Endoplasmic Reticulum Stress Is Associated With Autophagy and Cardiomyocyte Remodeling in Experimental and Human Atrial Fibrillation

Marit Wiersma, PhD,* Roelien A. M. Meijering, PhD,* Xiao-Yan Qi, PhD; Deli Zhang, PhD; Tao Liu, MD; Femke Hoogstra-Berends, BSc; Ody C. M. Sibon, PhD; Robert H. Henning, MD, PhD; Stanley Nattel, MD; Bianca J. J. M. Brundel, PhD

Background—Derailment of proteostasis, the homeostasis of production, function, and breakdown of proteins, contributes importantly to the self-perpetuating nature of atrial fibrillation (AF), the most common heart rhythm disorder in humans. Autophagy plays an important role in proteostasis by degrading aberrant proteins and organelles. Herein, we investigated the role of autophagy and its activation pathway in experimental and clinical AF.

Methods and Results—Tachypacing of HL-1 atrial cardiomyocytes causes a gradual and significant activation of autophagy, as evidenced by enhanced LC3B-II expression, autophagic flux and autophagosome formation, and degradation of p62, resulting in reduction of Ca2+ amplitude. Autophagy is activated downstream of endoplasmic reticulum (ER) stress: blocking ER stress by the chemical chaperone 4-phenyl butyrate, overexpression of the ER chaperone-protein heat shock protein A5, or overexpression of a phosphorylation-blocked mutant of eukaryotic initiation factor 2α (eIF2α) prevents autophagy activation and Ca2+-transient loss in tachypaced HL-1 cardiomyocytes. Moreover, pharmacological inhibition of ER stress in tachypaced Drosophila confirms its role in derailing cardiomyocyte function. In vivo treatment with sodium salt of phenyl butyrate protected atrial-tachypaced dog cardiomyocytes from electrical remodeling (action potential duration shortening, L-type Ca2+-current reduction), cellular Ca2+-handling/contractile dysfunction, and ER stress and autophagy; it also attenuated AF progression. Finally, atrial tissue from patients with persistent AF reveals activation of autophagy and induction of ER stress, which correlates with markers of cardiomyocyte damage.

Conclusions—These results identify ER stress–associated autophagy as an important pathway in AF progression and demonstrate the potential therapeutic action of the ER-stress inhibitor 4-phenyl butyrate. (J Am Heart Assoc. 2017;6:e006458. DOI: 10.1161/JAHA.117.006458.)

Key Words: 4PBA • atrial fibrillation • autophagy • Drosophila • drug research • Endoplasmic Reticulum stress • HSPA5 • molecular biology • structural biology • tachypacing

Atrial fibrillation (AF) is the most common persistent clinical tachyarrhythmia.1 Many patients experience clinical symptoms, including palpitations, fatigue, and weakness; AF also puts patients at risk for cardiac morbidity and mortality and often necessitates life-long anticoagulant therapy.1 When AF persists, sinus rhythm (SR) reversion and maintenance becomes progressively more difficult. Central to this self-perpetuating nature of AF is the remodeling of cardiomyocytes as a consequence of the increased atrial activation rate, resulting in disturbances of electrophysiological features and contraction and structural damage.2 Therapeutic strategies that limit cardiomyocyte remodeling would improve the success of...
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folding and preservation of contractile proteins.8,9 Proteins (HSPs), whose chaperone function subserves correct dysfunction as a consequence of induction of heat shock revealed by the attenuation of cardiomyocyte remodeling and countered disease progression in a dog model of AF.

Recent work shows that the mammalian target of rapamycin (mTOR) pathway12,13 and endoplasmic reticulum (ER) stress15 which plays a major role in the cardiac stress response.15 Macroautophagy (hereafter “autophagy”) is critically involved in maintaining proteostasis.10 Autophagy is an evolutionarily conserved protein-degradation pathway that removes damaged or expired proteins and organelles by sequestration in damaged or expired proteins and organelles by sequestration in microtubule network and cardiomyocyte structural remodeling.3,4,7 The importance of proper proteostasis is also depicted disease progression in both in vitro and in vivo AF models. Thus, our study points to ER stress as a potential novel druggable target to attenuate cardiac remodeling in AF.

Methods

HL-1 Atrial Cardiomyocyte Cell Culture, Transfections, and Constructs

HL-1 atrial cardiomyocytes derived from adult mouse atria were obtained from Dr William Claycomb (Louisiana State University, New Orleans).20 The cardiomyocytes were maintained in complete Claycomb Medium (Sigma) supplemented with 10% fetal bovine serum, 100 U/mL penicillin, 100 µg/mL streptomycin, 4 mmol/L L-glutamine, 0.3 mmol/L L-ascorbic acid, and 100 µmol/L norepinephrine. HL-1 cardiomyocytes were cultured on cell culture plastics or on glass coverslips coated with 0.02% gelatin in a humidified atmosphere of 5% CO2 at 37°C. Where indicated, HL-1 cardiomyocytes were transiently transfected with the LC3B–green fluorescent protein (kind gift of Professor T. Johansen),21 HSPA5 (kind gift of Professor H. Kampinga), pcDNA3.1+ (empty), eIF2α wild type, eIF2α S51A, or eIF2α S51D plasmid, by the use of Lipofectamine 2000.

Tachypacing of HL-1 Cardiomyocytes and Calcium Transient Measurements

HL-1 cardiomyocytes were subjected to tachypacing, as described before.3 In short, HL-1 cardiomyocytes were subjected to 1 Hz (normal pacing) or 6 Hz (tachypacing; Table 1), 40 V, and 20-millisecond pulses, for a maximal duration of 8 hours via the C-Pace EP Culture Stimulator. These frequencies were used to standardize the control firing frequency (1 Hz is the average spontaneous beating rate of HL-1 cardiomyocytes) and to produce a similar frequency increment with tachypacing (6-fold increase) to that which occurs during AF in humans. To measure Ca2+ transients (CaTs), HL-1 cardiomyocytes were incubated for 30 minutes with 2 µmol/L Ca2+-sensitive dye, Fluo-4-AM. Fluo-4–loaded cardiomyocytes were excited by a 488-nm laser with emission at 500 to 550 nm and were visually recorded with a 40× objective, using a Solamere-Nipkow-Confocal-Live-Cell-Imaging system (based on a DM IRE2 inverted microscope). The live recording of CaT in HL-1 cardiomyocytes was...
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performed at 1-Hz stimulation at 37°C. Live recordings were further processed by use of the software ImageJ. The relative value of fluorescence signals between experiments was determined using the following calibration: Fcal = F1/F0, where F1 is the fluorescent dye signal at any given time and F0 is the fluorescent signal at rest. Mean values and SEMs from each experimental condition were based on 7 consecutive CaTs in at least 50 cardiomyocytes.

Drug Treatment

Pepstatin A, bafilomycin A1 (BAF), tunicamycin, rapamycin, and 4PBA were dissolved, according to manufacturer’s instructions. HL-1 cardiomyocytes were treated with 4PBA (10 mmol/L), tunicamycin (5 μg/mL), and rapamycin (50 mmol/L) 8 hours before pacing. Pepstatin A (10 μmol/L) and BAF (10 mmol/L) were added 30 minutes before normal or tachypacing.

Drosophila Stocks, Tachypacing, and Heart Wall Contraction Assays

For all experiments, w 1118 strains were used. All flies were maintained at 25°C on standard medium. After fertilization, adult flies were removed and drugs were added to the medium containing fly embryos. Drosophila embryos and larvae were treated with 4PBA (100 mmol/L), pepstatin A (100 μmol/L), or BAF (100 mmol/L) during development. Controls were treated with the vehicle, 2% dimethyl sulfoxide. After 2 days, prepupae were selected for tachypacing, as previously described. 22 Groups of at least 5 prepupae were subjected to tachypacing (5 Hz for 20 minutes, 20-V and 5-millisecond pulses; Table 1) with a C-Pace EP Culture Stimulator. Before and after tachypacing, videos of spontaneous heart wall contractions in whole prepupae were recorded for 30 seconds. Heart wall contractions were analyzed with IonOptix software.

Western Blot Analysis

Western blot analysis was performed, as previously described. 3 Briefly, equal amounts of total protein in SDS-PAGE sample buffer were separated on SDS-PAGE 4% to 20% Precise Tris-HEPES gels. After transfer to nitrocellulose membranes, membranes were incubated with primary antibodies, followed by incubation with horseradish peroxidase–conjugated anti-mouse or anti-rabbit secondary antibodies. Signals were detected by the Western Lighting Ultra method and quantified by densitometry via the software Gene Gnome, Gene tools. The following antibodies were purchased: rabbit anti–phosphorylated protein kinase B (Akt; Ser473), rabbit anti-Akt, rabbit anti-LC3B, rabbit anti-SOSTM1/p62, rabbit anti–phosphorylated elf2α (Ser51), rabbit anti–phosphorylated S6 ribosomal protein (Ser235/236), mouse anti-S6 ribosomal protein, rabbit anti–phosphorylated mTOR (Ser2448/2481), rabbit anti-mTOR, mouse anti-elf2α, mouse anti-HSPA5, mouse anti–β-actin, and mouse anti-GAPDH; rabbit anti–β-myosin heavy chain 7 (MHC) was a kind gift of Professor J. Van der Velden.

Quantitative Real-Time Polymerase Chain Reaction

Total RNA was isolated from HL-1 cardiomyocytes using the nucleospin RNA isolation kit. First-strand cDNA was generated by M-MLV reverse transcriptase and random primers. Relative changes in transcription level were determined using the CFX384 Real-Time System C1000 Thermocycler in combination with SYBR green ROX-mix. Calculations were performed using the comparative threshold cycle method, according to User Bulletin 2. Fold inductions were adjusted for GAPDH levels.

Primer pairs used included the following: ATF4, GTCCGTTA-CAGCAACACTGC (forward) and CCACCATGCGTATTAGGG (reverse); ATF6, AAGAGAAGGCCTGTCACTG (forward) and GGCCTGTAGTGCTGAAT (reverse); CHOP, GACCGAGTTCTGC TTCAGG (forward) and CAGCGACAGCCAGAATAA (reverse); HSPA5, ATCTTGTTGTCTTGCTG (forward) and ATGAAGGAGACTGCTGAGGC (reverse); autophagy gene 12, TCTCCAGACCCATTCTTGG (forward) and AACTCCGGGAGAC ACCAAG (reverse); and GAPDH, CATCAAGAAGGTGGTAAGC (forward) and ACCACCCGTGCTGCTAG (reverse). Polymerase chain reaction efficiencies for all primer pairs were between 90% and 110%.

Immunofluorescent Staining and Confocal Analysis

HL-1 cardiomyocytes were untransfected or transiently transfected with green fluorescent protein–LC3B for 48 hours and paced at 1 Hz (normal pacing) or 6 Hz (tachypacing). This was followed by fixation with 4% formaldehyde for 15 minutes at room temperature and washing 3 times with PBS; then, they were permeabilized and blocked with 0.3% Triton X-100 and 5% fetal bovine serum in PBS (1 hour at room

<table>
<thead>
<tr>
<th>Model</th>
<th>NP Group, Hz (bpm)</th>
<th>TP Group, Hz (bpm)</th>
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<tbody>
<tr>
<td>HL-1</td>
<td>1 (60)</td>
<td>6 (360)</td>
</tr>
<tr>
<td>Drosophila</td>
<td>1.5 (90)</td>
<td>5 (300)</td>
</tr>
<tr>
<td>Dog</td>
<td>1.3 (80)</td>
<td>10 (600)*</td>
</tr>
</tbody>
</table>

Bpm indicates beats per minute; NP, normal-paced; and TP, tachypaced.

*Please note that 600 bpm atrial tachypacing in the dog induces atrial fibrillation (AF), with atrial-tissue responses at ≈6 to 8 Hz. This model is intentionally used to produce sustained AF, thereby mimicking the clinical situation.

Table 1. Comparison of the Different Models Used

DOI: 10.1161/JAHA.117.006458

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temperature). Endogenous LC3B was visualized by the anti-LC3B antibody and a secondary Alexa-488–labeled anti-rabbit antibody; endogenous MHC was visualized by the anti-MHC (kind gift of Professor J. Van der Velden) and a secondary fluorescein isothiocyanate–labeled anti-rabbit antibody. Endogenous LC3B, green fluorescent protein–LC3B puncta, indicative of autophagosomes, and MHC were visualized by confocal microscopy and captured at ×125 magnification. The number of puncta was counted manually from at least 2 independent experiments using ImagePro. Mean values and SEMs from each experimental condition were based on at least 20 cardiomyocytes in case of transfection and at least 50 in case of drug treatment.

**In Vivo Dog Model for AF**

Adult mongrel dogs were divided into 3 groups: nonpaced, atrial tachypaced (ATP; Table 1) to maintain AF, or ATP with sodium salt of PBA (Na-PBA) treatment (300 mg/kg per day, orally). All dogs underwent the same surgical procedure, AF induction measurements, and cardiomyocyte contractility and electro-physiological measurements. The dogs were anesthetized with acepromazine (0.07 mg/kg IM), ketamine (5.3 mg/kg IV), and isoflurane (1.5%); then, they were intubated and ventilated. One bipolar pacing lead was fixed into the right atrial (RA) appendage via the left jugular vein under fluoroscopic guidance. The tip was connected to a programmable pacemaker. Results in 7 ATP dogs with Na-PBA were compared with 7 tachypaced dogs without treatment and 7 nonpaced control dogs. Na-PBA was given orally (300 mg/kg per day), starting 3 days before and continuing throughout ATP. For the ATP and ATP with Na-PBA groups, the pacemakers were turned on 24 hours after surgery to stimulate the RA at 600 beats per minute for 7 successive days. The ECG was checked daily to ensure AF during pacing. At the end of the study, all dogs were anesthetized with morphine (2 mg/kg SC) and α-chloralose (120 mg/kg IV bolus, followed by 29.25 mg/kg per hour IV infusion); then, they were intubated and ventilated. Body temperature was maintained at 37°C. After midline sternotomy, the pericardium was opened and 2 bipolar electrodes were fixed to the RA appendage (1 for pacing and 1 for signal recording). For AF induction, the RA was paced at 50 Hz for 10 seconds. A total of 5 to 10 AF episodes were recorded to calculate the mean AF duration in each dog. An AF episode >10 minutes was considered sustained, and the electrophysiological study was terminated. Cardioversion was avoided to prevent tissue damage, which precludes further cellular and molecular studies.

**Atrial Cardiomyocyte Isolation**

After electrophysiological study, the heart was excised and immersed in oxygen-saturated Tyrode solution (in mmol/L): 

- NaCl 136, KCl 5.4, MgCl₂ 1, CaCl₂ 2, NaH₂PO₄ 0.33, HEPES 5, and dextrose 10 (pH 7.35), by NaOH. The left atrium (LA) was isolated from the heart with an intact blood supply. The left circumflex coronary artery was cannulated and perfused with Ca²⁺ (1.8 mmol/L), followed by Ca²⁺-free Tyrode solution perfusion for 10 minutes. All leaking branches were ligated. The tissue was then perfused with Ca²⁺-free Tyrode solution containing 150 U/mL collagenase and 0.1% BSA for 60 minutes. Digested LA tissue was harvested and carefully stirred. Isolated cells were centrifuged (500 rpm, 3 minutes) to separate cardiomyocytes from fibroblasts. Cardiomyocytes were stored in Tyrode solution containing 200 μmol/L Ca²⁺ for Ca²⁺-imaging studies.

**Cardiomyocyte Ca²⁺ Imaging and Cellular Contractility Assessment**

Isolated cardiomyocytes were stimulated at 1 Hz, and all measurements were performed at 35±2°C. Cell-Ca²⁺ recording was obtained, as previously described, with the use of Indo-1 AM.²⁻³ Cells were exposed to UV light (wavelength, 340 nm), and the exposure was controlled with an electronic shutter to minimize photographic bleaching. Emitted light was reflected into a spectral separator, passed through parallel filters at 400 and 500 nm (±10 nm), detected by matched photomultiplier tubes, and electronically filtered at 60 Hz. Background fluorescence was removed by adjusting the 400- and 500-nm channels to 0 over an empty field of view near the cell. Fluorescence signal ratios (R) were recorded and converted to [Ca²⁺]i following the equation developed by Grynkiewicz et al:²⁴

\[
\frac{[\text{Ca}^{2+}]_i}{K_d} = \frac{R - R_{\text{min}}}{R_{\text{max}} - R}
\]

where K_d is the ratio of the 500-nm signals at low and saturating [Ca²⁺]i. Intracellular K_d for Indo-1 was 844 nm. Cell and sarcomere contractility was detected by automatic edge detection, and 5 successive beats were averaged for each measurement.

**Cell Electrophysiological Recordings**

Borosilicate glass electrodes filled with pipette solution were connected to a patch-clamp amplifier. Electrodes had tip resistances of 2 to 4 MΩ. For perforated-patch recording, nystatin-free intracellular solution was placed in the tip of the pipette by capillary action (≈30 seconds); then, pipettes were back-filled with nystatin-containing (600 μg/mL) pipette solution. Data were sampled at 5 kHz and filtered at 1 kHz. Whole cell currents are expressed as densities (pA/pF). Junction potentials between bath and pipette solutions averaged 10.5 mV and were corrected for APs only. Tyrode solution contained the following (in mmol/L): NaCl 136, CaCl₂ 1.8, KCl 5.4, MgCl₂ 1, NaH₂PO₄ 0.33, dextrose 10, and HEPES 5, titrated to pH 7.3 with NaOH. The pipette solution for AP
recording contained the following (mmol/L): GTP 0.1, potassium-aspartate 110, KCl 20, MgCl₂ 1, MgATP 5, HEPES 10, sodium-phosphocreatine 5, and EGTA 0.005 (pH 7.4, KOH).

The extracellular solution for Ca²⁺-current measurement contained the following: tetraethylammonium chloride 136, CsCl 5.4, MgCl₂ 1, CaCl₂ 2, NaH₂PO₄ 0.33, dextrose 10, and HEPES 5 (pH 7.4, CsOH). Niflumic acid (50 μmol/L) was added to inhibit Ca²⁺-dependent Cl current, and 4-aminopyridine (2 mmol/L) was added to suppress Ito. The pipette solution for Ca²⁺-current recording contained the following (in mmol/L): CsCl 120, tetraethylammonium chloride 20, MgCl₂ 1, EGTA 10, MgATP 5, HEPES 10, and Li-GTP 0.1 (pH 7.4, CsOH).

Patient Material

Before surgery, 1 investigator assessed patient characteristics (Table 2), as described before. All patients were euthyroid and had normal left ventricular function. RA and LA appendages were obtained from all patients. After excision, the atrial appendages were immediately snap frozen in liquid nitrogen and stored at −85°C. The study conforms to the principles of the Declaration of Helsinki. The institutional review board approved the study, and patients gave written informed consent. Because of the low tissue yield per patient, not all experiments could be performed with each tissue sample. Therefore, at least 5 samples per group were used for experiments.

Table 2. Demographic and Clinical Characteristics of Patients With PeAF and Control Patients in SR

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>SR Group (n=17)</th>
<th>PeAF Group (n=28)</th>
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<tbody>
<tr>
<td>RAA</td>
<td>17 (100)</td>
<td>25 (89)</td>
</tr>
<tr>
<td>LAA</td>
<td>14 (82)</td>
<td>27 (96)</td>
</tr>
<tr>
<td>Age, mean±SEM, y</td>
<td>58±5</td>
<td>61±3</td>
</tr>
<tr>
<td>Duration of AF, median (range), mo</td>
<td>...</td>
<td>8 (0.1–56)</td>
</tr>
<tr>
<td>Underlying heart disease/surgical procedure</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Lone AF/maze</td>
<td>0 (0)</td>
<td>7 (25)*</td>
</tr>
<tr>
<td>CAD/MVI</td>
<td>17 (100)</td>
<td>21 (75)*</td>
</tr>
<tr>
<td>Medication</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Digoxin</td>
<td>1 (6)</td>
<td>13 (46)</td>
</tr>
<tr>
<td>Calcium antagonists</td>
<td>9 (53)</td>
<td>10 (36)</td>
</tr>
<tr>
<td>Blockers</td>
<td>17 (100)</td>
<td>9 (32)</td>
</tr>
</tbody>
</table>

Values are represented as number (percentage) of patients unless otherwise indicated. AF indicates atrial fibrillation; CAD, coronary artery disease; LAA, left atrial appendage; maze, atrial arrhythmia surgery; MVI, mitral valve insufficiency; PeAF, persistent AF; RAA, right atrial appendage; and SR, sinus rhythm.

Statistical Analysis

Results are expressed as mean±SEM of at least 3 independent experiments. Statistical analysis was performed using a Student t test for single comparison between 2 groups. For analysis involving >2 groups, statistical comparison was performed using a 1-way ANOVA. When showing significance, individual group differences were assessed using a Bonferroni-corrected t test. Correlations were estimated using Pearson correlation and tested to be significantly nonzero using Pearson correlation tests. All P values were 2 sided. P≤0.05 was considered statistically significant. SPSS version 20 was used for all statistical evaluations.

Results

Tachypacing of Cardiomyocytes Induces Autophagy

To explore whether tachypacing induces autophagy, the autophagy markers p62 and LC3B were tested. p62 is sequestered to autophagosomes during autophagy and degraded on fusion with the lysosome; reduced levels of p62 are an indication of autophagic activation. LC3B-II is a protein produced from LC3B-I on activation of autophagy and is also incorporated into autophagosomes; LC3B-II levels correlate with the induction of autophagy. Tachypacing of HL-1 atrial cardiomyocytes, in which 8-hour tachypacing produces changes resembling those reported in persistent AF (PeAF) in humans, activates autophagy, as demonstrated by a time-dependent decrease in the expression of p62 and increase in LC3B-II levels (Figure 1A through 1C). No such changes were noted in cardiomyocytes paced at 1 Hz (ie, their average intrinsic firing frequency; Figure S1). For the sake of clarity, results for normal-paced cardiomyocyte data shown in the figures without other specification were obtained after 8 hours of pacing. Furthermore, HL-1 atrial cardiomyocytes show normal morphological characteristics and are viable during normal pacing and tachypacing, as assessed by bright-field microscopy (Figure S2). Tachypacing also induces a clear redistribution of LC3B into discrete perinuclear puncta in both untransfected cardiomyocytes and LC3B-green fluorescent protein transfected cardiomyocytes (Figure 1D through 1F, Figure S3), supporting autophagosome formation. Next, we determined the autophagic flux, to discriminate between the induction of autophagy and decreased degradation of autophagosomes, by blocking autophagosome-lysosome fusion with BAF pretreatment further increases LC3B-II levels of tachypaced cardiomyocytes, but did not affect normal-paced cardiomyocytes (Figure 1G and 1H, Figures S4 and S5). This finding, together with reduced p62 levels,
Figure 1. Tachypacing (TP) induces autophagosome formation and enhanced activation of autophagy. A, Representative Western blot of TP-induced autophagy markers p62 (molecular weight [MW], 62), LC3B-I and LC3B-II (MWs, 14 and 16, respectively), and loading control GAPDH (MW, 37). HL-1 cardiomyocytes were normal paced (NP) or TP for the duration indicated. B, Quantified data showing a significant reduction in p62 levels after 6 or more hours of TP (N=4). C, Quantified data showing a significant increase in LC3B-II levels, beginning after 2 hours of TP (N=5). D, Confocal images of TP HL-1 cardiomyocytes, for the period as indicated, transfected with LC3B–green fluorescent protein (GFP) plasmid. E, Confocal images of TP HL-1 cardiomyocytes for the period as indicated. Endogenous LC3B was visualized by immunostaining. Green puncta indicate autophagosomes. F, Quantified data showing accumulation of LC3B-GFP punctae/cardiomyocytes during TP (n/N=35/3). G, Representative Western blot of HL-1 cardiomyocytes NP vs TP for the duration, as indicated, in the presence or absence of bafilomycin A1 (BAF). H, Quantification of the autophagic flux by determining the difference in LC3B-II levels in the presence vs absence of BAF (N=4). Note that all NP data are shown after 8 hours of observation. *P≤0.05, **P≤0.01, ***P≤0.001 vs NP.
indicates that tachypacing increases functional autophagic activity in HL-1 atrial cardiomyocytes.

**ER Stress Is Associated With Autophagy in a Tachypaced Cardiomyocyte Model**

To investigate which upstream pathway activates autophagy, we first examined the mTOR. mTOR assembles into 2 complexes, mTOR complex (mTORC) 1 and mTORC2; both complexes become activated by mTOR phosphorylation, although at different sites, after which they attenuate autophagy. To test whether tachypacing-induced autophagy results from the inhibition of mTORC2, we next examined involvement of ER stress signaling in tachypacing-induced autophagic flux; Akt S473 phosphorylation is observed during ER stress, and ER stress is an important regulator of autophagy. A role of ER stress was suggested by the finding that tachypacing strongly increases phosphorylation of its downstream effector eIF2α (Figure 3A and 3B, Figure S6), which, on phosphorylation, induces transcription of ER stress and autophagy genes (ie, ATF4, ATF6, CHOP, HSPA5, and ATG12; Figure 3C). In addition, tachypacing gradually induced protein levels of HSPA5 (Figure S7), an endogenous ER chaperone-protein induced by ER stress. These results suggest that ER stress

**Figure 2.** Tachypacing (TP)—induced autophagy does not involve mammalian target of rapamycin complex (mTORC) signaling. Top panels: Western blots of proteins within mTORC signaling. Bottom panels: Quantified data of the ratio of phosphorylated proteins normalized for basal protein levels. Phosphorylated mTOR S2448 (mTORC1; molecular weight [MW], 289; N=3; A), phosphorylated mTOR S2481 (mTORC2; MW, 289; N=3; B), phosphorylated ribosomal protein S6 (S6RP) S235/236 (downstream of mTORC1; MW, 32; N=3; C), and phosphorylated protein kinase B (Akt) S473 (downstream of mTORC2 and endoplasmic reticulum stress; MW, 60; N=3; D) in response to TP for the duration, as indicated, compared with normal pacing (NP). Note that all NP data shown are after 8 hours of observation. **P≤0.01, ***P≤0.001 vs NP.

DOI: 10.1161/JAHA.117.006458

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is the upstream activator of autophagy in the tachypaced cardiomyocyte model.

To extend the findings to human AF, we examined autophagy, ER stress, and markers of cardiac remodeling in atrial appendages of patients with PeAF along with control patients in SR. Patients with PeAF showed an accumulation of autophagosomes and autolysosomes and the presence of myolysis (degradation of sarcomeres) on electron microscopic examination, which is absent in patients in SR (Figure 4A through 4D). Autophagy is further evidenced in patients with AF by enhanced LC3-II induction and decreased levels of p62 compared with patients in SR (Figure 4E). The ER stress chaperone-protein HSPA5 showed a trend towards increased expression in patients with PeAF compared with patients in SR. Previously, we reported on structural remodeling involving degradation of contractile proteins in these patients. Involvement of autophagy in structural remodeling and AF progression is substantiated by the correlation of p62

**Figure 3.** Tachypacing (TP) augments levels of endoplasmic reticulum (ER) stress markers and the autophagy gene **ATG12.** A, Representative Western blot of phosphorylated eIF2α S51 (molecular weight [MW], 38), an ER stress marker, basal eIF2α (MW, 36), and GAPDH levels during normal pacing (NP) or in response to TP for the indicated duration. B, Quantified data of the ratio of phosphorylated eIF2α S51 normalized for basal eIF2α protein levels (N=3). C, Quantitative real-time polymerase chain reaction of ER stress markers ATF4, ATF6, CHOP, and heat shock protein (HSP) A5 and the autophagy-related gene **ATG12** in response to TP for the indicated duration relative to NP (N=3). Note that all NP data shown are after 8 hours of observation. **P≤0.01, ***P≤0.001 vs NP. DOI: 10.1161/JAHA.117.006458

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expression with cardiac troponins (I and T) and α-tubulin expression in patients with PeAF and those in SR (Figure 4F through 4H); there was an inverse correlation with the amount of myolysis (Figure 4I). Levels of p62 also correlated with HSPA5 levels (Figure 4J), suggesting that an ER stress response is associated with autophagy and AF progression. Although 1 data point seems to be an outlier, statistical analysis showed it did not qualify as such. Nevertheless, to exclude undue influence of this data point on correlations, we have repeated the analyses, omitting this data point. For most analyses, correlation remained statistically significant, with the exception of cardiac troponin T and myolysis (Figure 4F: \( R = 0.43, P = 0.075 \); Figure 4G: \( R = 0.74, P < 0.001 \); Figure 4H: \( R = 0.49, P < 0.05 \); Figure 4I: \( R = -0.46, P = 0.056 \); Figure 4J: \( R = -0.62, P < 0.05 \)).

The correlation of autophagy markers with degradation of contractile proteins and the amount of structural remodeling suggest a biologically relevant contribution of this pathway to AF-induced derailment of cardiomyocyte proteostasis and disease progression.

**Inhibition of ER Stress Attenuates Autophagy and Protects From Cardiac Remodeling**

The contribution of AF-induced ER stress and the subsequent enhanced autophagic flux to derailment of cardiomyocyte proteostasis and disease progression was tested by pharmacological and genetic manipulations in experimental model systems for AF. The orphan drug, 4PBA, in clinical use to treat urea cycle disorders,\(^{35-37}\) has recently been recognized as an
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DOI: 10.1161/JAHA.117.006458

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Inhibitor of ER stress by virtue of its chemical chaperone properties.38,39 To explore its potential as a therapeutic agent in AF, we examined its properties in tachypaced cardiomyocytes, Drosophila, and a dog model of AF.

In tachypaced HL-1 cardiomyocytes, 4PBA limits ER stress and prevents activation of autophagy, as demonstrated by normalization of phosphorylated eIF2α expression and LC3B-II and attenuation of p62 breakdown (Figure 5A and 5B). In addition, 4PBA treatment prevents tachypacing-induced accumulation of the contractile protein MHC in perinuclear puncta in HL-1 cardiomyocytes (Figure 5C and 5D). The protective 4PBA effects are mediated via upstream ER stress inhibition, because downstream inhibition of the autophagic process by pepstatin A (a lysosomal cathepsin D/E inhibitor) or BAF (a lysosomal fusion inhibitor) attenuated p62 degradation but did not normalize the phosphorylation of eIF2α, LC3B-II expression, and the formation of perinuclear MHC puncta on tachypacing (Figure 5A through 5D).

Next, we determined whether ER stress results in tachypacing-induced contractile dysfunction. HL-1 cardiomyocytes were pretreated with 4PBA, which caused protection against loss of CaTs in 8-hour tachypaced cardiomyocytes (Figure 5E and 5F, Figure S8A and S8B, Videos S1 through S4). Similar protective effects against CaT loss were observed in tachypaced HL-1 cardiomyocytes overexpressing the endogenous ER chaperone-protein HSPA5, indicating that ER stress is involved in contractile dysfunction (Figure 5G and 5H, Videos S5 through S8). To directly assess whether ER stress is associated with autophagy and contractile dysfunction, HL-1 cardiomyocytes were transfected with eIF2α mutants (wild type, constitutively phosphorylated [S51D], or constitutively nonphosphorylated [S51A]), followed by tachypacing. Tachypaced HL-1 cardiomyocytes overexpressing the nonphosphorylated eIF2α mutant were protected from CaT loss, in contrast to cardiomyocytes overexpressing the wild-type or constitutively phosphorylated eIF2α mutants (Figure 5I and 5J).

Normal-paced HL-1 cardiomyocytes, transfected with the eIF2α mutants, showed no differences in CaT amplitude compared with nontransfected cardiomyocytes (Figure S8C and S8D). The findings indicate that activation of the ER stress pathway is an important modulator of contractile dysfunction. In addition, inhibition of autophagic flux by preincubating HL-1 cardiomyocytes with the autophagy inhibitors pepstatin A and BAF was also protective against tachypacing-induced CaT loss. This again emphasized the role of ER stress–associated autophagy in contractile function (Figure 5E and 5F, Figure S8A and S8B, Videos S9 through S12). Pepstatin A and BAF effects are not conveyed via indirect modulation of ER stress, because neither of the drugs influenced HSPA5 expression levels, as suggested before (Figure S9).40

To extend these findings to a multicellular experimental animal model for tachypacing-induced contractile dysfunction, similar experiments were conducted in Drosophila.3,22 Comparable to findings in tachypaced HL-1 cardiomyocytes, inhibition of ER stress (4PBA) and autophagy (BAF) attenuates tachypacing-induced dysfunction in heart wall contractions in Drosophila (Figure 5K and 5L, Videos S13 through S18), whereas pepstatin A is not protective and toxic at the concentrations applied. Moreover, activators of ER stress (tunicamycin) and autophagy (rapamycin) resulted in ER stress and contractile dysfunction in tachypaced HL-1 cardiomyocytes and Drosophila (Figure S10).

ER Stress Attenuation Relieves Autophagy and Protects From Cardiac Remodeling in an in Vivo Animal Model

To obtain proof of concept that ER stress is involved in AF promotion in a large animal model for AF, dogs were subjected to 7 days of ATP (equal to persistent human AF41), to induce AF-associated atrial remodeling, and were treated with the orally administered Na-PBA (300 mg/kg per day). In isolated atrial cardiomyocytes, Na-PBA treatment protects from tachypacing-induced electrical changes, including shortening of action potential duration and reductions in L-type Ca²⁺ channel current (Figure 6A and 6B). In addition, Na-PBA treatment prevents tachypacing-induced abnormalities in Ca²⁺ handling and associated hypocontractility of isolated atrial cardiomyocytes (Figure 6C through 6F). Finally, Na-PBA conserved the effective refractory period at various sites at both RA and LA and significantly attenuated the vulnerability to AF induction (Figure 6G and 6H). In addition, the protective effect of Na-PBA was not via modulation of HDAC activity, because HDAC levels were not altered by Na-PBA treatment, as has been suggested before (Figure S11).42 Furthermore, Na-PBA reduces markers of ER stress and autophagy in LA tissue of tachypaced dogs, as demonstrated by an increase in p62 level and reductions in LC3B-II and HSPA5 levels compared with nontreated tachypaced dogs (Figure 7A through 7C). Moreover, Na-PBA treatment protected against MHC reduction in tachypaced dogs (Figure 7D), suggesting that attenuation of ER stress results in conservation of contractile protein expression.

Thus, tachypacing induces ER stress–triggered autophagic flux, which plays a prominent role in cardiomyocyte remodeling and AF progression (Figure 8).43 Findings from a clinically relevant dog model for AF indicate that the chemical chaperone 4PBA protects the heart against AF, making 4PBA a potentially interesting drug candidate for treating clinical AF.

Discussion

In the current study, we report that ER stress–associated enhanced autophagic flux appears to constitute an important
mechanism of cardiac remodeling in tachypaced cardiomyocytes, *Drosophila*, dogs, and atrial biopsy specimens from patients with AF. We provide data to show that blocking ER stress, by the chemical chaperone 4PBA, or overexpressing a phosphorylation-blocked mutant of eIF2α inhibits activation of autophagy and, thereby, suppresses cardiomyocyte remodeling in both in vitro and in vivo AF models. Thus, our study points to ER stress as a potential novel druggable target for

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**Figure 5.** Inhibition of endoplasmic reticulum (ER) stress and autophagy protects against tachypacing (TP)-induced contractile dysfunction in HL-1 cardiomyocytes and *Drosophila melanogaster*. A, Representative Western blot of ER stress marker (eIF2α-PS51) and autophagy markers (LC3B-II and p62) in HL-1 cardiomyocytes pretreated with dimethyl sulfoxide (DMSO; control [C]), the autophagy modulator pepstatin A (PepA) or bafilomycin A1 (BAF), or the molecular chaperone 4-phenyl butyrate (4PBA). B, Quantified data showing that HL-1 cardiomyocytes treated with 4PBA reveal attenuation of TP-induced increase in eIF2α-PS51, LC3B-II induction, and reduction in p62. PepA and BAF inhibit lysosomal cathepsin D/E and lysosomal fusion, respectively, and therefore result in an induction of LC3B-II levels and attenuation of p62 reduction without affecting upstream eIF2α-PS51 levels. Open bars represent normal-paced (NP) cardiomyocytes, whereas closed bars represent TP cardiomyocytes after 8 hours of observation. N=3. C, Confocal images of NP and TP HL-1 cardiomyocytes after 8 hours of observation, stained for myosin heavy chain with DMSO (C), 4PBA, BAF, or PepA pretreatment. D, Quantified data showing the number of puncta for the conditions as indicated, all obtained after 8 hours of observation. 4PBA pretreatment protects against the formation of perinuclear puncta (n/N=60/3). E, Representative Ca^2+^ transients (CaT; 5 seconds) of HL-1 cardiomyocytes after NP or TP. HL-1 cardiomyocytes were pretreated with the autophagy modulators PepA or BAF, or the chemical chaperone 4PBA, followed by NP or TP and measurement of CaT. F, Quantified CaT amplitude of HL-1 cardiomyocytes after NP or TP (n/N=60/4). HL-1 cardiomyocytes were pretreated with the autophagy modulators PepA or BAF or the ER chaperone 4PBA. G, Representative CaT (5 seconds) of HL-1 cardiomyocytes transfected with empty plasmid (C) or ER chaperone heat shock protein (HSP) A5, followed by NP or TP. H, Quantified CaT amplitude of NP and TP HL-1 cardiomyocytes transiently transfected with empty plasmid or HSPA5 (n/N=30/3). I, Representative CaT (5 seconds) of HL-1 cardiomyocytes transfected with empty plasmid (C), eIF2α wild-type, nonphosphorylated (S52A), or phosphorylated mimetic (S52D) mutant and followed by NP or TP. J, Quantified CaT amplitude of NP and TP cardiomyocytes transiently transfected with empty plasmid (C), eIF2α wild-type, constitutively nonphosphorylated (S52A), or constitutively phosphorylated (S52D) mutant (n/N=30/3). K, Representative heart wall contractions of *Drosophila* monitored before TP (sinus rhythm [SR]) and after TP with DMSO (C) or PepA, BAF, or 4PBA pretreatment. L, Quantified data showing heart wall contraction rates of *Drosophila* before and after TP with DMSO (C) or PepA, BAF, or 4PBA treatment. Open bars represent NP (in HL-1 cardiomyocytes) or spontaneous heart rate (SR; in *Drosophila*), and closed bars represent TP HL-1 cardiomyocytes or *Drosophila*. N=9 to 15 prepupae for each group. Note that all NP data shown are after 8 hours of observation. *P≤0.05, **P≤0.01, ***P≤0.001 vs control NP or before TP, #P≤0.05, ##P≤0.01, ###P≤0.001 vs control (after) TP.
cardiac remodeling in AF and proposes that 4PBA may emerge as a novel lead compound for the development of agents to attenuate AF progression.

Prominent Role of ER Stress–Associated Autophagy in Cardiomyocyte Remodeling

Although it is recognized that the increased atrial activation rate constitutes a major driving force for cardiac remodeling in AF,1 molecular events leading to remodeling have been poorly identified. We examined both experimental and clinical AF models that equal persistent human AF, which show reversible electrical and irreversible structural remodeling.3,8,28,44,45 We were able to reveal a prominent role for ER stress–associated enhanced autophagic flux in cardiomyocyte remodeling and AF progression using various pharmacological and genetic manipulations of the ER stress pathway (Figure 8). First, tachypacing-induced contractile dysfunction of HL-1 cardiomyocytes coincided with activation of known key players of this pathway. These include both phosphorylation of the ER stress regulator eIF2α at S51 and downstream expression of the stress-responsive transcripts ATF4 and ATF6. In turn, ATF4 and ATF6 activate autophagy,10 via enhanced expression of CHOP and the autophagy genes ATG12 and LC3B, resulting in the elongation of autophagosomes and a sustained and excessive autophagic flux, as observed in tachypaced HL-1 cardiomyocytes.12,46–48 Second, our data demonstrate that induction of ER stress represents the upstream event in tachypacing-elicited contractile-protein accumulation and contractile dysfunction. Genetic overexpression of a phosphorylation-blocked eIF2α protein or the ER chaperone HSPA5 abrogated both autophagy and contractile dysfunction. Knockdown of autophagy genes, such as ATG5 or ATG7, was avoided because of the detrimental effects in healthy cardiomyocytes.15,49–51 A prominent role for ER stress–induced autophagy in AF promotion is supported by pharmacological interventions. We observed that inhibition of the autophagic process by pepstatin A (a lysosomal cathepsin D/E inhibitor) or BAF (a lysosomal fusion inhibitor) protected against contractile dysfunction, but did not prevent the ER stress response and accumulation of contractile proteins within the HL-1 cardiomyocytes. Pharmacological prevention of ER stress by the chemical chaperone 4PBA precluded ER stress–related autophagy and cardiac remodeling in tachypaced HL-1 cardiomyocytes and Drosophila, as well as in the tachypaced dog model of AF-associated remodeling. Finally,
ER stress and autophagy are also activated in clinical AF, as evidenced by the presence of autophagosomes and autolysosomes in atrial heart tissue, enhanced LC3B-II expression, and reduction in p62 levels in patients with PeAF. Although altered proteasome function can influence p62 expression, proteasome function is not altered in AF, suggesting that the reduction in p62 levels is a consequence of autophagic flux activation. Although there is only a trend towards an increase in HSPA5 expression in patients with AF, diminished protein synthesis or enhanced degradation, by either the

Figure 6. Sodium salt of phenyl butyrate (Na-PBA) protects against atrial remodeling in a dog model for atrial fibrillation (AF). Atrial tachypacing (ATP) induces atrial remodeling, measured as shortening of action potential duration (APD90; A), reduced L-type Ca\(^{2+}\) current (ICaL; B), and increased diastolic calcium levels in cardiomyocytes (n=15–40 cardiomyocytes; C). D, Representative calcium transient (CaT) and cell shortening (CS) tracers for the conditions, as indicated. Furthermore, ATP results in loss of CaT amplitude (E), loss of contractility (F), reduced adaptation of the effective refractory period (ERP) at different basic cycle lengths (BCLs; G), and increased duration of induced AF (H). All ATP-induced atrial remodeling end points were significantly attenuated by Na-PBA treatment. C indicates control. *P\(\leq\)0.05, **P\(\leq\)0.01, ***P\(\leq\)0.001 vs C; #P\(\leq\)0.05, ##P\(\leq\)0.01, ###P\(\leq\)0.001 vs ATP.
proteasome or autophagy, or exhaustion of the protein levels may be the underlying cause.53 There is ongoing debate about whether autophagy plays a beneficial or detrimental role in cardiac diseases.54,55 Excessive autophagy contributed to age-related cardiac disease development, including heart failure, hypertension-induced cardiac diseases, mitral regurgitation, and diabetic cardiomyopathy.16,19,56–58 Interestingly, all these cardiac diseases are recognized to represent a substrate for AF,59 suggesting a role for autophagic activation in AF development. This is supported by the presence of autophagosome accumulation in patients who developed postoperative AF.60 On the other hand, in inherited cardiomyopathies, autophagic activation was found to be beneficial.51–63 In inherited cardiomyopathies, proteotoxic mutant proteins are likely cleared by autophagy. This assumption is strengthened by a study showing that autophagy could be induced as a cellular defense mechanism against ER stress–mediated cell death by degrading protein aggregates.14 Hence, it is clear that autophagic activation in the diverse cardiac diseases is not uniform in its function, but depends on the origin of cardiac disease; it can have either a beneficial (inherited cardiomyopathies) or detrimental (age-related cardiac diseases) effect on the course and outcome of the disease. Because protein aggregation has not been observed in most age-related cardiac diseases, including AF,55 our findings suggest that the autophagic machinery becomes overengaged with maladaptive consequences, possibly because of divergent Ca2+ handling in the ER. The ER plays a prominent role in proper cell function, because at least one third of all proteins are synthesized in this organelle. ER

Figure 7. Sodium salt of phenyl butyrate (Na-PBA) protects against endoplasmic reticulum stress and autophagy in a dog model for atrial fibrillation. A, Top panel: Representative Western blot. Bottom panel: Quantified data revealing a significant reduction in p62 levels in atrial tachypacing (ATP), which was not significantly reduced by Na-PBA treatment compared with control (C) dogs. B, Representative Western blot of LC3B-I/II and loading control β-actin (molecular weight [MW], 43) in groups, as indicated. ATP causes significant induction in LC3B-II levels, which was significantly reduced in case of Na-PBA treatment. C, Representative Western blot of heat shock protein (HSP) A5, showing a trend (P=0.058) in induction of HSPA5, which was not altered in Na-PBA–treated group. D, Representative Western blot of myosin heavy chain (MHC; MW, 230) in groups, as indicated. ATP causes a significant reduction in MHC, which was not changed in case of Na-PBA treatment. N=7 dogs for each group. *P≤0.05 vs C, #P≤0.05 vs ATP.

DOI: 10.1161/JAHA.117.006458

Journal of the American Heart Association
chaperone proteins, especially HSPA5, assist in the correct folding of the newly formed proteins. Reduced levels of HSPA5, or calcium overload in the ER, cause proteins to unfold and produce an ER stress response. In turn, this results in the activation of the transcription factor ATF4, which regulates the expression of autophagy genes and LC3B, causing activation of autophagy by stimulating induction and elongation of autophagosomes. Initially, AF-induced activation of autophagy may preserve cardiomyocyte proteostasis; however, excessive stress-induced autophagy contributes to loss of contractile function and cardiac remodeling. AF stress–induced autophagy appears maladaptive, because inhibition of autophagy via 4-phenyl butyrate (4PBA), HSPA5, or nonphosphorylatable mutant eIF2α (S51A) overexpression, pepstatin A (PepA), and bafilomycin A1 (BAF) prevented AF-associated remodeling and progression in our studies. MHC indicates β-myosin heavy chain 7.

**Therapeutic Implications**

From a translational perspective, the current results identify a potential benefit of pharmacological inhibition of ER stress as a therapeutic strategy in clinical AF. Among the available compounds, 4PBA seems promising, because this compound is already approved for clinical use to treat urea cycle disorders and is available under the trade names Buphenyl (available in the United States since 1996) and Ammonaps (available in Europe since 1999). 4PBA acts as a chemical chaperone and alleviates ER stress by protecting from aggregation of misfolded proteins. There are several ongoing human trials with 4PBA, which target ER stress in various clinical diseases featuring protein misfolding, including amyotrophic lateral sclerosis (NCT00107770), Huntington disease (NCT00212316), spinal muscular atrophy (NCT00528268), proteinuric nephropathies (NCT02343094), and cystic fibrosis (NCT00016744). Data from patients with urea cycle disorders to date indicate that 4PBA is safe and displays minor adverse effects, although conventional dosing is high (maximum, 20 g/d). In addition, our results indicate that cardiac remodeling in AF may also be modulated by inhibitors of autophagy. However, treatment with autophagy inhibitors might not be suitable for all patients, and further research is needed to identify the optimal therapeutic strategy.
inhibitors may precipitate considerable toxicity, as reported for bafilomycin, or additional detrimental effects because of disruption of normal cell physiological characteristics by inhibition of basal autophagy. Application of inhibitors of autophagy in AF and other chronic conditions thus awaits development of selective inhibitors targeting excessive autophagy. On the basis of these considerations, 4PBA may serve as a useful compound to explore the benefits of repression of ER stress–associated autophagy in the attenuation of AF progression and improvement of cardioversion outcome in clinical AF. 4PBA may serve as a lead compound for the further development of autophagy inhibitors for clinical use.

Limitations

HL-1 atrial cardiomyocytes are derived from mouse atria and generally show similar features as adult cardiomyocytes. Despite potential differences, the ease of confirmation of specific molecular pathways conferring tachypaced-induced remodeling by use of genetic manipulation is an important advantage of the HL-1 model. Moreover, findings in the tachypaced HL-1 atrial cardiomyocyte model have been confirmed repeatedly in the tachypaced dog model and clinical human AF. Therefore, the tachypaced HL-1 model has merit to identify potential signaling pathways involved in AF remodeling. The patient groups differed in terms of medications prescribed, as expected on the basis of the different disease causes of each group. For example, patients with AF frequently received digoxin for rate control, whereas patients in SR almost never took digoxin. On the other hand, patients in SR requiring surgery for coronary artery disease, like our control group, almost always receive β blockers, whereas a minority of patients with PeAF take them for rate control. Adjustments for these differences in our overall population cannot be made because of too few individuals; effects of drugs may also differ between SR and PeAF populations. Nevertheless, changes in patient data were similar to those observed in the in vitro HL-1 atrial cardiomyocyte model and the in vivo dog model.

Sources of Funding

This study was supported by the Dutch Heart Foundation (2013T096 and 2013T144), LSH-Impulse grant (40-43100-98-008), The Netherlands Cardiovascular Research Initiative, Dutch Heart Foundation CVON2014-40 DOSIS and CVON-STW2016-14728 AFFIP, NWO VICI grant (865.10.012 to Sibon), and the Canadian Institutes of Health Research (Foundation grant). Nattel received support from the Quebec Heart and Stroke Foundation.

Disclosures

None.

References

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Figure S1. Normal pacing does not change protein expression.

Representative Western blot showing LC3B-I/II, p62, eIF2α-P_S51, and mTOR-P_S2481 expression in normal paced HL-1 cardiomyocytes. Normal pacing does not change the protein expression levels of any of these proteins.
Figure S2. Normal pacing and tachypacing does not change cardiomyocyte morphology.

Bright field microscopic images showing the morphology of the HL-1 atrial cardiomyocytes after non-pacing (0 h), normal pacing (1 Hz) or tachypacing (6 Hz). No morphological changes were observed between the conditions as indicated.
Figure S3. Tachypacing induces autophagosome formation.

Confocal images of multiple tachypaced HL-1 cardiomyocytes, for the period as indicated, (A) transfected with LC3B-GFP or (B) endogenous LC3B visualized by immunostaining. Green puncta indicate autophagosomes.
Figure S4. LC3B-II levels are not increased during normal pacing after BAF treatment.

Representative Western blot showing LC3B-I/II expression in normal paced HL-1 cardiomyocytes with and without BAF treatment. BAF doesn't increase the LC3B-II levels during normal pacing.
Figure S5. BAF treatment further increases LC3B-II levels in tachypaced HL-1 cardiomyocytes.

Representative Western blot showing LC3B-I/II expression in normal paced (NP) and 8 h tachypaced (TP) HL-1 cardiomyocytes with and without BAF treatment. BAF increases the LC3B-II levels after 8h tachypacing.
Figure S6. Normal pacing does not change eIF2α phosphorylation.

Representative Western blot showing eIF2α phosphorylation in normal paced (NP) and tachypaced (TP) HL-1 cardiomyocytes.
Figure S7. Tachypacing induces expression of the ER chaperone HSPA5.

Top panel: representative Western blot showing HSPA5 and GAPDH expression in normal paced (NP) and tachypaced (TP) HL-1 cardiomyocytes. Bottom: quantified data revealing significant increase in HSPA5 levels in tachypaced HL-1 cardiomyocytes (N=3). *P≤0.05 vs NP.
Figures S8. The effect of pharmacological and genetic modulation of autophagy and ER stress in normal paced and tachypaced HL-1 cardiomyocytes.

(A) Representative CaT (5 sec) of HL-1 cardiomyocytes after normal pacing (NP). HL-1 cardiomyocytes were pre-treated with the autophagy modulators PepA or BAF, or the chemical chaperone 4PBA, followed by normal and measurement of CaT. (B) Quantified CaT of HL-1 cardiomyocytes pretreated with pepstatin A, bafilomycin A1 or 4PBA and subjected to normal pacing, which does not change calcium transients (n/N=60/4). (C) Representative CaT (5 sec) of HL-1 cardiomyocytes transfected with empty plasmid (Control), eIF2α wild-type, non-phosphorylated (S52A) or phospho-mimetic (S52D) mutant and followed by NP. (D) Quantified CaT of HL-1 cardiomyocytes transiently transfected with empty plasmid (Control), eIF2α wild-type, non-phosphorylated (S52A) or phospho-mimetic (S52D) mutant and subjected to normal pacing, which does not change calcium transients (n/N=30/3).
Figure S9. Pharmacological modulation of autophagy in HL-1 cardiomyocytes does not change HSPA5 expression.

Top panel: representative Western blot showing HSPA5 and GAPDH levels in HL-1 cardiomyocytes pretreated with various autophagy modulators as indicated. Lower panel: quantified data showing no significant changes in HSPA5 levels for the conditions as indicated (N=3).
Figure S10. Activators of ER stress and autophagy do not protect against tachypacing-induced contractile dysfunction in HL-1 cardiomyocytes and *Drosophila*.

(A) Representative Western blot showing eIF2α-P$_{S51}$, p62, LC3B-I/II and GAPDH levels in HL-1 cardiomyocytes pretreated the ER stress-inducer tunicamycin (TM), the autophagy-inducer rapamycin (RP), autophagy inhibitors BAF and pepstatin A and the chemical chaperone 4PBA. RP and TM show no protection against tachypacing-induced changes in eIF2α-P$_{S51}$, p62 or LC3B-I/II expression. (B) Representative CaT of HL-1 cardiomyocytes after normal pacing (NP) or tachypacing (TP), pre-treated with TM or RP. (C) Quantified CaT amplitude of NP and TP HL-1 cardiomyocytes, each from groups as indicated (n/N=50/3). There is no significant decrease in CaT amplitude for either RP or TM at NP, due to the non-toxic concentrations applied.
Nevertheless, RP and TM did not protect against contractile dysfunction. (D) Representative heart wall contractions of Drosophila monitored before TP and after TP with DMSO (Control), TM or RP pretreatment. (E) Quantified data showing heart wall contraction rates from groups as indicated. RP and TM did not protect against contractile dysfunction. White bars represent normal paced (NP in HL-1 cardiomyocytes) or spontaneous heart rate (SR in Drosophila) and black bars represent tachypaced HL-1 cardiomyocytes or Drosophila. N=9 to 15 prepupae for each group. **P≤0.01, ***P≤0.001 vs control SR, ###P≤0.001 vs control TP
Figure S11. 4PBA has no effect on HDAC activity in dogs.

Atrial tachypacing of dogs results in a borderline significant induction ($P=0.06$) of HDAC activity, which was not altered by 4PBA treatment ($N=7$ dogs for each group).
Supplemental Video Legends:

**Video S1.** Time-lapse video shows CaT after 8 hours normal pacing (1Hz) of HL-1 cardiomyocytes. Images were acquired at 2 ms intervals.

**Video S2.** Time-lapse video shows CaT after 8 hours tachypacing (6Hz) of HL-1 cardiomyocytes. Images were acquired at 2 ms intervals.

**Video S3.** Time-lapse video shows CaT after 8 hours tachypacing (1Hz) of HL-1 cardiomyocytes pretreated with 4PBA. Images were acquired at 2 ms intervals.

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**Video S5.** Time-lapse video shows CaT after 8 hours tachypacing (1Hz) of HL-1 cardiomyocytes transfected with empty plasmid. Images were acquired at 2 ms intervals.

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**Video S7.** Time-lapse video shows CaT after 8 hours tachypacing (1Hz) of HL-1 cardiomyocytes transfected with HSPA5 construct. Images were acquired at 2 ms intervals.

**Video S8.** Time-lapse video shows CaT after 8 hours tachypacing (6Hz) of HL-1 cardiomyocytes transfected with HSPA5 construct. Images were acquired at 2 ms intervals.
Video S9. Time-lapse video shows CaT after 8 hours tachypacing (1Hz) of HL-1 cardiomyocytes pretreated with pepstatin A. Images were acquired at 2 ms intervals.

Video S10. Time-lapse video shows CaT after 8 hours tachypacing (6Hz) of HL-1 cardiomyocytes pretreated with pepstatin A. Images were acquired at 2 ms intervals.

Video S11. Time-lapse video shows CaT after 8 hours tachypacing (1Hz) of HL-1 cardiomyocytes pretreated with BAF. Images were acquired at 2 ms intervals.

Video S12. Time-lapse video shows CaT after 8 hours tachypacing (6Hz) of HL-1 cardiomyocytes pretreated with BAF. Images were acquired at 2 ms intervals.

Video S13. Time-lapse video shows heart wall contractions of a prepupae of a W1118 genetic background before tachypacing, and after tachypacing (5Hz, Video S14).

Video S15. Time-lapse video shows heart wall contractions of a prepupae of a W1118 genetic background pretreated with BAF before tachypacing, and after tachypacing (5Hz, Video S16).

Video S17. Time-lapse video shows heart wall contractions of a prepupae of a W1118 genetic background pretreated with 4PBA before tachypacing, and after tachypacing (5Hz, Video S18).
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Supplemental Video Legends:

**Video S1.** Time-lapse video shows CaT after 8 hours normal pacing (1Hz) of HL-1 cardiomyocytes. Images were acquired at 2 ms intervals.

**Video S2.** Time-lapse video shows CaT after 8 hours tachypacing (6Hz) of HL-1 cardiomyocytes. Images were acquired at 2 ms intervals.

**Video S3.** Time-lapse video shows CaT after 8 hours tachypacing (1Hz) of HL-1 cardiomyocytes pretreated with 4PBA. Images were acquired at 2 ms intervals.

**Video S4.** Time-lapse video shows CaT after 8 hours tachypacing (6Hz) of HL-1 cardiomyocytes pretreated with 4PBA. Images were acquired at 2 ms intervals.

**Video S5.** Time-lapse video shows CaT after 8 hours tachypacing (1Hz) of HL-1 cardiomyocytes transfected with empty plasmid. Images were acquired at 2 ms intervals.

**Video S6.** Time-lapse video shows CaT after 8 hours tachypacing (6Hz) of HL-1 cardiomyocytes transfected with empty plasmid. Images were acquired at 2 ms intervals.

**Video S7.** Time-lapse video shows CaT after 8 hours tachypacing (1Hz) of HL-1 cardiomyocytes transfected with HSPA5 construct. Images were acquired at 2 ms intervals.

**Video S8.** Time-lapse video shows CaT after 8 hours tachypacing (6Hz) of HL-1 cardiomyocytes transfected with HSPA5 construct. Images were acquired at 2 ms intervals.
**Video S9.** Time-lapse video shows CaT after 8 hours tachypacing (1Hz) of HL-1 cardiomyocytes pretreated with pepstatin A. Images were acquired at 2 ms intervals.

**Video S10.** Time-lapse video shows CaT after 8 hours tachypacing (6Hz) of HL-1 cardiomyocytes pretreated with pepstatin A. Images were acquired at 2 ms intervals.

**Video S11.** Time-lapse video shows CaT after 8 hours tachypacing (1Hz) of HL-1 cardiomyocytes pretreated with BAF. Images were acquired at 2 ms intervals.

**Video S12.** Time-lapse video shows CaT after 8 hours tachypacing (6Hz) of HL-1 cardiomyocytes pretreated with BAF. Images were acquired at 2 ms intervals.

**Video S13.** Time-lapse video shows heart wall contractions of a prepupae of a W1118 genetic background before tachypacing, and after tachypacing (5Hz, **Video S14**).

**Video S15.** Time-lapse video shows heart wall contractions of a prepupae of a W1118 genetic background pretreated with BAF before tachypacing, and after tachypacing (5Hz, **Video S16**).

**Video S17.** Time-lapse video shows heart wall contractions of a prepupae of a W1118 genetic background pretreated with 4PBA before tachypacing, and after tachypacing (5Hz, **Video S18**).
Endoplasmic Reticulum Stress Is Associated With Autophagy and Cardiomyocyte Remodeling in Experimental and Human Atrial Fibrillation

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*J Am Heart Assoc.* 2017;6:e006458; originally published October 24, 2017;
doi: 10.1161/JAHA.117.006458

The *Journal of the American Heart Association* is published by the American Heart Association, 7272 Greenville Avenue, Dallas, TX 75231
Online ISSN: 2047-9980

The online version of this article, along with updated information and services, is located on the World Wide Web at:
http://jaha.ahajournals.org/content/6/10/e006458

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