

# Effects of β-adrenoceptor stimulation in human atrial repolarizing currents

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### INTRODUCTION

Atrial fibrilation (AF) is the most prevalent arrhythmia and the main risk factor associated with myocardial-related cerebrovascular events (1). Nowadays, pharmacological treatment of AF is clearly suboptimal (2), mainly due to rapid changes (4 to 6 hours after the onset) in the electrical properties of the atria (electrical remodeling) induced by the arrhythmia itself (3). This electrical remodeling promotes the maintenance and recurrence of AF (4), and it is characterized by a marked shortening of the atrial action potential duration (APD) and refractoriness as a consequence of changes in Ca<sup>2+</sup> and K<sup>+</sup> channel density (5). Our group has described that chronic AF (CAF) reduced the transient outward (I<sub>to1</sub>) and the ultrarapid delayed rectifier (I<sub>Kur</sub> or I<sub>sus</sub>) K<sup>+</sup> action potential united and the reaction (Ar I) and refraction diseased a large and the classified entering the contract and the classified entering the classified entering the contract and the classified entering the contract and the classified entering the classified entering the classified entering the contract and the classified entering t Associated with an increased atrial sympathetic innervation (8), suggesting that autonomic remodeling may be part of atrial substrate for AF. Stimulation of β-adrenoceptors inhibited I<sub>tot</sub> in dog Purkinje myocytes (9), but increased I<sub>sus</sub> in human RA myocytes (10) and I<sub>ks</sub> in guinea-pig ventricular myocytes (11). Furthermore, it has been shown that the increase of the L-type Ca<sup>2+</sup> current induced by β-adrenoceptor stimulation is potentiated by CAF (12). However, data on the effects of β-adrenoceptor stimulation on voltage-dependent K<sup>+</sup> repolarizing currents in patients with CAF are unavailable. Thus, in this study we analyzed the effects of isoproterenol, a β-adrenoceptor agonist, on I<sub>tot</sub>, I<sub>kur</sub>, and I<sub>ks</sub> recorded in isolated myocytes obtained from RA and LA appendages (RAA and LAA, respectively) obtained from sinus rhythm (SR) and CAF patients.

### **MATERIAL & METHODS**

- Human atrial myocytes were enzymatically isolated from RAA and LAA samples obtained from SR and CAF patients that underwent cardiac surgery at the Hospital Gregorio Marañón in Madrid (6,13-17).

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- at the end of the puse,  $\frac{1}{16}$  as the diinefence between the current amplitudes at the end of the 3+5 depointing and in end of a 4+5 depointing and puse and  $\frac{1}{16}$  was measured as the diinefence between the puse,  $\frac{1}{16}$  as the diinefence between the puse,  $\frac{1}{16}$  as the diinefence between the puse,  $\frac{1}{16}$  as the diinefence of pushed in the pushes,  $\frac{1}{16}$  as the diinefence of pushes and  $\frac{1}{16}$  as the diinefence of pushes and  $\frac{1}{16}$  as the diinefence of pushes and  $\frac{1}{16}$  and  $\frac{1}{16}$  as the diinefence between the peaks of the current amplitudes at the end of the 30+0-15 pushes and  $\frac{1}{16}$  an
- 8, KCI 40, Mg-AI P 5, EG IA 5, C&O; と, G I P U.1, and TEPES 10 UPT (-A, WILL TAUT).

  \*\*To simulate the shapes of human atrial action potentials, a mathematical model previously validated and used for identical purposes was employed (20)

  \*\*mRNA was isolated from human atrial appendages and quantitative reverse transcription polymerase chain reaction (qPCR) analysis was performed (6)

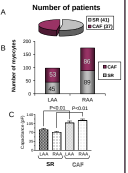


Figure 1. A-C, Distribution of patients (A), number (B), and mean capacitance values (C) of LAA and RAA myocytoobtained from SR and CAF patients.

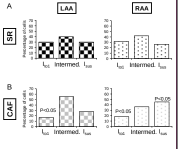


Figure 2. Types of cells according the predominant  $K^{\star}$  current during plateau. A and B, Bar graphs showing the percentage of cells

that exhibited I<sub>to1</sub>-predominant, I<sub>sus</sub>-predominant, and intermediate patterns in SR (A) and CAF (B) myocytes.

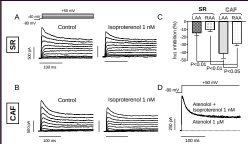


Figure 3. Isoproterenol inhibits human atrial I<sub>to1</sub>. A and B, Effects of isoproterenol on K\* currents elicited in two RAA cells obtained from an SR (A) and a CAF (B) patient. **C**, Percentage of isoproterenol-induced I<sub>to1</sub> inhibition at +30 mV in LAA and RAA myocytes from SR and CAF patients. Each bar represents the mean±SEM of n>8. **D**, Effects of isoproterenol in the presence of atenolol on K+ currents recorded in an RAA myocyte from a CAF patient.

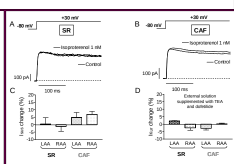


Figure 4. Isoproterenol (1 nM) does not modify the I. Effects of isoproterenol on outward K<sup>+</sup> currents recorded in I<sub>sus</sub> predominant RAA cells obtained from an SR (A) and a CAF (B) patient. C and D, Percentage of I<sub>sus</sub> (C) and I<sub>Kur</sub> (D) change at +30 mV induced by isoproterenol on LAA and RAA cells from SR and CAF patients. Each point represents the mean±SEM of n>10.

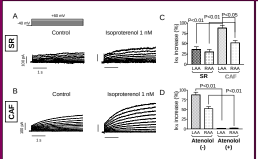


Figure 5. Isoproterenol increases human atrial  $I_{KS}$ . A and B, Effects of isoproterenol on 2 mM 4-AP-resistant K $^{+}$  currents elicited in two RAA cells obtained from an SR (A) and a CAF (B) patient. **C**, Percentage of isoproterenolinduced I<sub>kg</sub> increase at +30 mV in LAA and RAA myocytes from SR and CAF patients. D, Percentage of isoproterenol-induced I<sub>kg</sub> increase at +30 mV in LAA and RAA myocytes from CAF patients in the absence and presence of atenolol. Each bar represents the mean-SEM of n>8.

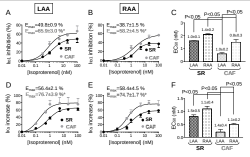


Figure 6. Concentration-dependent effects of isoproterenol on  $I_{tot}$  and  $I_{Ks}$ - A-F  $I_{tot}$  density reduction (A and B) and  $I_{ks}$  density increase (D and E) at +30 mV as a function of isoproterenol concentration in LAa (A and D) and RAa (B and E) myocytes from SR and CAF patients. Continuous lines represent the fit of a Hill equation to the data. "P-0.05 vs SR. EC<sub>50</sub> values for the isoproterenol-induced  $I_{\rm tot}$  inhibition (C) and  $I_{\rm Ks}$  increase (F). In C and F, Hill coefficients appear over the data bar. Each point/bar represents the mean±SEM of n>8.

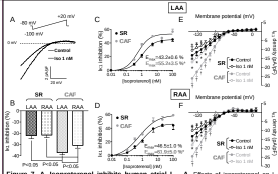


Figure 7. A, Isoproternol inhibits human atrial I<sub>K1</sub>. A, Effects of isoproternol on I<sub>K1</sub> recorded by applying a voltage-ramp (800 ms) in a LAA mycoyte from a CAF patient. B, toproternol-induced I<sub>K1</sub> inhibition at -100 mV in LAA and RAA mycoytes from SR and CAF patients. C and D, Concentration-dependent I<sub>K1</sub> inhibition produced by isoproterenol at -100 mV in LAA (C) and RAA (D) mycoytes from SR and CAF patients. E and F, Effects of isoproterenol on I<sub>K1</sub> density-voltage curves obtained in LAA (E) and RAA (F) mycoytes from SR and CAF patients. Each point/bar represents the mean±SEM of n-8. \*P<0.05 vs. control. †P<0.05 vs SR.

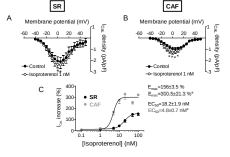
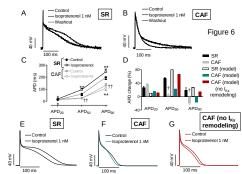


Figure 8. Isoproterenol increases human atrial  $I_{\rm Cal.}$  A and B, Effects of 1 nM isoproterenol on the current density-voltage relationships for  $I_{\rm Cal.}$  recorded in RAA myocytes from SR (A) and CAF (B) patients. \*P<0.05 vs control. The protocol to obtain current-voltage relationships of  $l_{\rm CaL}$  consisted of 500-ms pulses that were imposed in 5 mV increments between -40 and +50 mV. C, Percentage of isoproterenol-induced  $l_{\rm caL}$  increase at +5 mV in RAA myocytes from SR and CAF patients. Continuous lines represent the fit of a Hill equation to the data. \*P<0.05 vs SR



100 ms 100 ms 9. CAF modifies the effects of isoproterenol on human atrial action potentials (APs), A B, Effects of isoproterenol on APs recorded in RAA mycotyes obtained from an SR (A) and a (B) patient, C, Effects of isoproterenol on APs The measured at 20%, 50%, and 90% of artication. "P-0.05 vs control. "P-0.01 vs control." TP-0.01 vs Control TP-0.01 vs SR. D, Percentage of change in PD produced by isoproterenol in SR and CAF mocycles. "P-0.05 vs SR. In C and D, each har represents the mean-SEM of n>5. E and F, Mathematically modeled steady-state APs end in an SR (C), a CAF (F), and a CAF mycotyce without considering the CAF-induced increase on I<sub>Ks</sub> (G) in absence and presence of isoproterenol at a frequency of 1 Hz

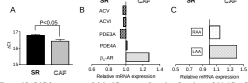


Figure 10. CAF increases atrial β<sub>1</sub>-RA expression. A<sub>1</sub> ΔCt values of β1-RA mRNA measured by qPCR in samples obtained from SR and CAF patients (pooled data). B<sub>1</sub>, Relative expression level of ACV, ACVI, PDE3A, PDE4A, and β<sub>1</sub>-RA mRNA in SR and CAF samples. C<sub>1</sub>, Relative expression level of β1-AR mRNA in SR and CAF samples when considering LAA and RAA separately. Each bar represents the mean±SEM of n>5.

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- CONCLUSIONS CAF potentiates the inhibition of the  $\rm I_{to1}$  and the increase of the  $\rm I_{Ks}$  produced by β-AR stimulation, this effect being greater in LAA than in RAA myocytes.
- CAF potentiates the  $\beta$ -adrenergic-induced increase of the  $I_{Cal}$ .
- $\beta$ -adrenergic stimulation does not modify the  $I_{Kur}$  either in SR or in CAF myocytes and inhibits  $I_{K1}$  only at potentials negative to the equilibrium notential for K+
- The CAF-induced potentiation of the  $\beta$ -adrenergic effects on human atrial ion currents can be attributed to an increase in the β1-AR expression. Moreover, the mRNA expression of the \$1-AR is higher in LAA than in RAA samples.
- The increase in  $\beta$ 1-AR expression as well as the ion channel derangements produced by CAF, could account for the different effects produced by the  $\beta\text{-AR}$ stimulation on the APD in myocytes from SR (prolongation) and CAF patients (shortening).
- The CAF-induced increase on  $I_{\mbox{\scriptsize Ks}}$  is critical to account for the  $\beta\mbox{\scriptsize 1-AR-induced}$ shortening of APD in CAF myocytes.
- The CAF-induced potentiation of the effects of \( \beta 1\)-adrenoceptor stimulation on human atrial K+ currents could contribute to the shortening of APD observed in CAF and, thus, to promote reentry.

## **FUNDING**