also significantly higher in aging WKY rats compared to both young and



Figure 3. Age-dependency of left atrial mRNA expression levels for selected genes with putative roles in atrial arrhythmogenicity. Data are expressed as means ± SE.

adult (1251.2 \pm 34.0 in aging versus 441.0 \pm 4.3 in adult versus 286.9 \pm 0.4 in young WKY rats, P = 0.01) WKY rats (*Figure 3*), and were significantly positively correlated with age (Spearman r = 0.94, P = 0.001).

At the ventricular level, no transcriptional change could be related to a pro-arrhythmic pattern.

Discussions

The incidence of both atrial and ventricular arrhythmias increases progressively in the senescent heart (1, 2), even in the absence of apparent structural abnormalities (14). However, molecular bases underlying the increased propensity to arrhythmogenesis of the elderly remain incompletely understood. Transcriptome analyses seem well suited for providing deeper insights into the molecular pathophysiology of aging-related cardiac arrhythmogenesis and identifying potential targets for novel antiarrhythmic therapies.

Steady expression profiles for the majority of studied genes

Transcriptomic analysis of left atrial samples revealed a static expression profile for almost 45% of the studied genes with advancing age. With 64 genes out of the 89 studied genes (71.9%) presenting similar ventricular expression levels in the three stages of the aging process, ventricular transcriptome displayed an even more stable profile. The large majority of genes presented stable mRNA levels in both the atria and the ventricles of rats of different ages. These genes, including genes encoding for calsequestrin 1, connexin-45, serine/threonine-specific protein kinase, or N-lysine methyltransferase, may indicate a panel of genes that are 'resistant' to both atrial and ventricular aging-related molecular remodeling.

Transcripts with putative roles in aging-related cardiac arrhythmogenesis

Our results indicate that the induction of an up-regulation transcriptional response is a common feature of atrial, but not ventricular, cellular aging. Moreover, none of the genes with significantly altered expression levels within the atrial tissue presented the same change within the ventricular tissue. These observations suggest that the two chambers of the heart undergo different molecular remodeling processes.

The most common arrhythmia found in the elderly is atrial fibrillation (AF), with a 100-fold higher prevalence in octogenarians compared to those younger than 55 years of age (15-17). Two main electrophysiological mechanisms seem to underlie the occurrence of AF: focal ectopic activity and reentry circuits, favored by either altered electrical properties or by fixed structural changes of the atrial tissue. While ectopic activity can arise from increased cell automaticity outside the normal electrical system of the heart and/or the occurrence of afterdepolarizations, reentry has been related to decreased conduction velocity and/or atrial refractoriness shortening (18).

Increased automaticity originating in, but not confined to the pulmonary veins sleeve myocardium, plays an important role in AF genesis. Although the exact mechanisms that initiate abnormal automaticity are not well understood, it is well recognized that unusual presence of the pacemaker current I_f, which is responsible for the spontaneous diastolic depolarization in sinus node cells but normally lacks in cells outside the electrical system of the heart, as well as abnormal up-regulation of I_{Ca-T} in these cells, play a dominant role in this process (19). In the present study, one of the most obvious alterations in the aging atria was the up-regulation of Hcn4, encoding for the I_f current. Moreover, aging atria also displayed up-regulated expression of Cac*nalg*, encoding for I_{Ca-T} . Both these molecular abnormalities could underlie the higher propensity to atrial automaticity observed in the elderly.

Additionally, aging atria displayed up-regulated expressions of genes encoding for $\boldsymbol{I}_{\text{Ca-L}}$ and I_{Na} currents, which could cause increased inward Ca²⁺ and/or Na⁺ movement during action potential phases 2 or 3, favoring the genesis of early afterdepolarizations. Fibrillating atria have been shown to exhibit significant I_{Cal} down-regulation, probably as an adaptive, protective mechanism designed to prevent the cytotoxic effects of intracellular Ca²⁺ overload (20). This mechanism appears to be involved in atrial refractoriness shortening and reentry. On the other hand, up-regulated expression of Cacnalc, encoding for I_{Cal}, observed in our study, seems to characterize aging, but non-fibrillating atria, and may underpin intracellular Ca2+ overload, facilitating the formation of early afterdepolarization in aging atrial myocytes (21). Similarly, fibrillating atrial myocytes usually display a significant reduction in I_{Na} , which could contribute to decreased conduction velocity and reentry (22). However, aging non-fibrillating atrial myocytes seem to exhibit up-regulation of genes encoding for I_{Na} , which could further participate to the higher propensity to early afterdepolarizations of these cells. In fact, a gain of function mutation in the SCN5A gene encoding for $I_{_{\rm Na}}$ has been recently reported in a large Japanese family with AF (23), demonstrating the causative effect of $I_{N_{a}}$ up-regulation in AF onset.

The significant up-regulation of several genes encoding for outward K⁺ currents could favor atrial action potential and refractoriness shortening, predisposing aging atria to reentry. Atrial action potential and refractoriness shortening is often observed in patients with gain of function mutations in genes encoding for outward K⁺ currents, including KCND3, KCNJ11, and KCNK3 (24-26). Furthermore, *Col6a1* expression, encoding for non-fibrillar type 6 collagen, was significantly up-regulated in aging

atria, in concordance with previous studies that demonstrate increased atrial fibrosis in this context (27). Up-regulated *Col6a1* expression may be related to conduction slowing and thus further favor reentry.

At the ventricular level, no transcriptional change could be related to a pro-arrhythmic pattern. A number of genes presented significantly different ventricular expression levels in aging WKY rats compared to both adult and young WKY rats. However, ventricular transcriptomic profile seems to reflect the activation of a compensatory program in aging cells, with some genes encoding for a given ion channel being up-regulated while others are down-regulated, suggesting the activation of the repolarization reserve.

Our study demonstrates for the first time that a number of molecular features predisposing to atrial arrhythmias could be directly related to the aging process itself, explaining the higher propensity to atrial arrhythmias of the elderly. Contrarily, based on the panel of studied genes, it appears that the ventricular myocardium is much less 'sensitive' to molecular aging, explaining the lower propensity to ventricular than atrial arrhythmias of the elderly. However, despite the pro-arrhythmic atrial molecular profile, we previously demonstrated using continuous electrocardiographic recordings that aging WKY rats do not develop significant atrial arrhythmic events, contrary to aging spontaneously hypertensive rats, which displayed high atrial arrhythmogenicity (28). Taken together, these observations suggest that atrial molecular aging is not a sufficient condition but more likely a facilitator of atrial arrhythmogenicity, and that the presence of other factors, such as arterial hypertension, is needed in order to initiate the arrhythmias. A similar transcriptomic analysis in arrhythmic aging spontaneously hypertensive rats would therefore be of interest.

Potential therapeutic targets

The use of antiarrhythmic agents in AF treatment and prevention of recurrences relies on the ability of these drugs to block I_{Na} (class I antiarrhythmic agents), I_{Ca-L} (class IV antiarrhythmic agents), and outward K⁺ currents (class III antiarrhythmic agents). However, among the genes that were most markedly up-regulated in our study was *Hcn4*, encoding for the I_s current. Moreover, several other studies have demonstrated abnormal presence of I_f in atrial myocytes, with altered expression of HCN isoforms in human AF. In the study of Lai et al. mRNA levels of I_f were significantly correlated with left atrial filling pressure and AF (29), while Morel et al. demonstrated HCN4 expression in interstitial human pulmonary veins Cajal cells capable of generating electrical activity (30). Therefore, it seems reasonable to hypothesize that I_{e} blockade might carry promising therapeutic potential, at least in some conditions that increase AF susceptibility. Several experimental studies have already demonstrated that I_f blockade with Ivabradine is able to suppress spontaneous activities in isolated pulmonary veins (31, 32). Since I_r is stimulated by *beta*-adrenergic activation of the adenylate cyclase-cAMP system, the combination of *beta*-blockers and specific I_f current blockers is indeed expected to provide beneficial effects in patients with sympathetic AF. However, it seems unlikely for I_f blockade to be efficient in other AF cases, such as vagally-mediated AF (33). The use of I_{f} channel blockers in AF prevention and/or treatment is an emergent therapeutic strategy that deserves to be further studied.

Limitations

In this study, gene expression changes were only assessed by mRNA quantification. Although altered gene expression is expected to result in abnormal protein variations, other posttranscriptional or posttranslational regulatory mechanisms may influence protein level and function.

Conclusions

Our results indicate the induction of an up-regulation transcriptional response in atrial, but not ventricular myocytes with advancing age, suggesting that the two chambers of the heart undergo different molecular remodeling programs. Aging atria displayed a transcriptomic profile consistent with higher propensity to arrhythmias, including I_{p} , I_{Ca-L} , I_{Ca-T} , I_{Na} , outward K⁺ currents, and collagen up-regulation, while ventricular transcriptome did not seem to be significantly altered by aging. Taken together, these observations could explain the higher propensity to atrial than ventricular arrhythmias in the elderly with no concomitant cardiac disorders.

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Abbreviations

$$\begin{split} AF &= \text{atrial fibrillation} \\ cDNA &= \text{complementary DNA} \\ I_{Ca-L} &= L\text{-type } Ca^{2+} \text{ current} \\ I_{Ca-T} &= T\text{-type } Ca^{2+} \text{ current} \\ I_{K} &= \text{delayed rectifier } K^{+} \text{ current} \\ I_{K1} &= \text{inward rectifier } K^{+} \text{ current} \\ I_{K-Ach} &= \text{acetylcholine-activated } K^{+} \text{ current} \\ I_{K-ATP} &= ATP\text{-sensitive } K^{+} \text{ current} \end{split}$$

 I_{Na} = voltage-dependent Na⁺ current I_{to} = transient outward K⁺ current TGF β = transforming growth factor *beta* TLDA = TaqMan Low Density Array WKY = Wistar-Kyoto

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