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Identification of atrial-enriched lncRNA *Walras* linked to cardiomyocyte cytoarchitecture and atrial fibrillation

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Abstract

Atrial fibrillation (AF) is the most prevalent cardiac arrhythmia in humans. Genetic and genomic analyses have recently demonstrated that the homeobox transcription factor Pitx2 plays a fundamental role regulating expression of distinct growth factors, microRNAs and ion channels leading to morphological and molecular alterations that promote the onset of AF. Here we address the plausible contribution of long non-coding (lnc)RNAs within the Pitx2>Wnt>miRNA signaling pathway. In silico analyses of annotated lncRNAs in the vicinity of the Pitx2, Wnt8 and Wnt11 chromosomal loci identified five novel lncRNAs with differential expression during cardiac development. Importantly, three of them, Walaa, Walras, and Wallrd, are evolutionarily conserved in humans and displayed preferential atrial expression during embryogenesis. In addition, Walrad displayed moderate expression during embryogenesis but was more abundant in the right atrium. Walaa, Walras and Wallrd were distinctly regulated by Pitx2, Wnt8, and Wnt11, and Wallrd was severely elevated in conditional atrium-specific Pitx2deficient mice. Furthermore, pro-arrhythmogenic and pro-hypertrophic substrate administration to primary cardiomyocyte cell cultures consistently modulate expression of these lncRNAs, supporting distinct modulatory roles of the AF cardiovascular risk factors in the regulation of these lncRNAs. Walras affinity

Abbreviations: AF, atrial fibrillation; AngII, angiotensin II; ECG, electrocardiogram; GWAS, genome-wide association study; HTD, hyperthyroidism; HTN, hypertension; LC/MS/MS, liquid chromatography/mass spectrometry/mass spectrometry; lncRNA, long non coding RNA; MS, mass spectrometry; NE, norepinephrine.

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pulldown assays revealed its association with distinct cytoplasmic and nuclear proteins previously involved in cardiac pathophysiology, while loss-of-function assays further support a pivotal role of this lncRNA in cytoskeletal organization. We propose that lncRNAs *Walaa*, *Walras* and *Wallrd*, distinctly regulated by Pitx2>Wnt>miRNA signaling and pro-arrhythmogenic and pro-hypertrophic factors, are implicated in atrial arrhythmogenesis, and *Walras* additionally in cardiomyocyte cytoarchitecture.

KEYWORDS

atrial fibrillation, lncRNA, post-transcriptional regulation

1 | INTRODUCTION

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Atrial fibrillation (AF) is the most common cardiac arrhythmia, with an incidence of 2%-3% in the general population that rises up to 8%-10% in the elderly. Genetic linkage analyses have identified point mutations in several ion channels with key roles in the configuration of the cardiac action potential, suggesting that they are implicated in AF. However, these genetic defects only explain a minority (<10%) of all AF cases. Seminal work by Gudbjartsson et al.¹ using genome-wide association studies (GWAS) revealed that risk variants in the 4q25 locus were highly associated to lone AF, postulating that regulatory elements within this locus might influence expression of a neighboring gene encoding the homeobox transcription factor PITX2, in turn leading to impaired cardiac function and AF. Subsequent GWAS studies and meta-GWAS analyses have increased exponentially our understanding of the plausible genetic substrates of AF, with the identification of more than 90 risk variants associated to AF.² Importantly, in all cases the most significant risk variants were those in the 4q25 locus. Experimental studies in mice have provided evidence that 4q25 physically interacts with the *Pitx2* promoter,^{3,4} supporting key regulatory roles as suggested by Gudbjartsson et al.¹ In addition, systemic Pitx2 loss-of-function experiments revealed increased susceptibility to atrial arrhythmias,^{5,6} while conditional atrial-specific deletion of Pitx2, resulting in *Pitx2* insufficiency, led to spontaneous atrial arrhythmias.7

Subsequent studies demonstrated that *Pitx2* insufficiency led to remodeling of several meta-GWASassociated genes, such as that encoding Wnt8, which in turn modulate Wnt11 expression, leading to microRNA deregulation and hence impaired ion channels expression and function.^{7–9} These data demonstrate a pivotal role of the *Pitx2>Wnt>microRNA* pathway modulating AF. In addition, experimentally induced AF leads to *Pitx2* downregulation, supporting the notion of a self-perpetuating reduction of *Pitx2* during AF progression.¹⁰ Moreover, cardiovascular risk factors contributing to increased frequency of AF onset in the human population, such as hypertension, hyperthyroidism and antioxidant redox imbalance, consistently altered the *Pitx2>Wnt>microRNA* pathway, underscoring a molecular link between these AF risk factors and specific cellular processes.¹¹

The past two decades have revealed that gene regulatory networks are profoundly influenced by novel types of non-coding RNAs with regulatory potential. There is now extensive evidence that microRNAs are essential regulators of cardiovascular development and diseases, including seminal studies on the role of microRNAs regulating cardiac ion channels and AF itself. The discovery of long non coding RNAs (lncRNAs) has increased the complexity of the non-coding RNA regulatory roles. LncRNAs generally have no protein-coding potential, yet many of them are structurally similar to mRNAs, including the presence of 5' terminal cap and 3' terminal poly(A) tails and their transcription by RNA polymerase II.¹² Similarly, most lncRNA genes comprise exons and introns and are often spliced. LncRNAs display lower expression levels and are more tissue-specific than protein-coding mRNAs,¹³ supporting the notion that they may have tightly defined roles in different cellular events.^{14,15} LncRNAs can be localized in both cytoplasm and nucleus; cytoplasmic lncRNAs are preferentially involved in post-transcriptional regulation while nuclear lncRNAs are predominantly involved in transcriptional regulation. Interestingly, lncRNAs can translocate from the nucleus to the cytoplasm and act therein.16

Distinct studies have investigated the functional role of cardiac-enriched lncRNAs during heart development, i.e., *Carmen, Braveheart*, and *Fendrr. Carmen*, miR-143 and miR-145 are located within the same genomic locus, but they are independently expressed.^{17,18} Expression of *Carmen* in fetal and adult hearts is highly conserved in different mammalian species. Functionally, *Carmen* plays

a pivotal role at the earliest lineage commitment steps, modulating cardiac differentiation from initial mesoderm by regulating *Mesp1* expression.¹⁷ Similarly, *Braveheart* is a key regulator in cardiac lineage commitment and its ablation leads to defective activation of key cardiac factors, heart development, and cardiomyocyte differentiation.¹⁹ *Fendrr* is transiently expressed at the posterior end of the forming lateral plate mesoderm, and is required for the proper development of the heart and body walls.^{20,21} We have recently demonstrated that these lncRNAs display a dynamic chamber-specific expression, are distinctly regulated by cardiac enriched transcription factors such as *Mef2c, Srf* and *Nkx2.5* and display distinct isoform usage during cardiogenesis.²²

In the context of AF, several reports have provided evidence of altered expression of certain lncRNAs^{23–26} but scarce information is available about their tissue distribution and regulatory mechanisms. Gore-Panter et al.²⁷ identified *PANCR* as an intergenic lncRNA expressed in the adult left atrium from a region adjacent to *PITX2*, and is expressed in coordination with *PITX2C* mRNA during cardiomyocyte differentiation. Importantly, *PANCR* is expressed in human tissues but no orthologues have been found in other species.²⁷

In this study, we have investigated the expression of lncRNAs via processes regulated by Pitx2>Wnt>miRNA. In silico analyses of annotated lncRNAs expressed from the vicinity of Pitx2, Wnt8 and Wnt11 chromosomal loci identified five novel lncRNAs showing differential expression levels during cardiac development in mice and some of them are also evolutionarily conserved in humans. Three of them displayed preferential atrial-specific expression during embryogenesis and were distinctly regulated by Pitx2, Wnt8 and Wnt11 as well as by microR-NAs such as miR-1, miR-133 and miR-29. Furthermore, pro-arrhythmogenic and pro-hypertrophic substrates such as angiotensin II, norepinephrine and thyroid hormone administration distinctly regulate their expression. Walras pulldown and loss-of-function assays further demonstrate a key role of this lncRNA in cardiomyocyte cytoskeletal organization. In summary, we have identified novel lncRNAs with enhanced atrial expression that are distinctly regulated by signaling pathways leading to atrial arrhythmogenesis.

2 | MATERIALS AND METHODS

2.1 | Mouse breeding and tissue sampling

CD1 mice were bred and embryos were collected at different embryonic developmental stages, ranging from



3 of 24

embryonic day (E) E12.5 to E18.5. Neonatal (1 day old) and adult (>6 months) hearts were also collected. Pregnant females, adult and neonatal mice were euthanized by cervical dislocation. Neonates were previously placed on ice. Subsequently, embryonic and postnatal hearts were dissected into right atrium, left atrium and ventricular chambers, pooled and stored in liquid nitrogen until used. Additionally, distinct tissues such as spleen, liver, lungs, small and large intestine, kidney and brain were also dissected from adult mice, pooled and stored in liquid nitrogen until used.

The Pitx2^{floxed} and NppaCre transgenic mouse lines and the generation of conditional atrial (NppaCre) mutant mice were previously described.^{7,8,28,29} Two different conditions were used for the NppaCrePitx2 mice: wildtype Cre controls (NppaCre⁻Pitx2^{fl/fl}) and atrial-specific homozygous (NppaCre⁺Pitx2^{-/-}). This investigation conforms with the Guide for the Care and Use of Laboratory Animals published by the US National Institutes of Health. The study was approved by the University of Jaén Bioethics Committee.

2.2 | Mouse genotyping and phenotyping

DNA for PCR screening was extracted from adult ear and/or tail samples. Screening of Cre and Pitx2 floxed alleles was routinely done using used specific primers as previously described.⁷ Cycling conditions for *Cre* DNA were as follows: 5 min at 95°C, 35 cycles of 30 s at 95°C, 30 s at 60°C and 90 s at 72°C: and for *Pitx2* DNA as follows: 5 min at 95°C, 40 cycles of 30 s at 95°C, 30 s at 60°C and 90 s at 72°C, followed by a final extension step of 10 min at 72°C. In addition, the expression levels of *Pitx2* mRNA in left atrial samples of wild-type Cre controls (NppaCre2Pitx2^{fl/fl}) and atrial-specific homozygous (NppaCre⁺Pitx2^{-/-}) were assessed by RT-qPCR analysis, displaying in all cases 60%–70% reduction in PITX2c expression in NppaCre⁺Pitx2^{-/-} samples.

2.3 | Mouse tissue samples

Genetically modified transgenic Pitx2 mice and their corresponding controls were sacrificed by cervical dislocation. Adult hearts were carefully dissected and briefly rinsed in Ringer's solution. Left atrium tissue samples were collected for each experimental condition, immediately snap-frozen in liquid nitrogen, and stored at -80° C until used. Pooled samples of at least three independent mice were processed for each condition, respectively. Three independent pooled samples were further processed for RNA isolation and RT-qPCR analysis.

2.4 | Human atrial biopsies

Atrial myocardial tissue samples were obtained from patients undergoing cardiac surgery at Hospital de la Santa Creu i Sant Pau (Barcelona). Specimens were obtained from the right atria just prior to atrial cannulation for cardiopulmonary bypass. After excision, samples were rapidly frozen in liquid nitrogen and stored at -80°C until analyzed. The atrial samples were classified as patients with (AF; n = 4) and without (No AF; n = 3) a recorded history of AF. Clinical data are presented in Table S1. Although the atrial tissue samples consisted of tissue that would normally be discarded during surgery, permission to be used in this study was obtained from each patient. The study was approved by the Ethical Committee of the Hospital de la Santa Creu i Sant Pau (Barcelona) and the investigation conforms with the principles outlined in the Declaration of Helsinki.

2.5 | RNA isolation and cDNA synthesis

Total RNA was isolated using Trizol (Roche) according to manufacture's guidelines and DNase treated using RNase-Free DNase (Roche) for 1 h at 30°C. In all cases, at least three distinct pooled samples were used to perform the corresponding RT-qPCR experiments.

First strand cDNA was synthesized at 50°C for 1 h using 1 μ g of RNA, oligo-dT primers and Superscript III Reverse Transcriptase (Invitrogen) according to manufacture's guidelines. Negative controls to assess genomic contamination were performed for each sample, without reverse transcriptase, which resulted in all cases in no detectable amplification product.

2.6 | qPCR analyses (mRNA and lncRNA)

RT-PCR was performed in Mx3005Tm QPCR System with an MxPro QPCR Software 3.00 (Stratagene) and SyBR Green detection system. Reactions were performed in 96well plates with optical sealing tape (Cultek) in 20 µl total volume containing SYBR Green Mix (Finnzymes) and the corresponding cDNA. Two internal controls, mouse *Gusb* and *Gapdh* mRNAs, were used in parallel for each run and represented as previously described.^{8,9} Amplification conditions were as follows: denaturalization step of 95°C for 10 min, followed by 40 cycles of 95°C for 30 s, 60°C for 30 s, 72°C for 30 s; with final elongation step of 72°C for 10 min. All primers were designed to span exon-exon boundaries using online Primer3 software Primer3input (http://bioinfo.ut.ee/primer3-0.4.0/). Primer sequences are provided in Table S2. No amplifications were observed in PCR control reactions containing only water as the template. Each PCR reaction was performed at least three times to obtain representative averages. The Livak method was used to analyze the relative quantification RT-PCR data³⁰ and normalized in all cases taking as 100% the wild-type (control) value, as previously described.⁷

2.7 | qPCR analyses (microRNA)

microRNA RT-qPCR was performed using Exiqon LNA microRNA reverse transcription (RT) followed by realtime quantitative (q) PCR primers and detection kit according to the manufacturer's guidelines. All reactions were always run in triplicate using *5S* as normalizing control, as recommended by the manufacturer. SyBR Green was used as quantification system on a Stratagene Q-Max 2005P RT-qPCR thermocycler. Relative measurements were calculated as described by Livak and Schmittgen³⁰ and control measurements were normalized to represent 100% as previously described.⁷

2.8 | Plasmid, siRNA, microRNA mimics cell transfections

HL1 cells (6×10^5 cells per well) were transfected with plasmids containing expression constructs for Pitx2c, Wnt8a (Addgene), Wnt11a (Addgene, Cambridge, MA, USA) and with pre-miR-1, pre-miR-133, pre-miR-29 (Exiqon) or siRNA-Pitx2c, siRNA-Wnt8a, siRNA-Wnt11a, siRNA-Gm44934/*Walras* (Sigma, Aldrich, Munich, Germany), respectively, as previously described.^{6,7} siRNA sequences are provided in Table S3.

2.9 | Immunofluorescence analyses

Control and experimentally siRNA treated cells were collected after the corresponding treatment, rinsed in PBS for 10 min at room temperature, and fixed with 1% PFA for 2 h at 4°C. After fixation, the samples were rinsed three times (10 min each) in PBS at room temperature and then permeabilized with 1% Triton X-100 in PBS for 30 min at room temperature. To block nonspecific binding sites, PBS containing 5% goat serum and 1% bovine serum albumin (Sigma) was applied to the cell cultures overnight at 4°C. Phalloidin-Atto 488 (Sigma) was used, diluted (5 mg/ ml) in PBS, and applied to each culture overnight at 4°C. Subsequently, the samples were rinsed three times (for 1 h each) in PBS to remove excess Phalloidin-Atto 488. Finally, they were incubated with DAPI (1:1000; Sigma) for 7 min at room temperature and rinsed three times in PBS for 5 min each. Alternatively to Phalloidin-Atto 488 staining, control and experimentally siRNA treated cells were immunofluorescently labelled to detect ACTN1, ACTN4 and MHY9 expression, respectively. Primary antibodies against ACTN1 (ab68194), ACTN4 (ab108198), MYH9 (ab75590) were used, diluted (1:200) in PBS, and applied to each culture overnight at 4°C, respectively. Subsequently, the samples were rinsed three times (for 1 h each) in PBS to remove excess primary antibody and incubated 2 hours at room temperature with Alexa-Fluor 546, 488 and 633 anti-rabbit (1:100; Invitrogen) as secondary antibody. Cell cultures were stored in PBS in darkness at 4°C until analysed using a Leica TCS SP5 II confocal scanning laser microscope.

2.10 | Cell culture and angiotensin II, norepinephrine and thyroid hormone treatment

Primary cultures of mouse fetal (E17.5) cardiomyocytes were isolated using standard procedures,³¹ cultured accordingly and treated with 1 μ M angiotensin II (AngII), 100 μ M norepinephrine (NE) and 10 nM thyroid T3/T4 hormone, respectively, as previously reported.^{9,32,33}

2.11 | Nuclear/cytoplasmic distribution

Cytoplasmic and nuclear RNA fractions from atrial and ventricles samples were isolated with Cytoplasmic & Nuclear RNA Purification Kit (Norgen, Belmont, CA, USA) following the manufacturer's instructions. After RNA isolation, RT-qPCR analysis for nuclear enriched *Rpb1* mRNA marker and cytoplasmic *Gapdh* mRNA marker were performed to validate enrichment on each subcellular fractions. RT-qPCR analysis of distinct lncR-NAs was subsequently performed as detailed above.

2.12 | IncRNA pull down assays

Biotinylated RNA of exon 1 and exon2 of Gm_44934 of Exon 1 and Exon 2 were synthesized from PCR fragment using specific forward primers that contained the T7 RNA polymerase promoter sequence [(T7), CCAAGCTTCTAATACGACTCACTATAGGGAGA]. After purifying the DNA template, biotinylated transcripts were synthesized using MaxiScript T7 kit (Ambion); whole-cell lysates (500 µg) from HL1 cells were incubated with 1 µg of biotinylated RNA for 2 h at room temperature. Complexes were isolated with Streptavidin-coupled Dynabeads (Invitrogen) and analyzed by MS.



For MS analysis, peptide mixtures from each sample were loaded onto a peptide trap cartridge and eluted onto a reversed-phase PicoFrit column (New Objective, Woburn, MA, USA). Eluted peptides were ionized and sprayed into the mass spectrometer, using a Nanospray Flex Ion Source ES071 (Thermo Scientific). The LC/MS/MS analysis of samples were carried out using a Thermo Scientific Q-Exactive hybrid Quadrupole-Orbitrap Mass Spectrometer and a Thermo Dionex UltiMate 3000 RSLCnano System. Proteins were identified using the Thermo Proteome Discoverer 1.4.1 platform. Database search against public mouse protein database from NCBI was performed through the Proteome Discoverer 1.4.1 platform.

2.13 | miRNA pull down assays

Biotinylated miR-29 was synthesized by IDT technologies. Whole-cell lysates (500 µg) from HL1 cells were incubated with 1 µg of biotinylated RNA for 2 h at room temperature. Complexes were isolated with Streptavidin-coupled Dynabeads (Invitrogen) following the protocol published by Panda et al.³⁴ RNA associated to biotinylated miR-29 was isolated using *Direct-zol*TM *RNA Miniprep Plus* (Zymo research) following the manufacturer's instructions and RT-qPCR was subsequently performed as detailed above. Biotinylated miR-23 was used as a negative control.

2.14 | RIP assay

For immunoprecipitation of endogenous ACTN1, ACTN4 and MHY9 cytoskeleton proteins, HL1 cells were lysed with PEB buffer for 10 min on ice and centrifuged at 10 000 g for 15 min at 4°C. The supernatants were incubated with protein A Sepharose beads (Abcam) coated with antibodies that recognized ACTN1, ACTN4, MYH9 or control IgG (Abcam) for 2 h at 4°C, respectively. The corresponding beads were washed with NT2 buffer (50 mM Tris-HCl [pH 7.5], 150 mM NaCl, 1 mM MgCl₂, 0.05% NP-40). Protein complexes were incubated with 20 units of DNase I (15 min at 37°C). In this step, an aliquot from each reaction was isolated for Western blot validation. Subsequently they were further incubated with 0.1% SDS/0.5 mg/ml Proteinase K (30 min at 55°C) to remove DNA and proteins, respectively. The RNA isolated from the IP materials were further assessed by RT-qPCR analysis.

2.15 | Western blot

For western blot analysis, 10% of total IP lysate was used. Primary antibodies recognizing ACTN1 (ab68194),

SEBJOURNAL

ACTN4 (ab108198), MYH9 (ab75590) from Abcam were used. After incubation with the rabbit secondary antibodies, protein signals were developed using HRP.

2.16 | Statistical analyses

For statistical analyses of datasets, unpaired Student's *t*-tests were used. Significance levels or *p* values are stated in each corresponding figure legend. p < .05 was considered statistically significant.

3 | RESULTS

3.1 | Identification of novel lncRNAs putatively involved in the Pitx2>Wnt>microRNA pathway

Previous studies have demonstrated that some long non coding RNAs act locally regulating neighboring genes.^{35–38} We therefore examined the mouse genome to identify lncRNAs in close vicinity to genes previously identified to play a pivotal role in the onset of AF, particularly Pitx2>Wnt>microRNA pathway.⁷⁻⁹ In particular, we searched for lncRNAs expressed from regions close to the Pitx2, Wnt8 and Wnt11 loci. We did not find novel lncRNAs near the Pitx2 locus, besides the previously reported *Playrr* lncRNA,³⁹ but we identified two lncRNAs adjacent to Wnt8 and four near Wnt11 genes, respectively (Figure 1A; Table 1). Three of them, i.e., Walras, Wallrd and Walce are evolutionarily conserved in humans (Figure S1). In particular, Walras displayed significant homology, ranging from 25% to 43%, with three distinct human lncR-NAs (AP0002340.1, AP000785.1 and LINC02761). Wallrd isoforms also displayed significant homology, ranging from 33% to 45%, with AC13382.1, AC13382.2 and AC104116.1, respectively. Finally, Walce display significant homology (35%) with Wnt11_03 human lncRNA (Figure S1).

We then measured the levels of expression of these lncRNAs in the right and left atria and in ventricular tissues of mouse embryos ranging from E12.5 to E18.5, neonate and adult tissues. Among the six lncRNAs interrogated, including *Playrr*, only five of them were detectable in cardiac tissues (Figure 1B–F). Three of them (*Walaa*, *Walras* and *Wallrd*) displayed a preferential embryonic expression confined to the atrial chambers with only low ventricular expression. Interestingly, *Walaa* displayed a similar atrial-specific expression in both right and left atrial chambers, peaking at E16.5 (Figure 1B), *Walras* displayed a left atrial-restricted expression (Figure 1D), with peak levels at E16.5, while *Wallrd* also displayed atrialrestricted expression, preferentially in the left atrium, at

E16.5 (Figure 1F). In all cases, expression in neonatal and adult heart was markedly lower than in embryonic heart. Importantly, Walrad displayed preferential expression in the right atrium in all stages analyzed (Figure 1C), including adulthood, but also presented a transient peak expression at neonatal stages in the ventricular chambers. By contrast, Walce was preferentially expressed in embryos but was not restricted to the atrial chambers (Figure 1E), showing a dynamic expression pattern in right atrium, left atrium and ventricular tissues at different developmental time points. Playrr displayed a prominent expression in the early developmental stages declining during development (Figure 1G). Curiously, expression of these lncRNAs is significantly different from the expression of Wnt8 and Wnt11 mRNAs expression during cardiogenesis, as illustrated in Figure 1H-I. Expression of these five distinct lncRNAs is not restricted to the heart, since all but Walce are also expressed in distinct adult tissues such as spleen, liver, lungs, gut, kidney and brain as depicted in Figure 2A-F. Walce expression is restricted to the heart and spleen (Figure 2D). Importantly, Wnt8 mRNA is also broadly expressed in distinct adult tissues (Figure 2H), while Wnt11 mRNA is primarily expressed in the heart and brain, with low, but detectable levels in spleen and lungs (Figure 2G). In summary, we identified novel lncRNAs with wide distribution in adult tissues and importantly, three of them, Walaa, Walras and Wallrd, display preferential expression in the embryonic atrial chambers and one of them, Walrad, with preferential expression in the adult right atrium during cardiogenesis. Furthermore, their expression profile, both in different adult tissues as well as during cardiogenesis, is markedly distinct from those of Wnt8 and Wnt11 mRNAs, supporting the notion that they might not directly regulate Wnt expression.

3.2 Differential spatio-temporal nuclear vs cytoplasmic localization of these novel lncRNAs support both transcriptional and post-transcriptional regulatory function

LncRNAs can reside in both the nucleus and the cytoplasm, exerting distinct regulatory functions in each compartment. We thus measured the relative abundance of these lncRNAs in the nucleus and cytoplasm by RT-qPCR analysis using *Rpb1* mRNA for normalization²² (Figure S2). The nuclear and cytoplasmic distribution was analyzed in three distinct cardiac compartments of E14.5 mouse embryos and neonatal mice: right atrium, left atrium and ventricles. *Walas* displayed similar nuclear *vs* cytoplasmic distribution in the left atrium in both embryonic and adult stages, while it was prominently nuclear in the embryonic and adult ventricles. Importantly, a rise in cytoplasmic levels of

FASEB JOURNAL

7 of 24

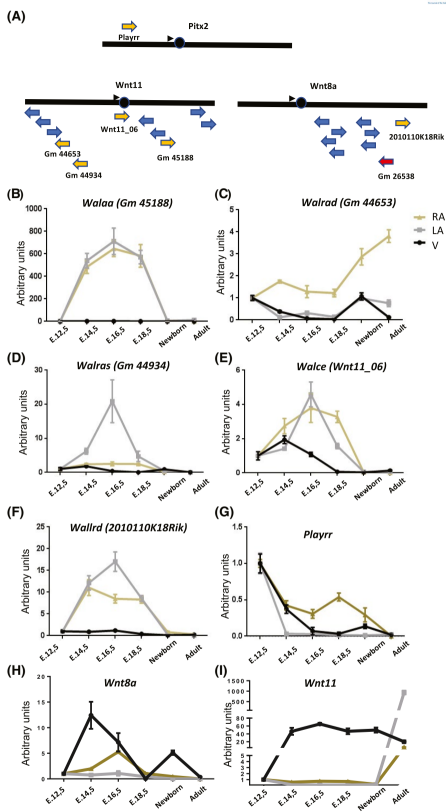


FIGURE 1 (A) Schematic representation of the mouse chromosomal localization of lncRNAs situated in the vicinity of *Pitx2*, *Wnt11* and *Wnt8* genes. (B–G) RT-qPCR analyses of *Walaa*, *Walras*, *Wallrd*, *Walrad*, *Walce* and *Playrr* expression levels. We note observe atrial expression of *Walaa*, *Walras* and *Wallrd* during cardiogenesis, while *Walrad* is mainly expressed in the left atrium in the adulthood. (H–I) Rt-qPCR analyses of Wnt11 and Wnt8a expression during cardiogenesis. Three embryonic hearts were pooled on each developmental time and three distinct neonatal and adult hearts were used for RNA isolation, respectively. Three independent biological assays were analyzed in each case

TABLE 1 Transcripts and gene localization data of novel identified lncRNAs

8 of 24

LncRNA name (new)	LncRNA name (previous)	Transcript number	Туре	Cromosome position	Isoforms	Bps
Walras	Gm 44934	ENSMUST00000208941.1	LincRNA	Chromosome 7: 98,734,747- 98,735,081 reverse strand	1	238
Walaa	Gm 45188	ENSMUST00000207592.1	LncRNA	Chromosome 7: 98,895,930- 98,899,545 forward strand	1	1369
Walrad	Gm 44653	ENSMUST00000208543.1	LncRNA	Chromosome 7: 98,720,057- 98,728,760 reverse strand	1	4610
Walce	Wnt11_06	ENSMUST00000208062.1	LncRNA	Chromosome 7: 98,835,112- 98,855,195 forward strand.	1	745
Wallrd	2010110K18Rik	ENSMUST00000181453.2	LncRNA	Chromosome 18: 34,751,809- 34,758,685 forward strand	3	2398
		ENSMUST00000181641.1 ENSMUST00000236940.1		54,750,005 for ward strand		678 490

Walas was observed from embryonic to neonatal stages in the right atrium (Figure 3A). Walrad was prominently nuclear in both embryonic and neonatal stages in the left and right atria, while similar nuclear/cytoplasmic distribution was observed in the embryonic and neonatal ventricles (Figure 3B). Walras displayed no significant differences in the left atrium at both stages analyzed, while in the right atrium and ventricle it was prominently nuclear in the embryonic stages whereas there were no significant differences in the neonatal stage (Figure 3C). Walce displayed a prominent nuclear distribution in the left atrium at both stages, but no significant differences in the right atrium and ventricles (Figure 3D). Wallrd displayed similar nuclear and cytoplasmic expression levels in the embryonic and adult left atrium and ventricles. Importantly, however, Wallrd shifted from being primarily cytoplasmic in the embryonic right atrium to being mostly nuclear in the adult ventricles (Figure 3E). Finally, Playrr displays similar nuclear vs cytoplasmic distribution at all tissues and stages analyzed except in the embryonic left atrium where it is prominently nuclear (Figure 3F). Overall, these data demonstrate distinct spatio-temporal and subcellular distribution for these lncRNAs, suggesting chamber-specific shifts in their transcriptional vs post-transcriptional regulatory roles during cardiac development.

3.3 | LncRNA regulatory roles exerted by Pitx2, Wnt8 and Wnt11

To dissect the functional roles of Pitx2, Wnt8 and Wnt11 in the regulation of these newly identified lncRNAs, overexpression and silencing experiments were carried out in atrial HL1 cardiomyocytes.

Pitx2c overexpression and silencing significantly upregulated *Walaa*, *Walrad* and *Walce*, downregulates *Wallrd* while *Walras* display no significant differences (Figure 4A). Wnt8a overexpression significantly decreased all lncRNAs analyzed except Wallrd. Wnt8a siRNA silencing significantly decreased Walaa and Walras, upregulated Walce while Walrad and Wallrd displayed no significant differences (Figure 4B). On the other hand, Wnt11 overexpression significantly upregulated all lncRNAs analyzed except Walaa, which was downregulated. Importantly, silencing Wnt11 using small interfering (si)RNA also upregulated all lncRNAs analyzed except Walras (Figure 4C). Control experiment of gain- and loss-of-function are provided in Figure 4A-C (right). These data indicate that Pitx2, Wnt8a and Wnt11 influenced the expression levels of these lncRNAs. Moreover, Pitx2 silencing upregulated Walrad and Walce, while Wnt8 and Wnt11 overexpression displayed complementary patterns on the regulation of Walrad and Walce expression, in line with our previous findings in $NppaCrePitx2^{-/-}$ mice.⁷ In addition, it is importantly to note that both Pitx2 overexpression and silencing similarly affected lncRNAs, demonstrating that even subtle changes in Pitx2 expression greatly influenced lncRNA abundance, although the precise regulatory molecular mechanisms remain to be elucidated.

To further support the notion that Pitx2 regulates the expression of these lncRNAs, we measured lncRNA levels of these lncRNAs in conditional Pitx2 mouse model $NppaCrePitx2^{-/-}$. Left atrial samples were used to verify Pitx2 loss-of-function by measuring *Pitx2c* mRNA levels using RT-qPCR analysis (Figure 4D). As previously reported, $NppaCrePitx2^{-/-}$ mice displayed 70%–80% reduction of *Pitx2c* mRNA expression levels in the left atrial chamber (Figure 4D, right), leading to Pitx2 insufficiency. In this context, *Walaa* was severely upregulated, *Wallrd* moderately upregulated, while *Walrad* was downregulated, and *Walras* and *Walce* displayed no significant differences in abundance (Figure 4D). These data support our earlier observations in HL1 cardiomyocytes that Pitx2 regulates *Walaa* but not *Walras*. The surprising finding that *Walrad*



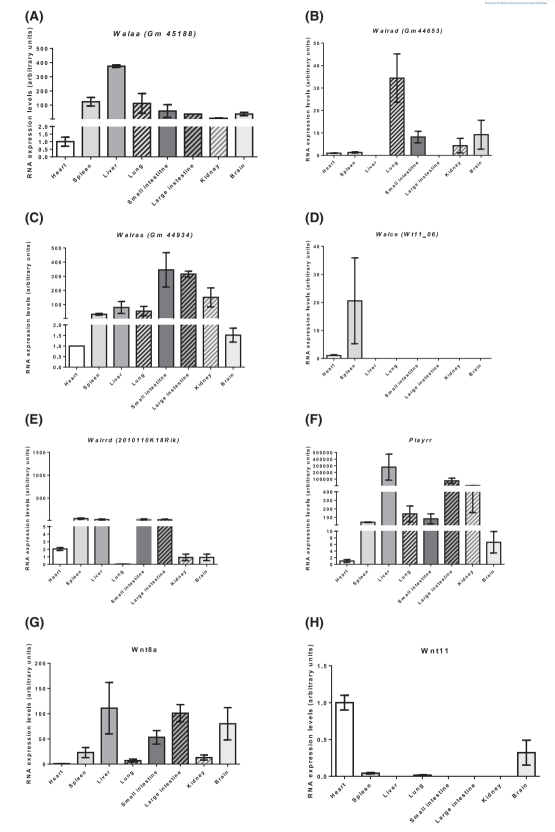


FIGURE 2 (A–H) RT-qPCR analyses of *Walaa*, *Walras*, *Walrad*, *Walrad*, *Walrad*, *Walce*, *Playrr*, *Wnt8a*, and *Wnt11* expression levels in different organs (heart, spleen, liver, lung, small intestine, large intestine, kidney, and brain) in the adult mouse. Note that *Walaa*, *Walras*, *Wallrd*, *Walrad*, *Playrr*, and *Wnt8a* are widely distributed in different organs, while *Walce* is essentially confined to the spleen and heart, and *Wnt11* to the heart and brain with just detectable levels in the spleen and lungs. Three independent adult tissues were used for RNA isolation, respectively. Three independent biological assays were analyzed in each case

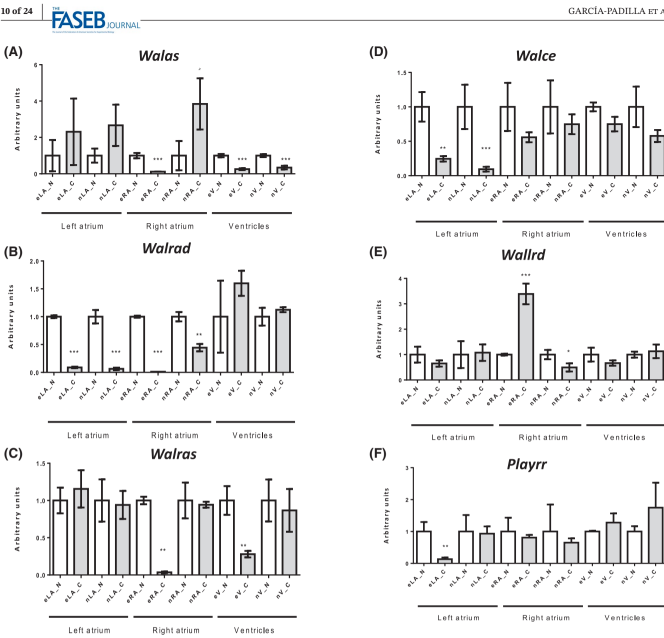


FIGURE 3 (A-F) Nuclear vs cytoplasmic distribution of Walas, Walrad, Walras, Walce, Wallrd and Playrr in the embryonic and neonatal right atrium, left atrium and ventricles, respectively. Differential nuclear vs cytoplasmic subcellular distribution was observed for Walas, Walras and Wallrd in the right atrium of both embryonic and neonatal hearts, while Walras only displayed it in the embryonic right atrium. Walas displayed enhanced nuclear localization in the embryonic and neonatal ventricles while Walras only did in the embryonic ventricle. Walrad and Walce displayed enhanced nuclear vs cytoplasmic localization in embryonic and neonatal left atrium. Five to ten embryonic right atrial, left atrial or ventricles were pooled on each developmental staged, respectively and used for RNA isolation. Three independent biological assays were performed on each case. Statistical analyses was performed using unpaired Student's t-test. *p < .05, ***p* < .01, ****p* < .005. eLA_N, embryonic left atrium, nuclear, eLA_C, embryonic left atrium, cytoplasmic, eRA_N, embryonic right atrium, nuclear, eRA_C, embryonic right atrium, cytoplasmic, eV_N, embryonic ventricle, nuclear, eV_C, embryonic ventricle, cytoplasmic, nLA_N, neonatal left atrium, nuclear, nLA_C, embryonic left atrium, cytoplasmic, nRA_N, neonatal right atrium, nuclear, nRA_C, neonatal right atrium, cytoplasmic, nV_N, neonatal ventricle, nuclear, nV_C, neonatal ventricle, cytoplasmic

and Wallrd display opposite patterns in NppaCrePitx2^{-/-} mice and HL1 Pitx2 siRNA assays demonstrate that additional studies are required to clarify this point.

Since we identified plausible human lncRNAs homologues for Walras, Wallrd and Walce, we also investigated if these conserved lncRNAs are deregulated in AF patients. Right atrial biopsies of AF and no AF patients were subsequently scrutinized. Our data demonstrate that AP00785.1 displayed no significant differences while AP002340.1 is significantly decreased while AC1040116.1 was significantly increased in AF as compared to no AF patients (Figure 4E). The other identified human homologues displayed no detectable expression (data not shown). Therefore these data further confirmed the upregulation

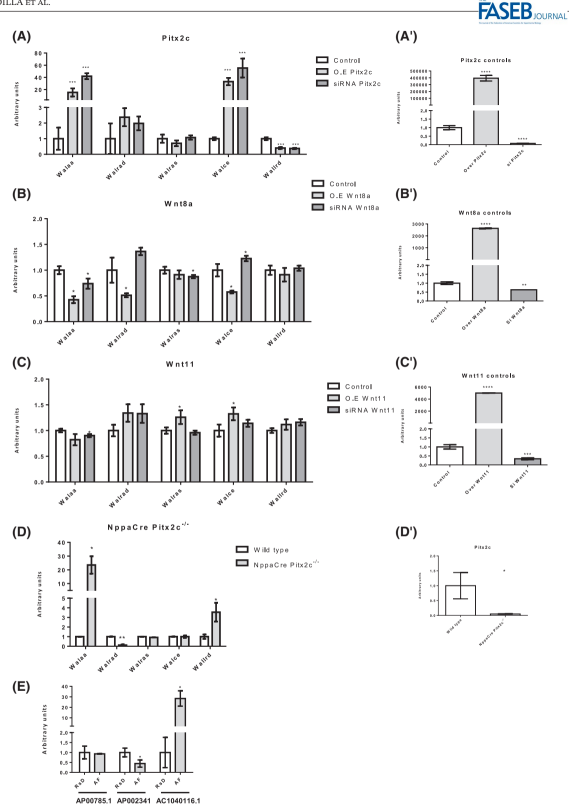


FIGURE 4 (A–C) *Right*, RT-qPCR analyses of the levels of *Walaa*, *Walrad*, *Walras*, *Walce* and *Wallrd* after overexpression or silencing of *Pitx2*, *Wnt8* and *Wnt11*, respectively. *Left*, levels of *Pitx2*, *Wnt8* and *Wnt11* mRNAs after overexpression and silencing. (D) *Right*, Expression levels of *Walaa*, *Walras* and *Wallrd* in NppaCrePITX2^{-/-} adult left atrial chambers; *Left*, *Pitx2c* mRNA levels in NppaCrePITX2^{-/-} adult left atrial chambers. (E) Expression levels of AP00785.1, AP0002340.1 (*Walras* human homologue) and AC1040116.1 (*Wallrd* human homologue) in human right atrial biopsies of patients with sinus rhythm (RsD) (i.e., no AF) and atrial fibrillation (AF), respectively. Three independent siRNA and overexpression assays were analyzed in each case. Similarly, at least three independent biopsies corresponding to each condition were analyzed in each case. Statistical analyses were performed using unpaired Student's *t*-test. **p* < .05, ***p* < .01, ****p* < .005, ***p* < .001

11 of 24

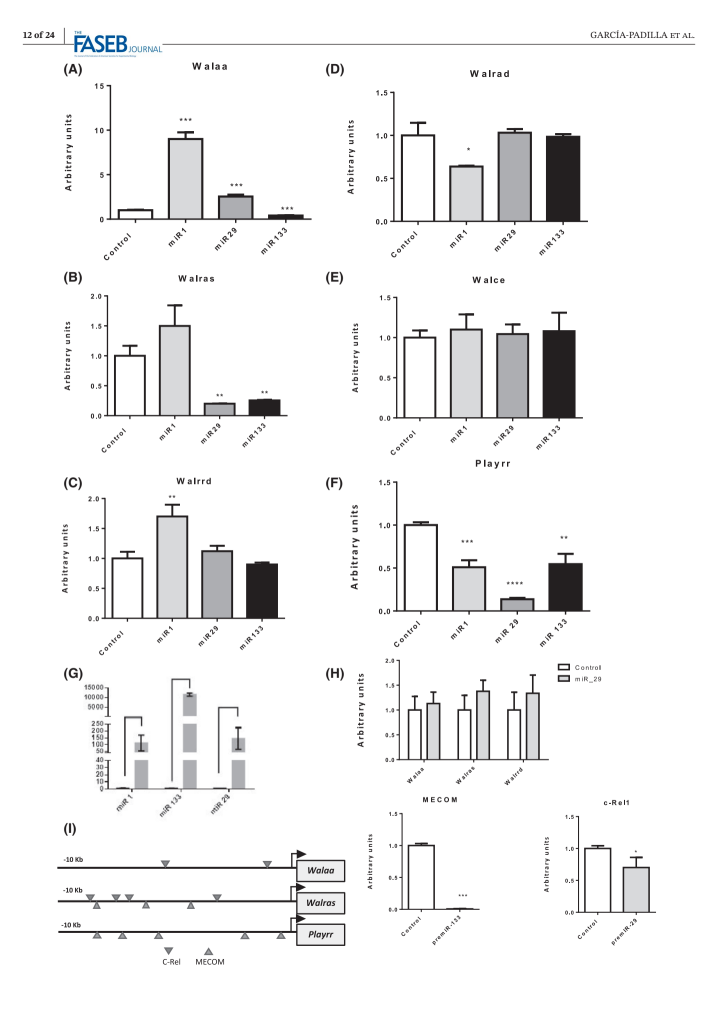


FIGURE 5 RT-qPCR analysis of the abundance of lncRNAs after elevating miR-1, miR-29 and miR-133 in HL1 atrial cardiomyocytes. (A–F) RT-qPCR analysis of *Walaa*, *Walras*, *Wallrd*, *Walca*, and *Playrr* in cells overexpressing miR-1, miR-29 and miR-133, respectively. miR-1 increased *Walaa* and *Wallrd* but decreased *Walrad* and *Playrr*. miR-29 overexpression elevated *Walaa* and reduced *Walras* and *Playrr*. miR-133 significantly suppressed *Walaa*, *Walras* and *Playrr*. (G) Levels of miR-1, miR-29, and miR-133 after ectopic overexpression. (H) represents miR29 pull-down assays, which did not influence *Walaa*, *Walras* and *Wallrd* interactions. (I) represents MECOM and C-Rel1 predicted binding sites on the upstream regulatory regions (-10 Kb) of *Walaa*, *Walras* and *Playrr*, respectively and their RT-qPCR expression after miR-133 and miR-29 overexpression. Three independent microRNA overexpression assays were analyzed in each case (A–E). Three independent pull-down assays were analyzed (F). Statistical analyses were performed using unpaired Student's *t*-test. **p < .01, ***p < .005, ***p < .001

of *Walrrd* in both experimental and human AF conditions, while *Walras* display downregulation in human AF but not in mice.

3.4 | AF related microRNAs modulate IncRNA expression

We have previously shown that signaling through Pitx2>Wnt regulates the expression of a large subset of microRNAs with pivotal roles regulating ion channel expression/function and thus has an important impact on the onset of atrial arrhythmogenesis.⁷⁻⁹ We therefore tested if AF-related microRNAs such as miR-1, miR-133 and miR-29 are capable of modulating the expression of these lncRNAs. It is important to highlight that microRNA-lncRNA interaction algorithms such as DIANA software (http://carolina.imis.athen a-innovation.gr/diana_tools/web/index.php?r=lncba sev2%2Findexpredicted&miRNAs%5B%5D=&lncRN As%5B%5D=&threshold=0.7&filters=0) failed to predict any putative binding of these microRNAs in these lncRNAs. We overexpressed these microRNAs in HL1 atrial cardiomyocytes and measured lncRNA expression levels by RT-qPCR analysis (Figure 5G). We found that ectopic expression of miR-1 and miR-29 increased the levels of Walaa, while miR-133 treatment decreased it (Figure 5A). Walrad was only downregulated by miR-1 while miR-29 and miR-133 resulted in no significant differences (Figure 5B). miR-133 and miR-29 administration reduced Walras levels in HL1 cardiomyocytes, while miR-1 had no effect on Walras abundance (Figure 5C). Walce was not altered by overexpression of any of the tested microRNAs (Figure 5D). Finally, miR-1 upregulates Wallrd expression while miR-133 and miR-29 administration leads to no significant changes (Figure 5E). In addition, we also found that *Playrr* was significantly downregulated by miR-1, miR-133 and miR-29 overexpression (Figure 5F). Pulldown analysis using biotinylated miR-29 revealed that the impact of miR-29 upon Walaa and Walras levels was indirect (Figure 5H). A plausible indirect mechanism might be that microR-NAs distinct regulate upstream regulatory transcription factors involved in lncRNA regulation. In this context,

we demonstrate that miR-133 and miR-29 significantly impaired the expression levels of MECOM and c-Rel, respectively. MECOM and c-Rel transcriptional binding sites are broadly identified within the proximal *Walras*, *Walras* and *Playrr* regulatory elements (10 Kb upstream the transcriptional start site) (Figure 5I). In sum, these data indicate that distinct microRNAs exert different regulatory properties upon these lncRNAs and further support a plausible functional role of these lncRNAs in the Pitx2>Wnt>microRNA pathways, previously reported to play a fundamental role in the onset of AF.

3.5 | Thyroid hormone administration distinctly regulated AF-associated lncRNAs

Impaired thyroid levels have been consistently reported as a cardiovascular risk factor associated with increased prevalence of atrial arrhythmias^{40,41} and hypertrophy.⁴²⁻⁴⁴ Therefore, we tested whether T3 and T4 administration in primary cultures of fetal cardiomyocytes might modulate the levels of these lncRNAs. Our data show that *Walaa* abundance is reduced following administration of T4 but not T3 (Figure 6A). On the other hand, administration of both T3 and T4 significantly increased the expression of *Walras* and *Walce* while it decreased the expression of *Wallrd* (Figure 6A). No significant differences were observed for *Walrad*, while *Playrr* was only downregulated by T4 administration (Figure 6A). Thus, thyroid administration modulates AF-associated lncRNAs, supporting the notion that these lncRNAs are involved in AF pathophysiology.

3.6 | Angiotensin II (AngII) and norepinephrine (NE) treatment significantly upregulated AF-associated lncRNAs

We previously reported that AF-associated cardiovascular risk factors, such as hyperthyroidism and hypertension, distinctly modulate signaling through Pitx2>Wnt>microRNA. Accordingly, we tested if these newly identified lncRNAs are regulated by AngII and/or NE administration. Primary cultures of fetal cardiomyocytes were treated with AngII

13 of 24

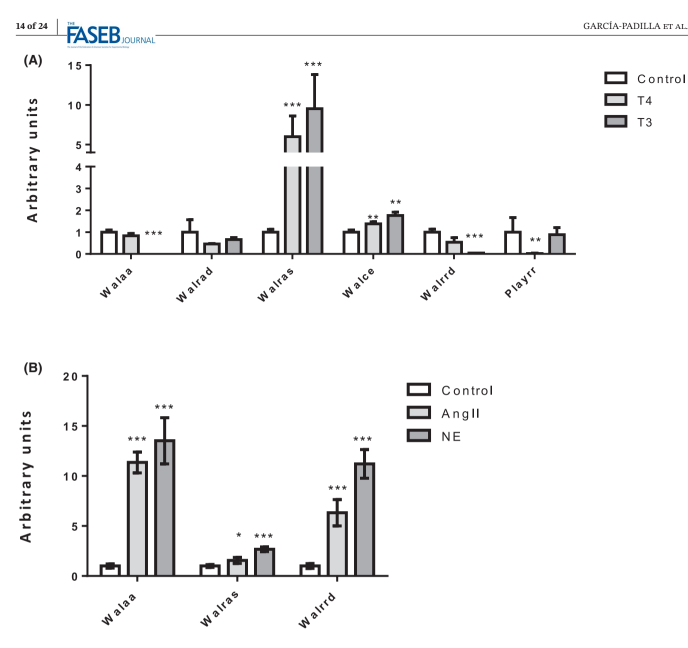


FIGURE 6 (A) RT-qPCR analyses of the levels of *Walaa*, *Walrad*, *Walras*, *Walce*, *Wallrd*, and *Playrr* after T3 or T4 thyroid hormone administration in HL1 atrial cardiomycoytes, respectively. T3 administration significantly suppressed *Walaa* and *Wallrd* levels and increased *Walras* and *Walce* levels while T4 administration increased *Walras* and *Walce* but downregulated *Playrr* while no changes are observed for *Walaa*, *Walrad* and *Wallrd*. (B) RT-qPCR analyses of the levels of *Walaa*, *Walras*, and *Wallrd* after angiotensin II (AngII) or norepinephrine (NE) administration. Both, AngII and NE treatments enhances *Walaa*, *Walras* and *Wallrd* expression. Three independent assays were analyzed in each case. Statistical analyses were performed using unpaired Student's *t*-test. **p* < .05, ****p* < .001

and NE, respectively, as previously reported,⁸ and lncRNA expression was assessed by RT-qPCR analysis. Our data indicate that AngII and NE administration, respectively, significantly enhanced the levels of *Walaa*, *Wallrd* and *Walras* (Figure 6B). In line with our previous findings after thyroid hormone administration, these data therefore support a plausible role of these lncRNAs in AF pathophysiology.

3.7 | Walras lncRNA pull down assays

To examine more deeply the function of the atriumspecific Walras lncRNA, we sought to identify the proteins interacting with it. To this end, we prepared lysates from HL1 cells and biotinylated *Walras*. After incubation, lncRNA-protein complexes were pulled down and the associated proteins were identified by LC-MS/ MS. A total of 40 proteins significantly interacting with *Walras* were identified (Table 2). Among them, ~40% were cytoplasmic proteins, 12% were nuclear and 20% mitochondrial, supporting a function dual role for *Walras* in the nucleus and cytoplasm, in line with our nuclear/ cytoplasmic distribution analyses (Figure 3C). Among the cytoplasmic proteins, *Walras* associated with three distinct myosin proteins, MYH6, MYH9 and MYH10; interestingly, MYH6 is a structural protein that forms part of

																							ASE			
Gene ontology (roles)	Actin cytoskeleton organization	Actin cytoskeleton reorganization	Actin filament-based movement	Angiogenesis	Glycolytic process	Actin filament bundle assembly	Aromatic compound catabolic process	ADP biosynthetic process	mRNA processing	Bone resorption	mRNA processing	Autophagosome assembly	Gluconeogenesis	Cholesteriol biosynthetic process	Actin crosslink formation	Glycolytic process	2-oxoglutarate metabolic process	Cellular response to DNA damage stimulus	Axonal fascilutaion	Peripheral nervous system axon regeneration	Electron transport chain	Axon extension	Alternative mRNA splicing, via spliceosome	Binding of sperm to zona pellucida	Activation of mitophagy in response to mitochondrial depolarization	Glutamate biosynthetic process
Subcellular location	Cytoplasm, cell projection, lamellipodium	Cytoplasm, cytoskeleton	Cytoplasm, myofibril. Note=Thick filaments of the myofibrils	Cytoplasm. Secreted	Cytoplasm	Cytoplasm	Microsome membrane; Single-pass type II membrane protein. ER membrane	Mitochondrion intermembrane space	Nucleus, nucleoplasm. Cytoplasm	Cytoplasm vesicle, phagosome membrane	Cytoplasm	Lysosome. Melanosome	Mitochondrion matrix	Isoform 1: ER membrane	Cytoplasm, cytoskeleton	Mitochondrion matrix	Cytoplasm	Cytoplasm	ER membrane	ER lumen. Melanosome	Mitochondrion inner membrane	Cell membrane	Nucleus. Cytoplasm. Shuttles continuously between the nucleus and the cytoplasm	Cytoplasm. Melanosome	Cytoplasm. Cell membrane. Cell projection, lamellipodium	Mitochondrion matrix
Short-name	Myosin-10	Myosin-9	Myosin-6	Glucose-6-phosphate isomerase	Phosphoglycerate kinase 1	Alpha-actinin-4	Epoxide hydrolase 1	Adenylate kinase 2, mitochondrial	Heterogeneous nuclear ribonucleoproteins A2/B1	Ras-related protein Rab-7a	Heterogeneous nuclear ribonucleoprotein K	Cathepsin D	Triosephosphate isomerase	NADH-cytochrome b5 reductase	Alpha-actinin-1	Phosphoglycerate mutase 1	Aspartate aminotransferase, cytoplasmic	Ubiquitin-like modifier-activating enzyme 1	Reticulon-4	Calumenin	Electron transfer flavoprotein-ubiquinone oxidoreductase, mitochondrial	Integrin beta-1	Heterogeneous nuclear ribonucleoprotein B	T-complex protein 1 subunit delta	Nucleoside diphosphate kinase B	Glutamate dehydrogenase 1, mitochondrial
Gene	MYH10	6НХМ	MYH6	G6PI	PGK1	ACTN4	НҮЕР	KAD2	ROA2	RAB7A	HNRPK	CATD	TPIS	NB5R3	ACTN1	PGAM1	AATC	UBA1	RTN4	CALU	ETFD	ITB1	ROA1	TCPD	NDKB	DHE3
UniprotKB-ID	Q61879	Q8VDD5	Q02566	P06745	P09411	P57780	Q9D379	Q9WTP6	O88569	P51150	P61979	P18242	P17751	Q9DCN2	Q7TPR4	Q9DBJ1	P05201	Q02053	Q99P72	035887	Q921G7	P09055	P49312	P80315	Q01768	P26443
Accessiona	71152969	205371802	3024204	146345422	146345481	13123946	341940814	224471907	124028629	46397834	48429104	115718	353526354	60390645	46395721	20178035	338817898	267190	94730421	5915871	52000730	124964	1350822	549057	266608	118542

15 of 24

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	(CONTINUACIO)					В
Accessiona	UniprotKB-ID	Gene	Short-name	Subcellular location	Gene ontology (roles)	JOURN
81896595	Q8BLF1	NCEH1	Neutral cholesterol ester hydrolase 1	Membrane	Lipid catabolic process	AL
29839593	Q91W90	TXND5	Thioredoxin domain-containing protein 5	ER lumen	Apoptotic cell clearance	
118105	P17742	PPIA	Peptidyl-prolyl cis-trans isomerase A	Cytoplasm. Secreted	Lipid particle organization	
61217662	Q9Z1N5	DX39B	Spliceosome RNA helicase Ddx39b	Nucleus	Cellular response to DNA damage stimulus	
50400807	035114	SCRB2	Lysosome membrane protein 2	Lysosome membrane	Cell adhesion	
6093768	Q60930	VDAC2	Voltage-dependent anion-selective channel protein 2	Mitochondrion outer membrane	Negative regulation of intrinsic apoptotic signalling pathway	
46577116	Q9D1G1	RAB1B	Ras-related protein Rab-1B	Cytoplasm	Autophagy	
71153826	Q7TPV4	MBB1A	Myb-binding protein 1A	Nucleus	Cellular response to glucose starvation	
1709998	P53994	RAB2A	Ras-related protein Rab-2A	ER-Golgi intermediate compartment membrane; Lipid-anchor. Melanosome	Golgi organization	
342187049	P26039	TLN1	Talin-1	Cell projection, rufile membrane	Cell-substrate junction assembly	
341940396	P52825	CPT2	Carnitine O-palmitoyltransferase 2, mitochondrial	Mitochondrion inner membrane	Fatty acid metabolic process	
66773808	Q9JHI5	ΠΛΙ	Isovaleryl-CoA dehydrogenase, mitochondrial	Mitochondrion matrix	Fatty acid beta-oxidation using acyl- CoA dehydrogenase	
51702234	P62897	CYC	Cytochrome c, somatic	Mitochondrion intermembrane space. Loosely associated with the inner	Apoptotic process	
51701449	Q9CQA3	SDHB	Succinate dehydrogenase [ubiquinone], iron- sulfur subunit, mitochondrial	Mitochondrion inner membrane	Respiratory electron transport chain	
<i>Note</i> : Observed that 40 inter and endoplasmic reticulum.	hat 40 interacting prot 5 reticulum.	teins were ide	entified, most of them are located in the cytoplasm, with a sr	Note: Observed that 40 interacting proteins were identified, most of them are located in the cytoplasm, with a small but significant fraction is also observed in other cellular organelles such as the nucleus, mitochondria and endoplasmic reticulum.	rganelles such as the nucleus, mitochondria	

TABLE 2 (Continued)

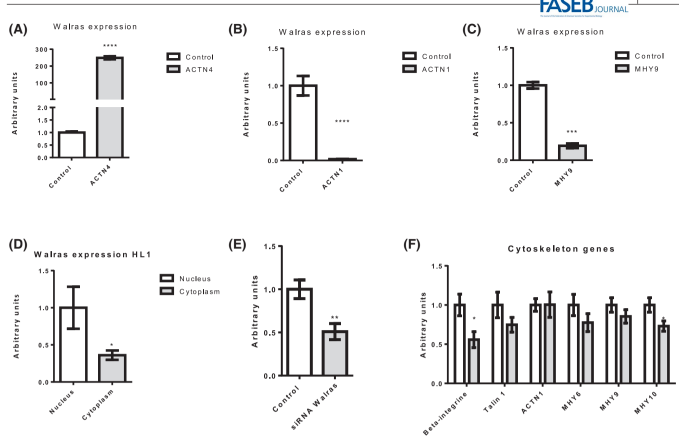


FIGURE 7 (A–C) RIP assays for ACTN4 (A), ACTN1 (B) and MHY9 (C), respectively. (D) RT-qPCR analyses of Walras nuclear and cytoplasmic distribution in HL1 atrial cardiomyoctes. (E) *Walras* levels 24 h after siRNA transfection. (F) RT-qPCR analyses of genes encoding for Walras interacting proteins in cells transfected with *Walras* siRNA (grey bars) as compared to controls (white bars)

the sarcomere and is frequently altered in cardiac pathological conditions such as cardiac hypertrophy^{45–47} as well as more recently with AF.⁴⁸ Several cytoskeletal proteins RAB1A, RAB2A, RAB7A, ACTN1, ACTN4, TLN1 and ITB1 are also associated to Walras. Importantly several of these proteins are linked to WNT non-canonical signaling pathway and at least one of them (ACTN4) has been recently linked to AF.49 Among the nuclear proteins, Walras was found to interact with ROA2, ROA1, DXC39B and MBBIA, suggesting that it can also play a nuclear role, possibly in transcription, although at present the functions of these interacting proteins are poorly understood. Surprisingly, Walras also interacted with a large array of mitochondrial proteins, such as KAD2, ETFD, DHE3, VDAC2, IVD, CYC and SDHB, as well as with proteins located in other subcellular organelles, particularly at the ER (RTN-4, CALU, TXDN5); the functional consequences of these interactions remain to be explored. In sum, these data revealed the complexity of Walras-interacting proteins and pave the ways for dissecting the functional role of this lncRNA in a cardiovascular context. Validation of the Walras pulldown assays were performed for a subset of these interacting proteins (Figure 7A-C). We demonstrated by RIP assays in HL1 cardiomyocytes that ACTN4,

but not ACTN1 and MHY9, directly interact with *Walras* (Figure 7A–C). Importantly, *Walras* is distributed in both, nuclear and cytoplasmic compartments in HL1 cardiomyocytes (Figure 7D), in line with our observations in embryonic and neonatal cardiac chambers (Figure 3C). Further support of *Walras* lncRNA-protein interactions is provided as qPCR analyses of distinct *Walras* interacting proteins failed to display differential expression in siRNA *Walras* treated cells at mRNA level, except for integrin beta 1 (*Itb1*).

To further support the plausible functional role of *Walras* in cardiomyocytes, we designed siRNAs and performed loss-of-function assays in HL1 cardiomyocytes. RT-qPCR analyses demonstrated significant knockdown of *Walras* in HL1 cardiomyocytes (Figure 7E). Immunohistochemical analyses in HL1 cardiomyocytes demonstrated that cells treated with *Walras* siRNAs display significantly decreased cell-cell interactions (Figure 8B,C) as compared to controls (Figure 8A,C) and cytoskeletal rearrangements (Figure 8D–C). Immunocytochemical assays of ACTN4 (Figure 8D–F), ACTN1 (Figure 8G–I) and MHY9 (Figure 8J–L) distribution evidenced a significant downregulation of these cytoskeletal proteins in *Walras* siRNA treated HL1 cardiomyocytes.

17 of 24

FASEB JOURNAL

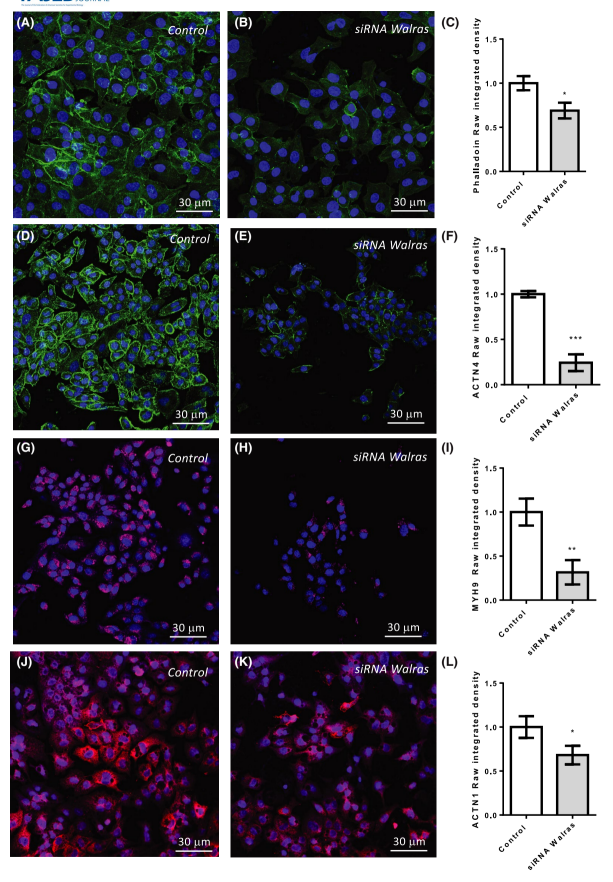


FIGURE 8 Immunohistochemical analyses of control (A,D,G,J) and *Walras* silenced (B,D,H,K) cardiomyocytes stained with phalloidin (A,B), anti-ACTN4 (D,E), anti-ACTN1 (G–H) and MHY9 (J–K). Note that cell-cell interactions are disrupted in all *Walras* siRNA conditions, and that phalloidin (C), ACTN4 (F), ACTN1 (I) and MHY9 (L) stainings are significantly decreased. DAPI staining (blue) identify cell nuclei

4 | DISCUSSION

AF is the most common cardiac arrhythmia in humans, with an estimated incidence of 2%–4% in the general population, but rising up to 8%–10% in the elderly. Mechanistically, AF is initiated by impaired electrical activity that is most frequently originated in the left atrium, particularly in the vicinity of the entrance of pulmonary veins. The onset of AF leads to structural and electrical remodeling of both the left and right atrial chambers leading to self-perpetuation.⁵⁰

Over the past few decades, our understanding of the genetic basis of AF has increased greatly. A large number of studies provided evidence on the causal relationship between distinct point mutations in multiple genes encoding ion channels and the occurrence of AF. More recently, a seminal paper by Gudbjartsson et al.¹ identified risk variants at 4q25 highly associated to lone AF and subsequently corroborated in distinct AF cohort studies.^{51–55} These variants are located 150 kb downstream of the homeobox transcription factor PITX2. Importantly, lossof-function assays in mice showed that impaired Pitx2 function leads to increased atrial arrhythmia susceptibility.⁵⁻⁷ Wang et al.⁵ demonstrated aberrant embryonic *Shox2* and Tbx3 expression in Pitx2 loss-of-function and related such impaired expression to AF susceptibility. Kirchhof et al.⁶ reported similar findings and demonstrated that the expression of multiple ion channels was impaired. Importantly, in both cases, no AF episodes occurred under basal conditions. Using an atrial-specific Pitx2 conditional transgenic mouse line, Chinchilla et al.⁷ demonstrated abnormal ECGs at rest as well as impaired SCN5A (I_{Na}), KCNJ2 (I_{K1}) and KCNJ12 (I_{K1}) expression and function. These observations were subsequently corroborated by Tao et al.⁵⁶ using a model of conditional Pitx2 expression in adult mice. Additional evidence was found for a role for Pitx2 in calcium homeostasis.^{8,57-60} These data therefore suggest that an embryonic impairment of Pitx2 might predispose to atrial arrhythmias while an adult deficiency will be already causative of AF.

4.1 | Expression and distribution of lncRNAs

In this study, we have identified four distinct Wntassociated lncRNAs with preferential expression in the atrial chambers (*Walaa*, *Walras*, *Wallrd* and *Walrad*), three of them confined to the embryonic stages (*Walaa*, *Walras* and *Wallrd*), and a fourth lncRNA that is also preferentially expressed in the adult right atrium (*Walrad*). Expression of these lncRNAs is widely distributed along several other tissues such as the liver, brain, lungs and gastrointestinal tract. Curiously they display discordant expression profiles as compared to Wnt8 and Wnt11, during development and adulthood, suggesting a lack of regulatory influence. Importantly, three of them (Walras, Wallrd and Walce) are conserved in humans. Walras display variable degree of homology with three distinct IncRNAs adjacent to the human WNT11 locus, being the most likely homologous one AP002340.1 lncRNAs since it displays 43% similarity and is roughly similar in size (238 nt vs 285 nt), while AP000785.1 and LINC02761 display lower similarity (25% and 31%, respectively) and are lager in size (613 nt and 388 nt, respectively). Mouse Walrrd presents three distinct isoform, each of them displaying homologue similarity to AC13382.1 (33%), AC13382.2 (45%) and AC1041116.1 (44%), while Walce displayed 35% similarity to Wnt11 03 lncRNAs. Such findings are in line with previous results of the modest similarity of lncRNAs between distinct species.

LncRNAs are found in the nucleus and cytoplasm, primarily functioning as transcriptional or posttranscriptional modulators,⁶¹⁻⁶⁴ respectively. Our analyses demonstrate that most lncRNAs studied display dual nuclear and cytoplasmic localization i.e., Walras, Walce, Wallrd and Playrr. Importantly, both temporal and spatial differences in the nuclear vs cytoplasmic localization was observed for several of these lncRNAs, with Walrad being primarily nuclear in the embryonic and neonatal right and left atrial chambers but not in the ventricles, and Walas displaying a prominent nuclear localization in the embryonic right atrium but similarly distributed in both atrial chambers at the neonatal stage. These data underscore the highly dynamic temporal and spatial distribution of these lncRNAs during cardiogenesis and support the notion that they can exert distinct transcriptional and post-transcriptional functions.

4.2 | Modulation of lncRNAs

Subsequently experiments revealed that Pitx2 enhanced the expression of *Walaa* and *Walce*, while it significantly decreased the expression of *Wallrd*, supporting a role for these lncRNAs in a Pitx2-regulated processes. Analyses in an atrium-specific Pitx2 conditional transgenic mice further supported a regulatory role of Pitx2 in raising the levels of *Walaa* and *Wallrd*. Importantly, such upregulation is also observed for human *WALLRD* (AC1040116.1) in AF patients. Moreover, *WALRAS* (AP002340.1) is downregulated in AF patients, further supporting a plausible role for these lncRNAs in Pitx2-induced AF. Surprisingly, divergent results are observed for *Wallrd* in atrium-specific Pitx2 conditional transgenic mice. It is possible that additional regulatory

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mechanisms are driven by Pitx2 in non-cardiomyocyte cells. Additional experiments will be required to reconcile these findings.

More recently, GWAS have identified additional genes linked to AF pathophysiology, such as KCNN3, ZFHX3, IL6R, CAV1, HCN4, SYNE2, SYNPOL2, PRRX1, and *WTN8A*, among others.^{1,65–68} To date more than 90 genes have been associated to AF by GWAS analyses.² Strategies to increase and reduce Pitx2, Lozano-Velasco et al.8 demonstrated that Pitx2 can drive expression of Wnt8a, but not Zfhx3. Interestingly, impaired Wnt8a expression is only reported in Pitx2 loss-of-function models with abnormal electrophysiological parameters at rest, supporting the notion that Wnt signaling is essential for the AF susceptibility vs triggering capacity. In this context it is important to highlight that risk variants associated with increased PR interval, a condition frequently associated to AF, have been identified at the WNT11 locus, neighboring WALRAS conserved human homologues (AP002340.1, AP000785.1 and LINC02761 lncRNAs).

In this study we therefore analyzed if *Wnt8* and *Wnt11* could modulate lncRNA expression as they exert synergistic and complementary regulatory mechanisms in the Pitx2>Wnt>microRNA signaling pathway as previously reported by Lozano-Velasco et al.^{8,9} Our data demonstrate that Wnt8 *vs* Wnt11 overexpression modulate *Walrad*, *Walras* and *Walce* expression in a complementary manner. Therefore, expression of *Walrad* and *Walce* is regulated by both Pitx2 and Wnt whereas *Walras* is exclusively regulated by Wnt signaling. Thus, these data reinforce the notion of a plausible role for these lncRNAs in Pitx2>Wnt signaling and thus in atrial arrhythmias.

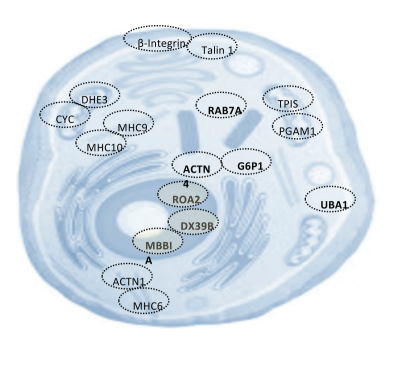
Different studies have reported that Pitx2-regulated microRNAs are involved in AF pathophysiology. Chinchilla et al.⁷ demonstrated abnormal regulation of *Kcnj2* and Kcnj12 by miR-1, leading to impaired resting membrane potential configuration in Pitx2 loss-of-function mice. Lozano-Velasco et al.8 identified several microRNAs regulated by Pitx2 that were previously linked to AF in humans. In this context, miR-1, miR-133, miR-21, miR-26, miR-29 and miR-106b have previously been associated with the regulation of calcium (CACNA1C, RYR2), sodium (SCN5A), potassium (KCNJ2, KCNE1, KCNB2), cation (HCN4) channel subunits, respectively (revised in Ref. [11]). Importantly, microRNA regulation by Pitx2 is exerted by a balanced interplay between Wnt8 and Wnt11 expression.8 Therefore, in the present study we tested if overexpression of Pitx2-regulated microRNAs influenced the expression of lncRNAs. Our data indicate that that microRNA overexpression can indeed modulate lncRNAs. In particular, miR-1 overexpression upregulated Walaa and Wallrd, but not Walras. In addition, miR-29 overexpression also significantly increased Walaa, while miR-133 significantly decreased *Walaa* and *Walras*. These data reveal that the expression of *Walaa* and *Wallrd* is modulated by Pitx2 and Wnt signalling, as well as by key pro-arrhythmogenic microRNAs. Importantly, such modulatory actions seems to be indirect, as several transcription factors potentially involved in *Walaa*, *Walras* and *Playrr* regulation are distinctly impaired by miR-133 and miR-29 overexpression. Together, these data support a plausible role for these novel lncRNAs in atrial arrhythmias.

There is wide demonstration that hypertension (HTN), hyperthyroidism (HTD), diabetes and obesity promote AF onset, respectively.⁶⁹⁻⁷¹ In addition, AF occurrence can be also increased in presence of hypertrophic cardiomyopathy and valvular heart diseases.^{70,72,73} Importantly, Pitx2 is upregulated in cardiac hypertophy⁷⁴ as well as in heart failure.¹⁰ We have recently provided evidence on the modulatory role of upstream pathways modulating Pitx2 in the setting of AF. HTD but not HTN leads to impaired Pitx2>Wnt>microRNA signaling causing thus abnormal ion channel expression. We herein tested whether AngII and NE treatment modulate lncRNA expression. Our data demonstrate that all lncRNAs analyzed are upregulated upon AngII and NE administration, supporting a plausible role of these lncRNAs in atrial arrhythmias as well as in cardiac hypertrophy.

In addition, we also analyzed if thyroid hormone influenced their expression. Surprisingly, thyroid hormone distinctly modulated lncRNA expression as compared to AngII/NE administration. Thus, *Walaa* was significantly downregulated after T3 administration as well as *Wallrd* after both T3 and T4 administration, respectively. On the other hand, *Walras* was significantly upregulated after both T3 and T4 administration, respectively. These data further strengthen a role for these lncRNAs in atrial arrhythmias as they display differential modulation by AngII/NE vs T3/T4 administration.

4.3 | Functional roles of lncRNAs

In order to start understanding the functional roles of these lncRNAs in cardiac development, pulldown assays for the lncRNA *Walras* were performed in HL1 atrial cardiomyocytes. A total of 40 proteins were identified that significantly interacted with *Walras*. Most of these proteins are cytoplasmic (>40%), suggesting a prominent extranuclear role for *Walras* (Figure 8). Two major sets of cytoplasmic proteins were found, myosin proteins (MHY6, MHY9 and MHY10) and cytoskeletal proteins (RAB1A, RAB2A, RAB7A, ACTN1, ACTN4 and TLN1). Importantly, we validated a direct protein-*Walras* interaction for ACTN4. Point mutations in MHY6 have been associated to cardiac hypertrophy, dilated cardiomyopathy, AF and congenital



Cytoskeleton and membrane cell proteins

21 of 24

- MHC10
- MHC9
- MHC6
- β-Integrine
- Talin 1
- ACTN1

Cytoplasm related proteins

- G6P1
- ACTN4
- RAB7A
- UBA1

Nuclear related proteins

- ROA2
- DX39B
- MBBIA

Mitochondria proteins

- TPIS
- PGAM1
- DHE3
- CYC

FIGURE 9 Schematic representation of the most prominent *Walras*-interacting proteins as identified by pulldown assays and mass spectrometry identification. Observe that *Walras* interacts with distinct cytoskeletal and cytoplasmic proteins as well as nuclear proteins mitochondrial proteins

heart diseases.^{75–79} In addition, SNV in the proximity of MHY6 have been associated with AF and impaired heart rate.⁴⁵ Experiments in mice further support the functional role of MHY6 in cardiac pathophysiology.⁸⁰ RAB7A has been implicated in controlling β-adrenergic receptor endosome recycling⁸¹ and impaired β-adrenergic stimulation is frequently associated to AF. Abnormal expression of ACTN1 and TLN1 leads to dilated cardiomyopathy.⁸²⁻⁸⁴ In particular, TLN1 knock-out display decreased beta1integrin expression (another Walras interacting protein), costameric instability and cardiac hypertrophy.⁸⁴ Thus, it is plausible that the interaction of Walras with TLN1 might contribute to stabilization of cytoskeletal proteins such as ACTN1, ACTN4 and MHY9 and MHY10, via ITG1 phosphorylation,^{83,84} a process that might be impaired in cardiac hypertrophy. In this context, we provided evidence that impairing Walras expression leads to decreased expression of ACTN1, ACTN4 and MHY9 in HL1 atrial cardiomyocytes. Importantly, two additional cytoplasmic Walras-interacting proteins are also impaired in cardiac hypertrophy, UBA1 and PPIA.^{85–87} Collectively, these lines of evidence support the idea that impairment of Walras might also lead to cardiac arrhythmias and/or hypertrophy by deregulating several interacting proteins (Figure 9).

Walras also showed affinity for nuclear proteins ROA2, ROA1, DXC39B and MBBIA, supporting a nuclear function for Walras. It is important to note that we found similar subcellular distribution patterns for Walras in other tissues analyzed, except in the embryonic right atrium and ventricles. At present, the functional roles of these interacting proteins are poorly understood. Surprisingly, Walras also interacts with a large array of mitochondrial proteins (KAD2, PGAM1, DHE3, CPT2, IVD, CYC, and SDHB) and endoplasmic reticulum proteins (HYEP, NB5R3, RTN-4, CALU, ETFD, and TXDN5). While the nature and functional consequences of such interactions remain to be explored, it is important to highlight the fact that impaired expression of SDHD, CPT2, RTN4, ETFD, and CATD has also been associated to hypertrophic and/ or dilated cardiomyopathy,^{88–91} reinforcing the notion that Walras might play a role in cardiac hypertrophy.

In summary, we have identified five novel lncRNAs that are differentially expressed in the developing and adult heart. Three of them displayed atrial expression during embryogenesis and a fourth one is expressed in the right atrium in adulthood. Signaling through Pitx2>Wnt>miRNAs modulates the expression of these lncRNAs. In addition, pro-hypertrophic and pro-arrhythmogenic pathways such FASEBJOURNAL

as those induced by AngII/NE and thyroid hormone administration distinctly regulated the expression of these lncRNAs. Pulldown and loss-of-function assays revealed that *Walras* interacts primarily with cytoplasmic proteins that, if impaired, are associated with cardiac arrhythmias and hypertrophy. Overall, these data suggest that role of these lncRNAs might plays a role in cardiac arrhythmogenesis and hypertrophy.

DISCLOSURES

The authors declare no conflicts of interest.

AUTHOR CONTRIBUTIONS

Diego Franco, Amelia E. Aránega, Myriam Gorospe, Leif Hove-Madsen designed research; Carlos García-Padilla, Jorge N. Domínguez, Valeria Lodde, Rachel Munk, Kotb Abdelmohsen performed research; Carlos García-Padilla, Jorge N. Domínguez, Valeria Lodde, Rachel Munk, Kotb Abdelmohsen analyzed data; Veronica Jiménez-Sábado, Antonino Ginel contributed with clinical data; Diego Franco wrote the draft paper; Carlos García-Padilla, Myriam Gorospe, Leif Hove-Madsen and Diego Franco edited final manuscript.

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