

# *Atrial Fibrillation: Upstream Therapies*

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# Effects of the RAAS on AF

- In human AF, tissue ACE is up-regulated correlating with increased atrial angiotensin II production<sup>1</sup>
- ACE inhibition has been shown to attenuate atrial structural remodeling (interstitial fibrosis) in a canine rapid ventricular pacing model<sup>2</sup> and reduce AF in post-MI LVD patients<sup>3</sup>
- There is an association of RAAS gene polymorphisms in patients with non-familial structural AF<sup>4</sup>
  - Are such patients more likely to benefit from ACEI and ARBs?

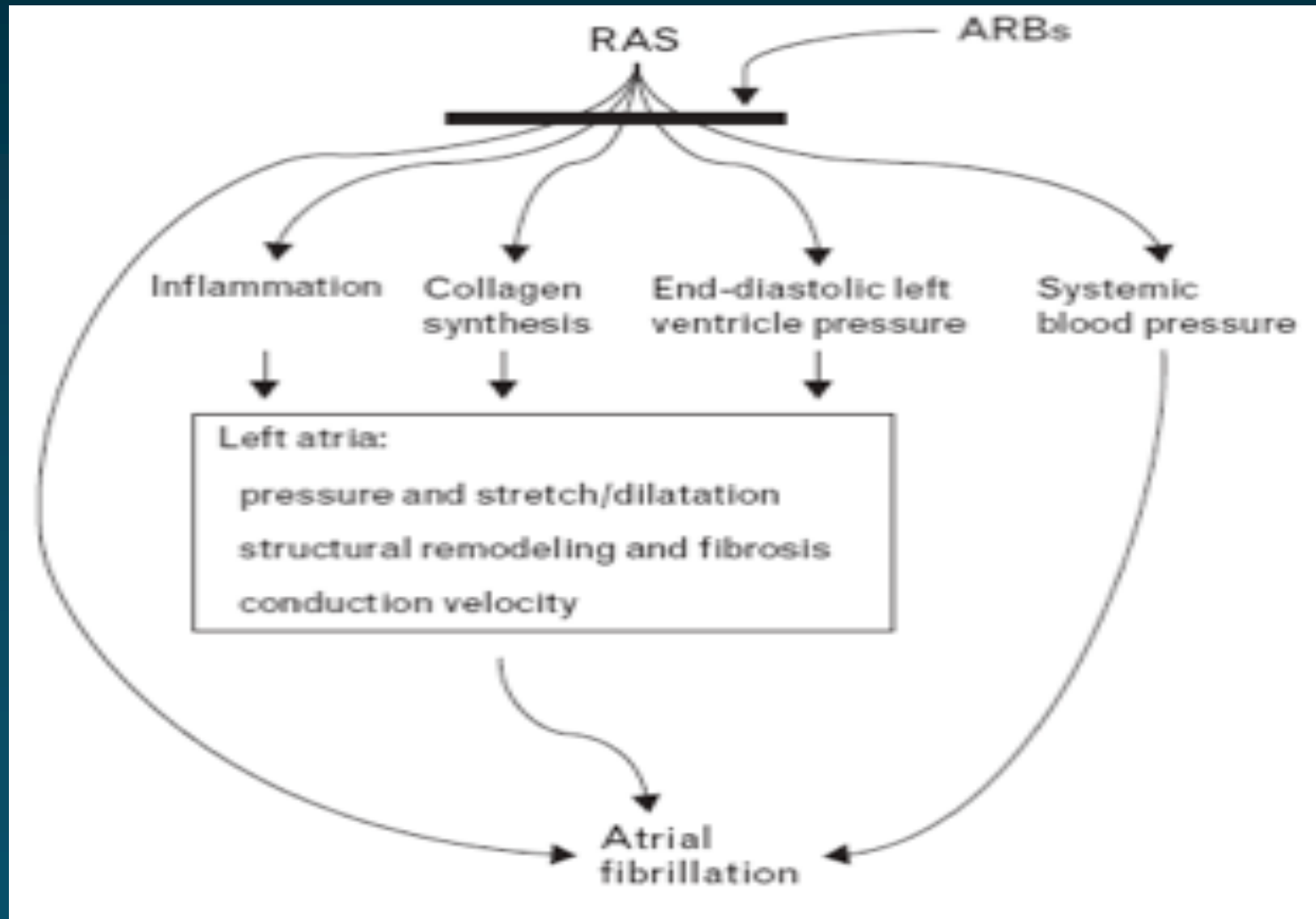
1. Goette A, et al. *J Am Coll Cardiol* 2000;35:1669-1677

2. Li D, et al. *Circulation* 1999;100:87-95

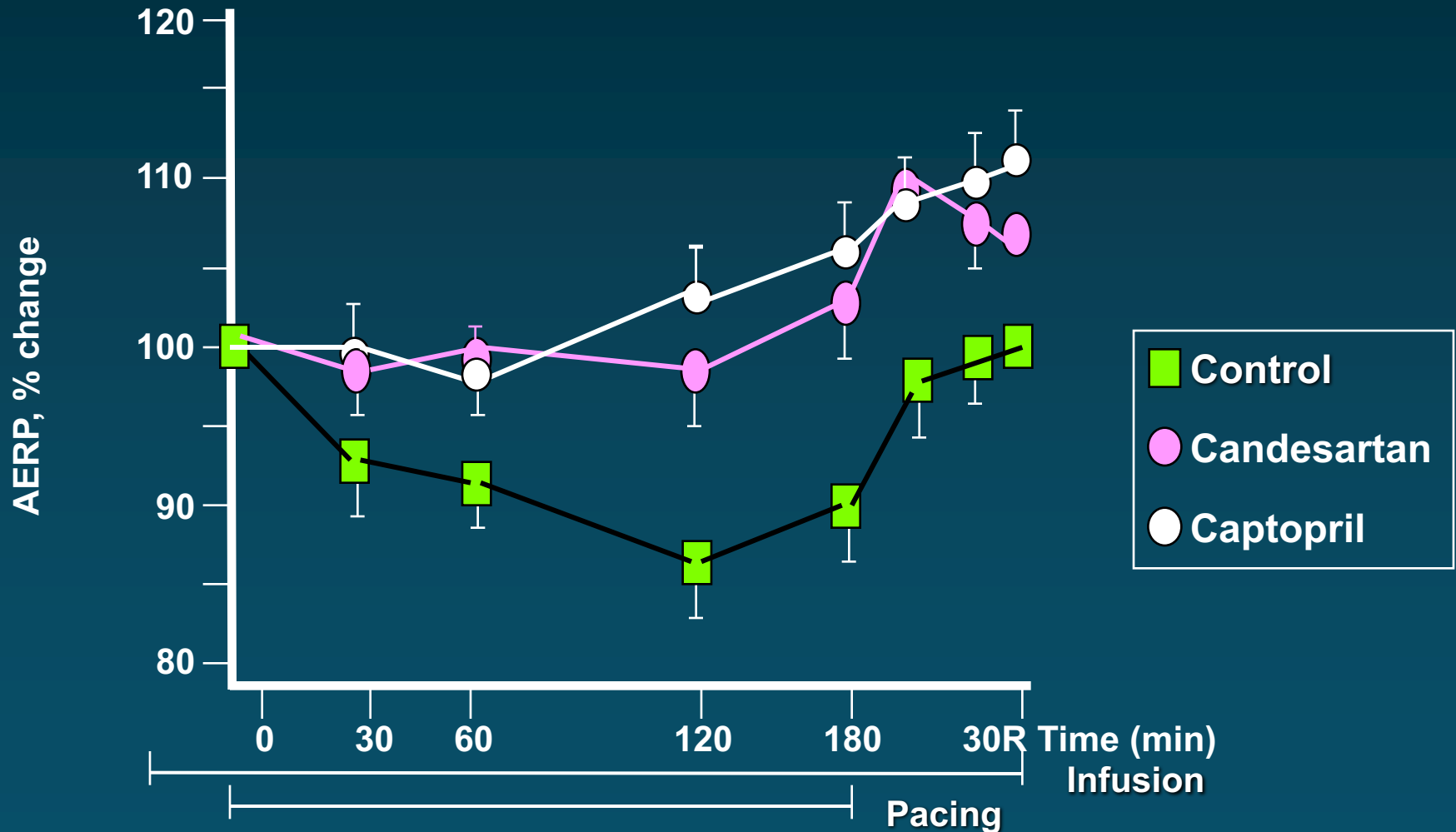
3. Pedersen et al. *Circulation* 1999;100:376-380

4. Tsai CT, et al. *Circulation* 2004;109:1640-1646

# Possible Mechanisms of ARBs in Preventing AF

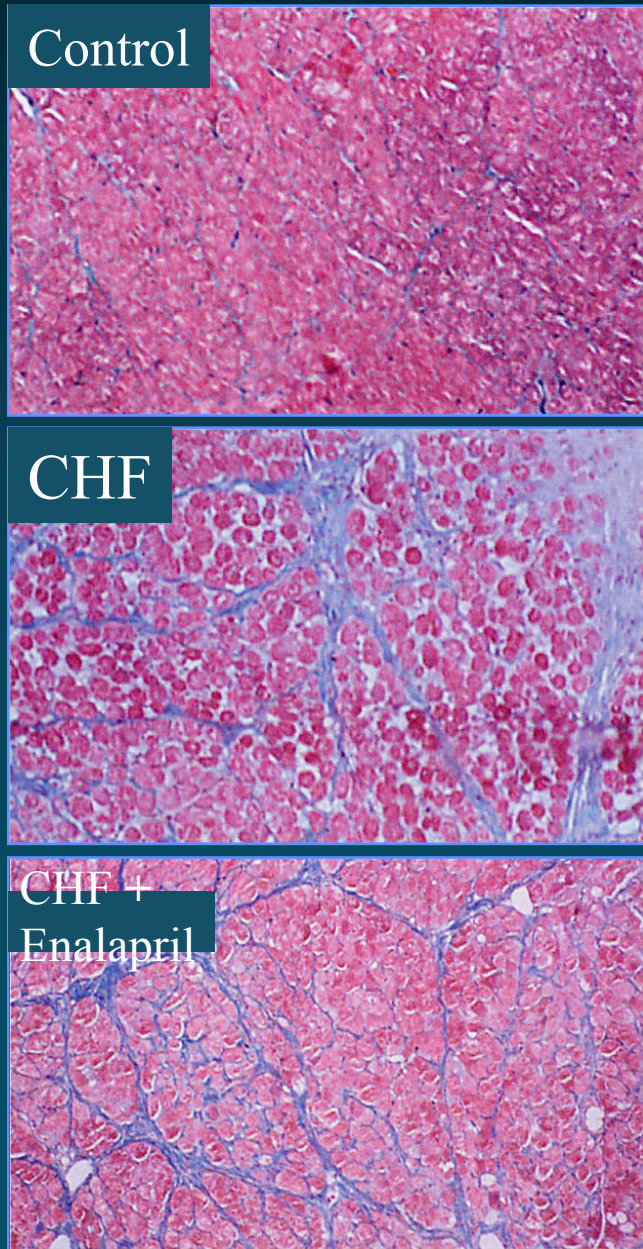


# Angiotensin II Antagonists Attenuate Electrical Remodeling in AF

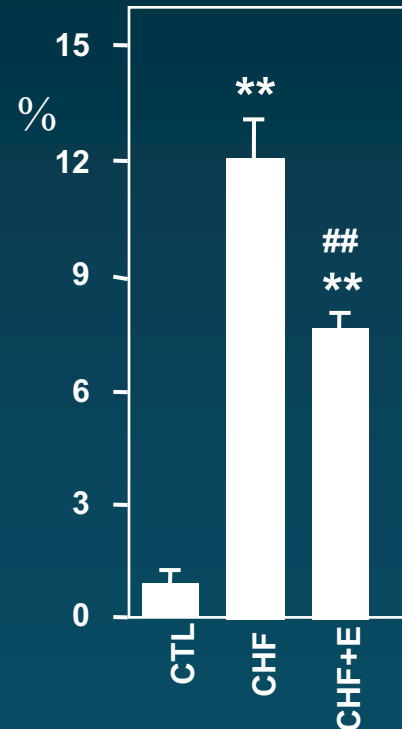




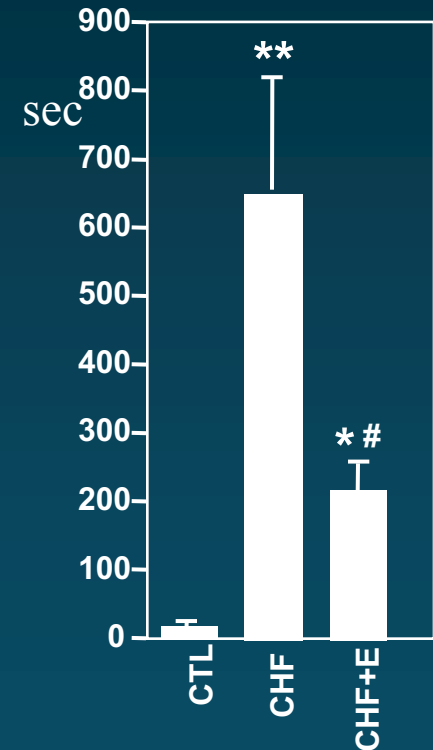
# ACEI: Prevention of Structural Remodeling



## Fibrosis



## AF Duration



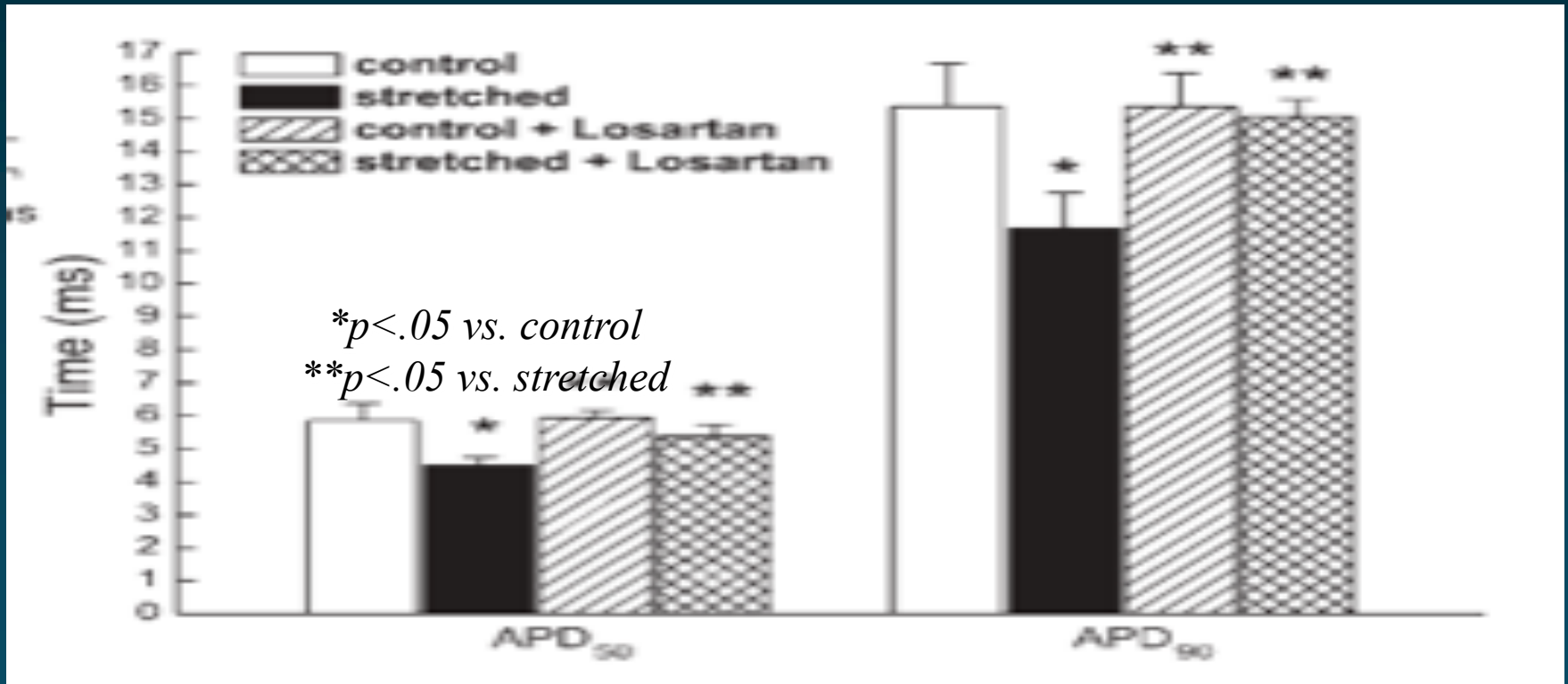
In another study, candesartan decreased duration of AF (4-5 weeks) ( $p < .05$ ) and % atrial fibrosis in atrial paced dog model ( $p < .001$ )

Li D, et al. *Circulation* 1999;100:87-95  
Kumagai K, et al. *JACC* 2003;41:2197-2204

# Does the RAAS Have Direct Electrophysiologic Effects?

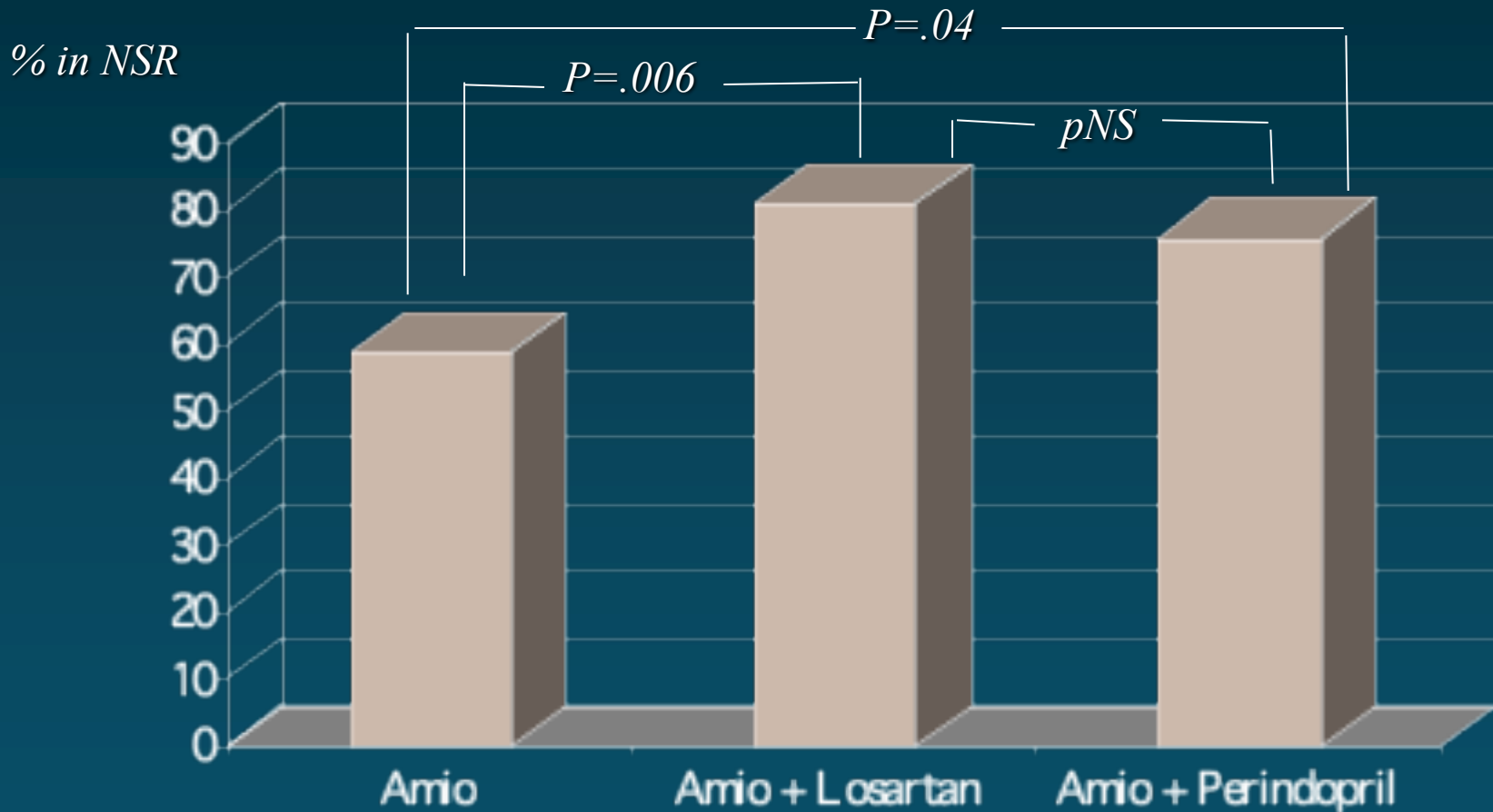
- In a canine RVP model, AF duration increases compared to controls both before and following recovery of pacing-induced atrial electrical remodeling
  - Tissue fibrosis creates a substrate for increased arrhythmia persistence independent of action potential duration
- Although enalapril attenuated the development of interstitial fibrosis, it had no impact on the AERP, conduction velocity or wavelength of conduction
- Angiotensin II may contribute to the development of atrial fibrosis, **but with minimal electrophysiologic effects**

# Losartan Prevents Stretch-Induced Atrial Remodeling *In Cultured Atrial Neonatal Myocytes*



- *Losartan prevented stretch -induced increases in the protein to DNA ratio, ANP mRNA expression*
- *Attenuated stretch-induced expression of IK1, IKur and Ito*
- *Thus preventing the stretch-induced abbreviation of atrial APD*

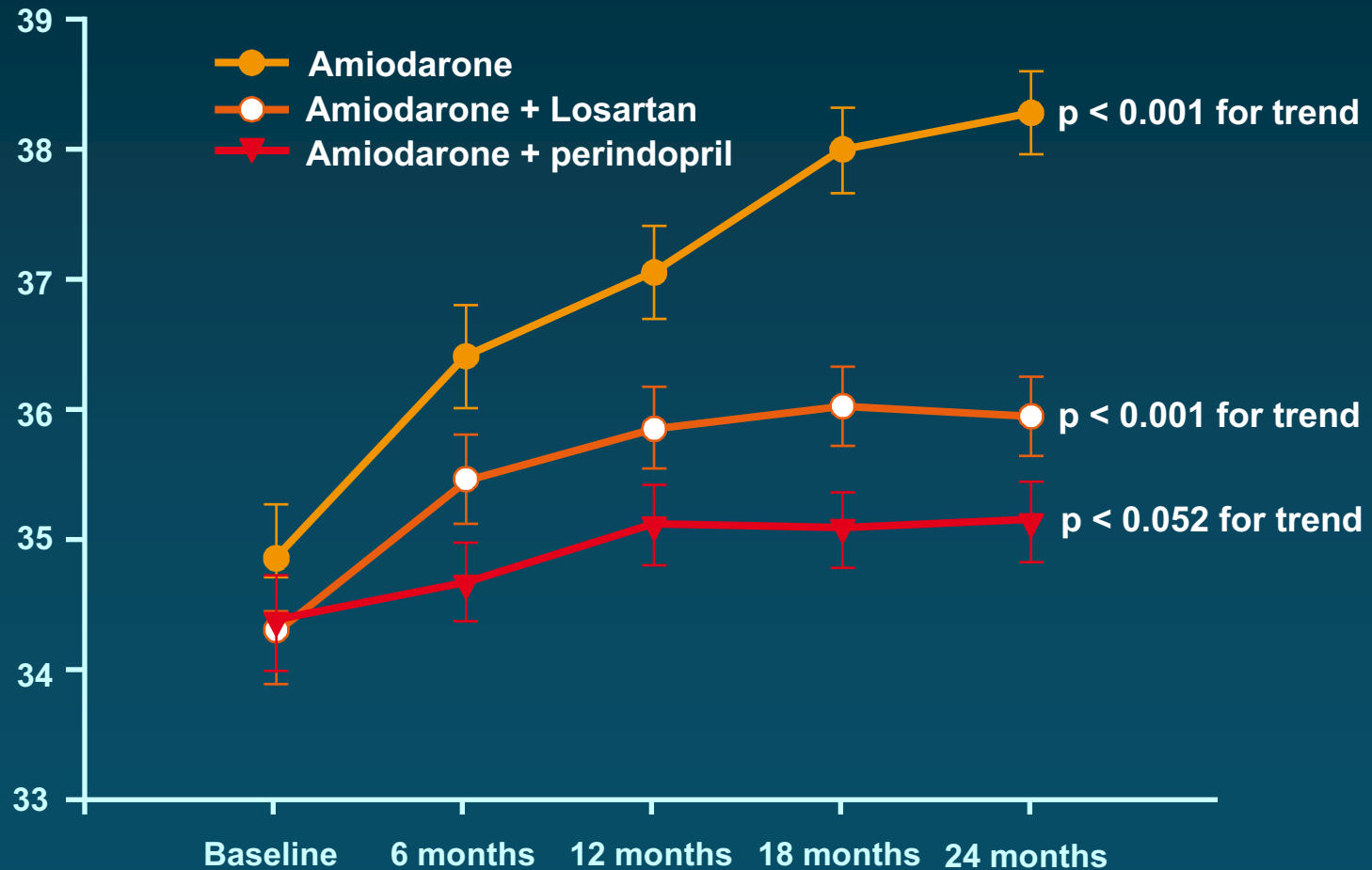
# Amiodarone + Losartan or Perindopril Maintains NSR Post-CV Better Than Amiodarone Alone



*59 pts in each group*

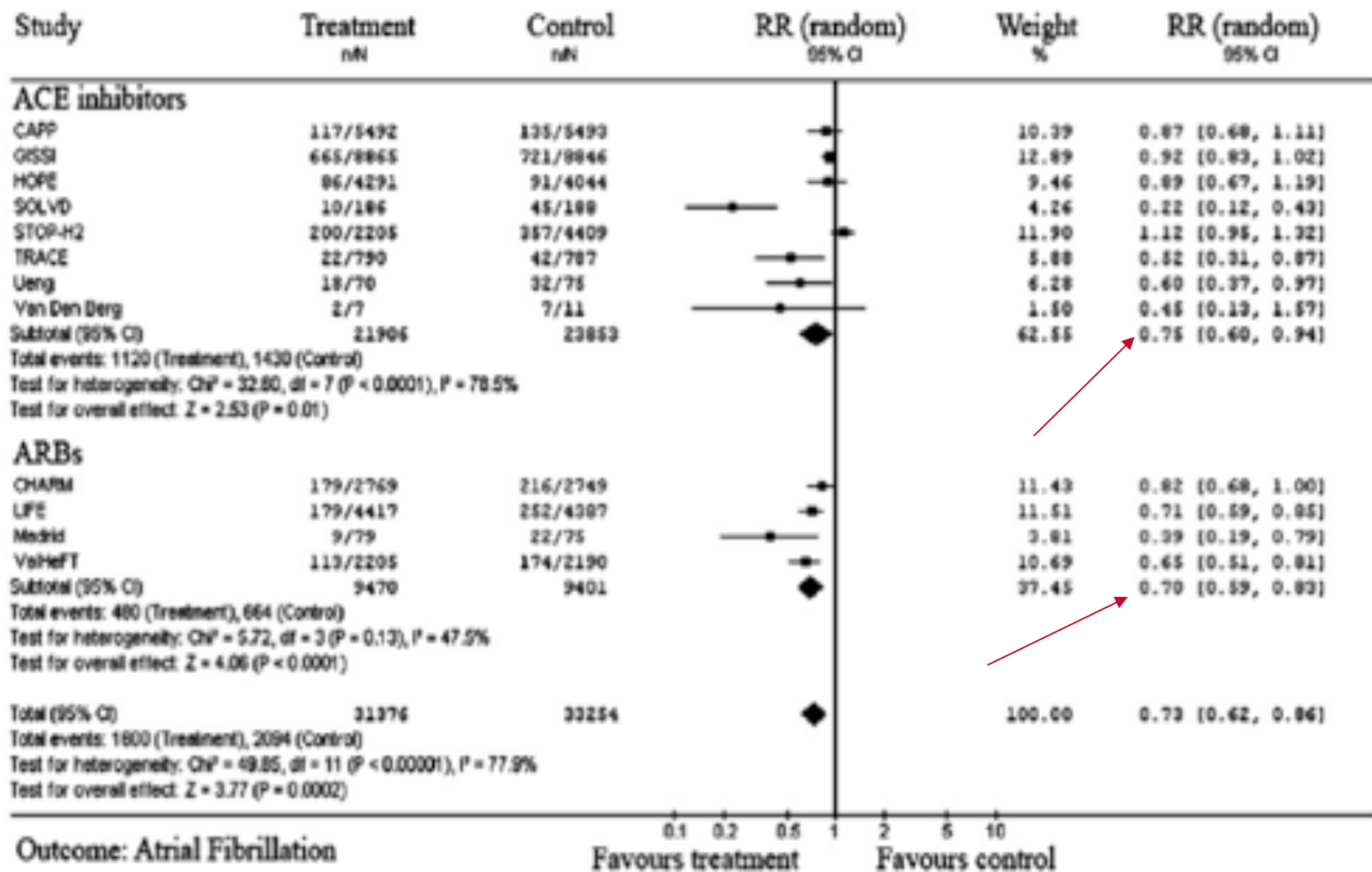
# ACEI and ARB Attenuate Time Dependent Increase in LA Diameter In AF

*LA Diameter in mm*

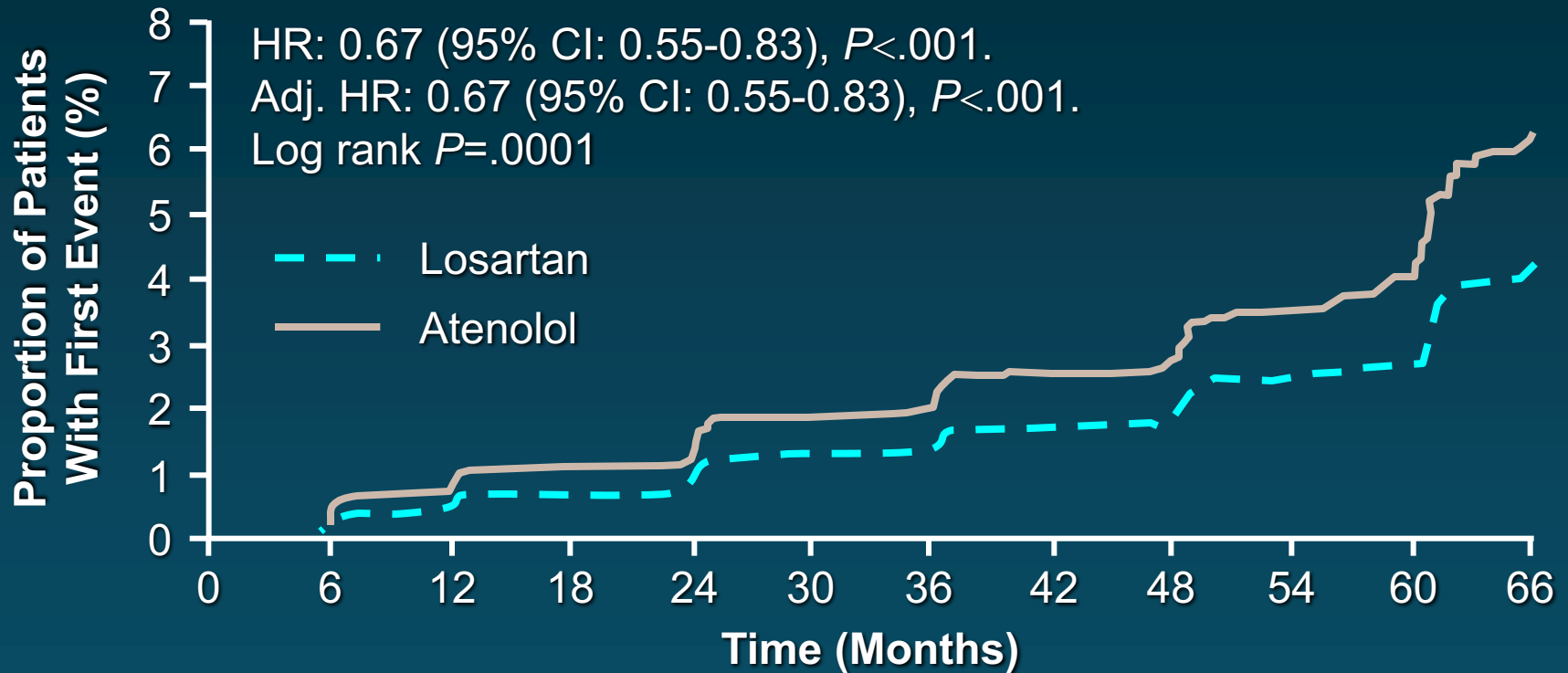




# AF Prevention: ARBs Vs. ACEI?



# Losartan Is Superior to Atenolol in Suppressing AF in the LIFE Study



- Losartan reduced AF compared to atenolol (HR 0.67;  $P < .001$ ) in 342 hypertensive patients with LVH and AF
- Losartan also reduced stroke (HR 0.49; CI 0.29-0.86;  $P = .01$ )

# Which Patient Groups Does RAAS Blockade Prevent AF?

- Hypertension  $\pm$  LVH
  - In **LIFE**, losartan reduced AF compared to atenolol (HR 0.67;  $P < .001$ ) in 342 hypertensive patients with LVH and AF (*Wachtell et al. J Am Coll Cardiol. 2005;45:712-719*)
- Diastolic Dysfunction
  - In CHARM, HR = 0.894 (0.618-1.295) (*Ducharme et al. Am Heart J. 2006;152:86-92*)
- Systolic Dysfunction
  - In **TRACE**, 5.3% of placebo vs. 2.8% of trandolopril group developed AF ( $p < .05$ ) (*Pedersen et al. Circulation 1999;100:376-380*)
  - In **SOLVD**, 5.4% enalapril vs. 24 placebo developed AF ( $p < .0001$ ) (*Vermes et al. Circ 2003; 107:2926-2931*)
  - In **VAL-HeFT**, Valsartan added to ACEI (93%) reduced AF by 35% (7.86% to 5.27%) (*Maggioni AP, et al. Circ 2003;24:504*)
- Diabetes
  - In hypertensive/diabetic patients, valsartan-amlodipine lowered 1 year AF to 14% from 41% with atenolol-amlodipine (*Fogari R, et al. Circulation 2006;114: II)-789*)
- Post-AF ablation
  - No benefit (*Richter et al. Am Heart J 2007;153:113-119*)
- All patients
  - No benefit in HOPE if LVEF  $\geq 40\%$  with only a 2% development of AF overall (HR 0.92, pNS) (*Salehian et al. Am Heart J 2007;154:448-453*)



# Ongoing Trials to Assess ARBs in AF

- **ACTIVE-I** (Irbesartan vs. placebo) - 9018 patients
  - Primary outcome: stroke, MI, vascular death + CHF hospitalization
- **ANTIPAF** (Olmesartan vs. placebo in PAF)
- **CAPRAF** (Candesartan vs. placebo)
- **DRAFT** (Diovan to Reduce post-CV recurrence of AF Trial)
- **GISSI-AF** (Valsartan vs. placebo) – 1442 patients
  - Primary endpoint: Time to 1<sup>st</sup> AF; total # AF episodes
- **I-PACE** (Irbesartan vs. placebo)
- **ON TARGET/TRANSCEND** (Telmisartan vs. ramipril vs telmisartan + ramipril vs. placebo)

**Some of the above studies will tell us if ARBs will prevent AF on top of ACEI**

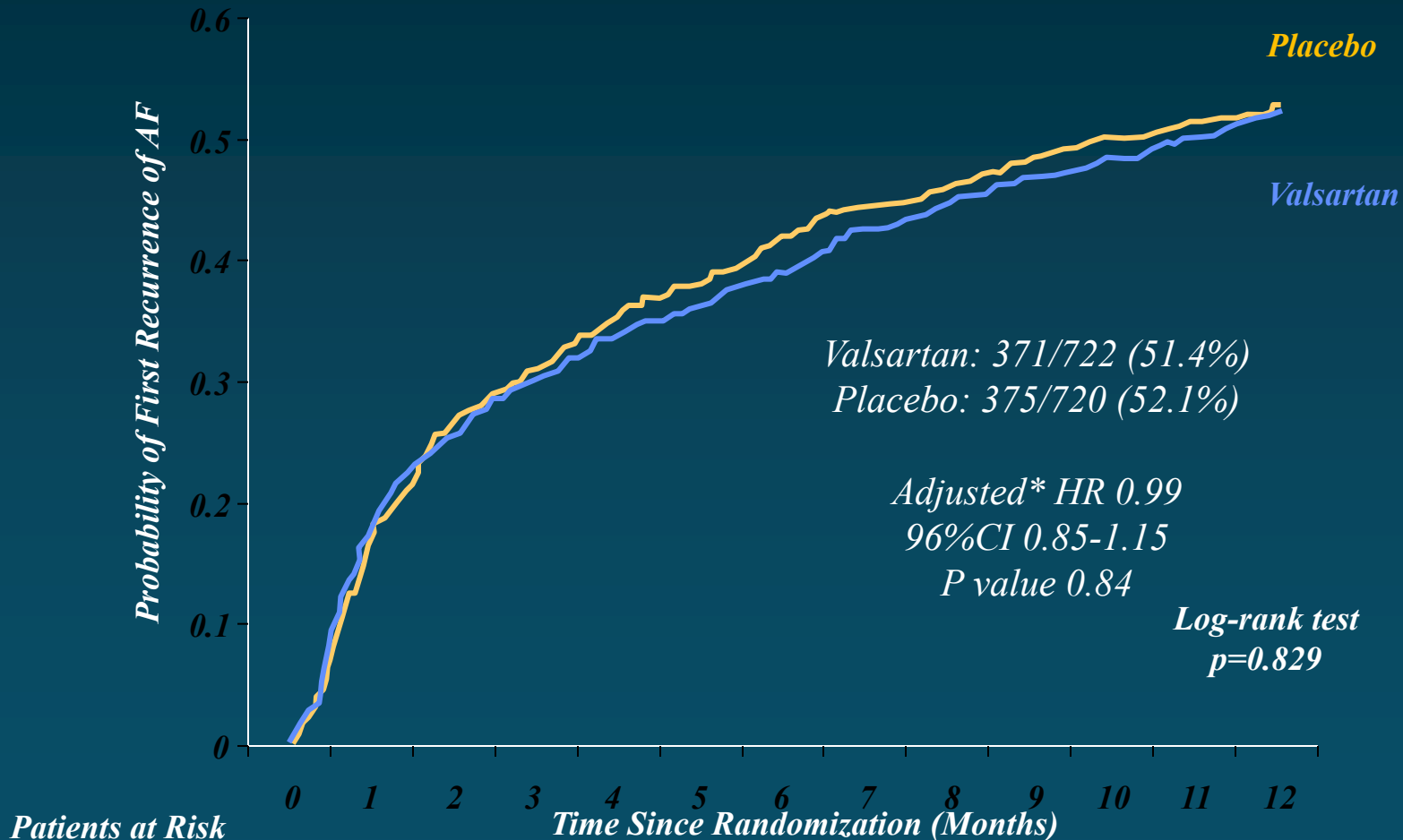
*Connolly S, et al. Am Heart J 2006;151:1187-1193*  
*Disertori m, et al. J Cardiovasc Med 2006;7:29-38*  
*Teo K, et al. Am Heart j 2004;148:52-61*  
*Aksnes TA, et al J Hyperten 2007;25:15-23*

# Preliminary Data ACTIVE-I

- 9018 patients (Irbesartan vs. placebo)
- **Primary outcome: stroke, MI, vascular death + CHF hospitalization**
- 65.2% permanent AF, 20.1 % PAF, 14.5% persistent AF
- 60.4% already on ACEI
- Largest trial of BP lowering in AF
  - 88.7% hypertensive
  - 4-7 mm Hg decrease in SBP over course of study
- Follow-up complete 5/08
  - Mean follow-up 3 years

# GISSI AF

## Study Results: Probability of First Recurrence of AF



# Effects of Aldosterone Antagonists on AF

- Serum aldosterone levels have been reported to be elevated in AF patients with levels returning to normal with restoration of sinus rhythm<sup>1</sup>
- In a rat model of heart failure following MI, spironolactone decreased atrial fibrosis and atrial P-wave duration but ACEI and beta-blockers did not<sup>2</sup>
- In the RVP heart model, eplerenone further prolonged right atrial appendage and left posterior AERP but no effect in Bachmann's bundle
  - No effect of Angiotensin II blockade<sup>3</sup>

1. Goette A et al. *Am J Cardiol* 2001;88:906-909  
2. Milliez P, et al. *Eur Heart J* 2005;26:2193-2199  
3. Schrott SC, et al. *JCVEP* 2006;17:534-541

# Statins and AF

- In a tachycardia induced AF dog model, simvastatin suppressed RAP remodeling effects (shortening of atrial ERP) and induced AF duration (>1000 sec with placebo vs 40 sec with statin)\*
- Tachy-pacing downregulated L-type calcium channel alpha subunit expression was greatly attenuated by simvastatin.
- Retrospective meta-analysis have suggested a beneficial effect in AF in man
- Statin therapy has also been found to reduce the risk for first ventricular arrhythmia in pts with CAD and an ICD.\*\*

\* Shiroshta-Takeshita et al. *Circulation* 2004; 110:2313-19

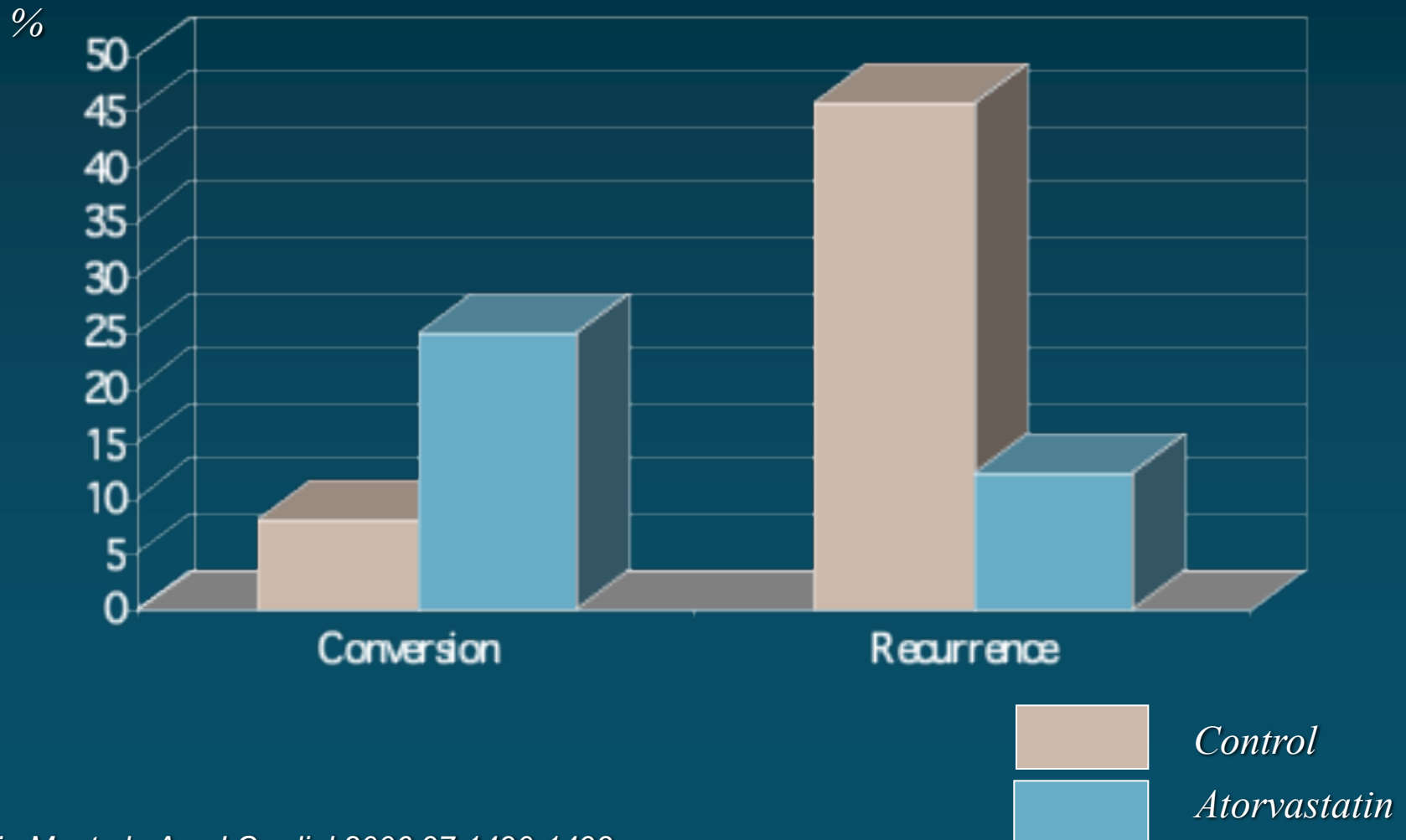
\*\* Chiu et al. *Am J Cardiol* 2005; 95:490-91



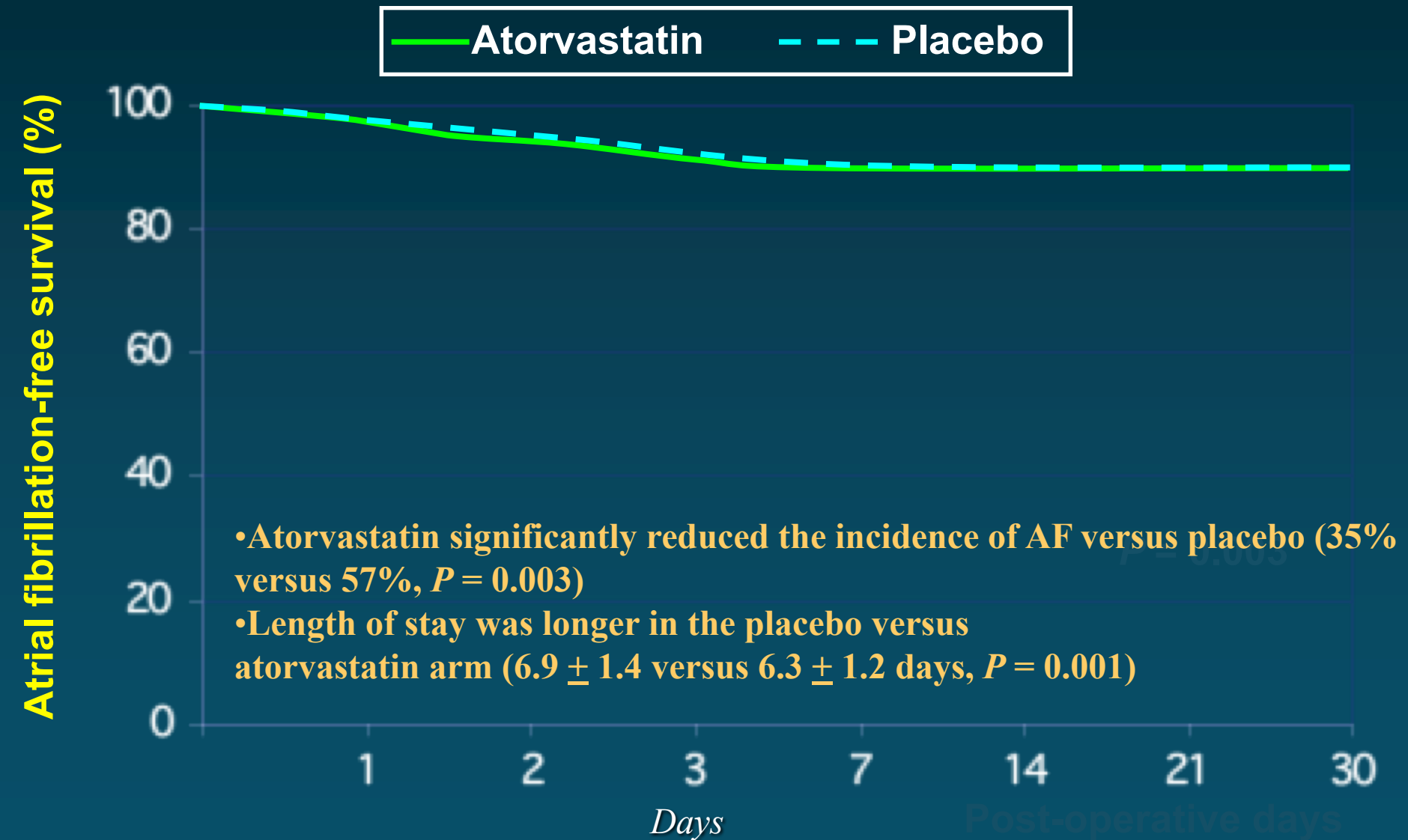
# Statins May Reduce AF in CAD

- 449 pts (age 40-87 yrs) with chronic stable CAD without AF were followed prospectively for an average of 5 yrs in a large out-patient cardiology practice
- The association between statin use and the development of AF was evaluated
- 52 pts (12%) developed AF
- Statins, used by 59% of the subjects, reduced the probability of developing AF (HR 0.49; CI 0.28-0.83;  $p < 0.05$ )
- This remained significant after adjustment for age, HTN, LVEF, CHF, ACU, baseline TC, change in TC (HR 0.37; CI 0.18-0.76)

# Effect of Atorvastatin 10 mg/day in Persistent AF



# ARMYDA-3 (Atorvastatin for Reduction of MYocardial Dysrhythmia After Cardiac Surgery): AF Free Survival





# Statins: AF Recurrence After Cardioversion

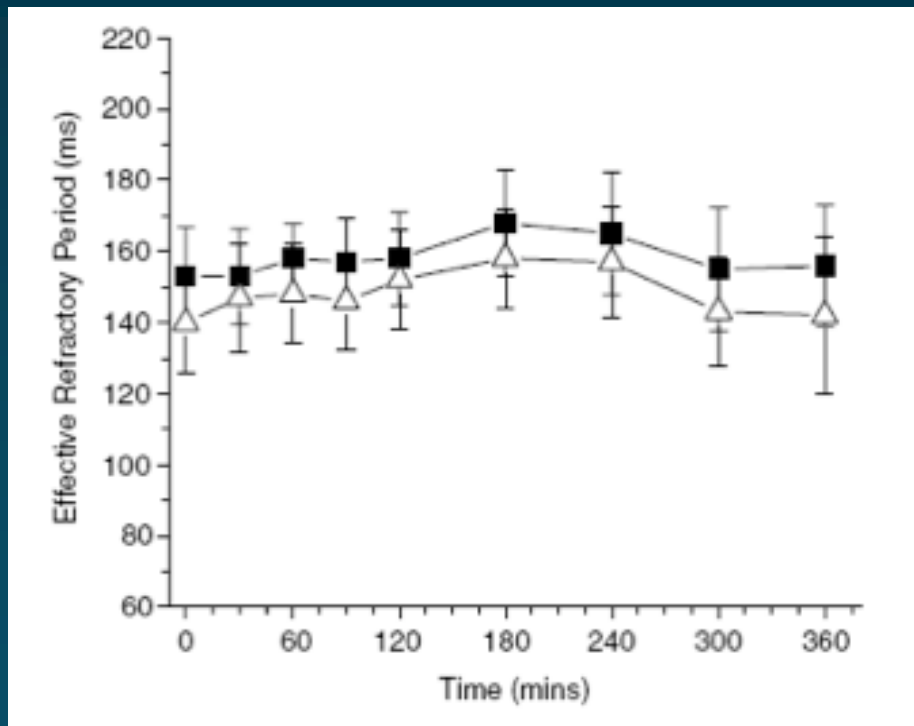
- 62 patients with lone, persistent AF underwent successful DC cardioversion.
- 10/62 were on statin therapy for hyperlipidemia (starting pre-CV). They were also older but had no difference in SHD or AF therapy.
- In a follow up of 44 months on average the use of statins in a retrospective analysis was found to significantly decrease the number with recurrent AF:
  - 40% vs 82%,  $p=0.007$

# Summary of Basic EP Effects of Fish Oil

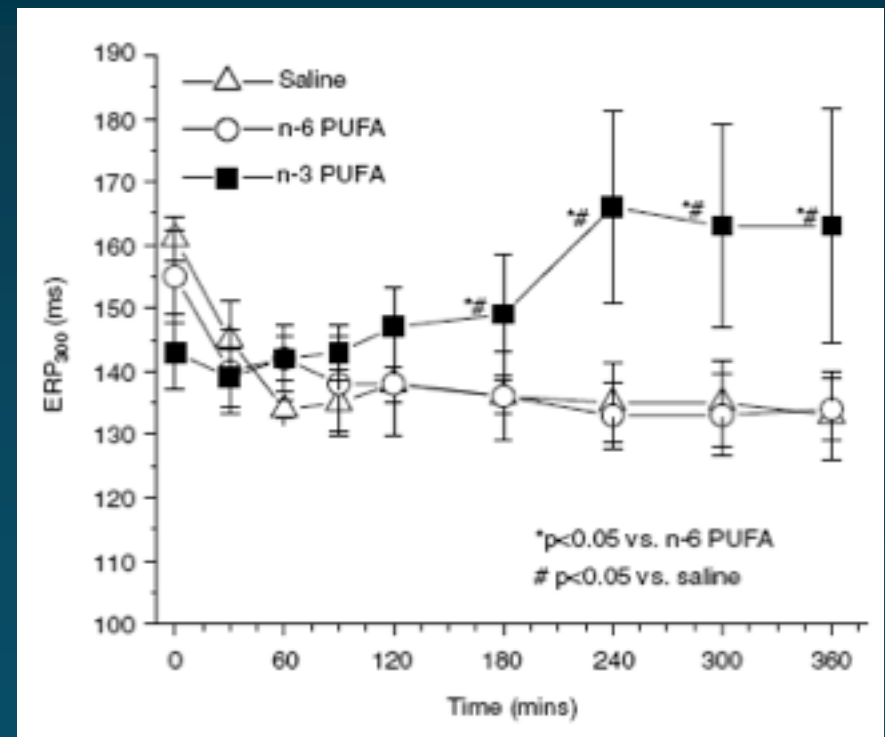
- EPA prolongs the QTc in the Langendorff rabbit model
- Fish oils block L-type calcium channels
- EPA and DHA suppress Na channels in cardiomyocytes and DHA slows Na channel dependent longitudinal conduction in the perfused heart model
- DHA and EPA raise the threshold to elicit an extrasystole
- In humans, fish oils slow heart rate, increase the PR and decrease the likelihood of a prolonged QTc
- Ninio et al noted that feeding rabbits 5% tuna oil for 12 weeks significantly increased the atrial pressure necessary to induce sustained AF compared to controls
  - The decline in AERP produced by increasing atrial pressure was less attenuated in the tuna oil group
- Mozzaferian et al showed that consumption of tuna or other broiled/baked fish lowered AF by 31% if intake was  $\geq 5$  times per week compared to  $< 1$  per month ( $p=.004$ )

# N-3 PUFAs Prevent Acute Electrical Remodeling in Acute Atrial Tachypacing Canine Model

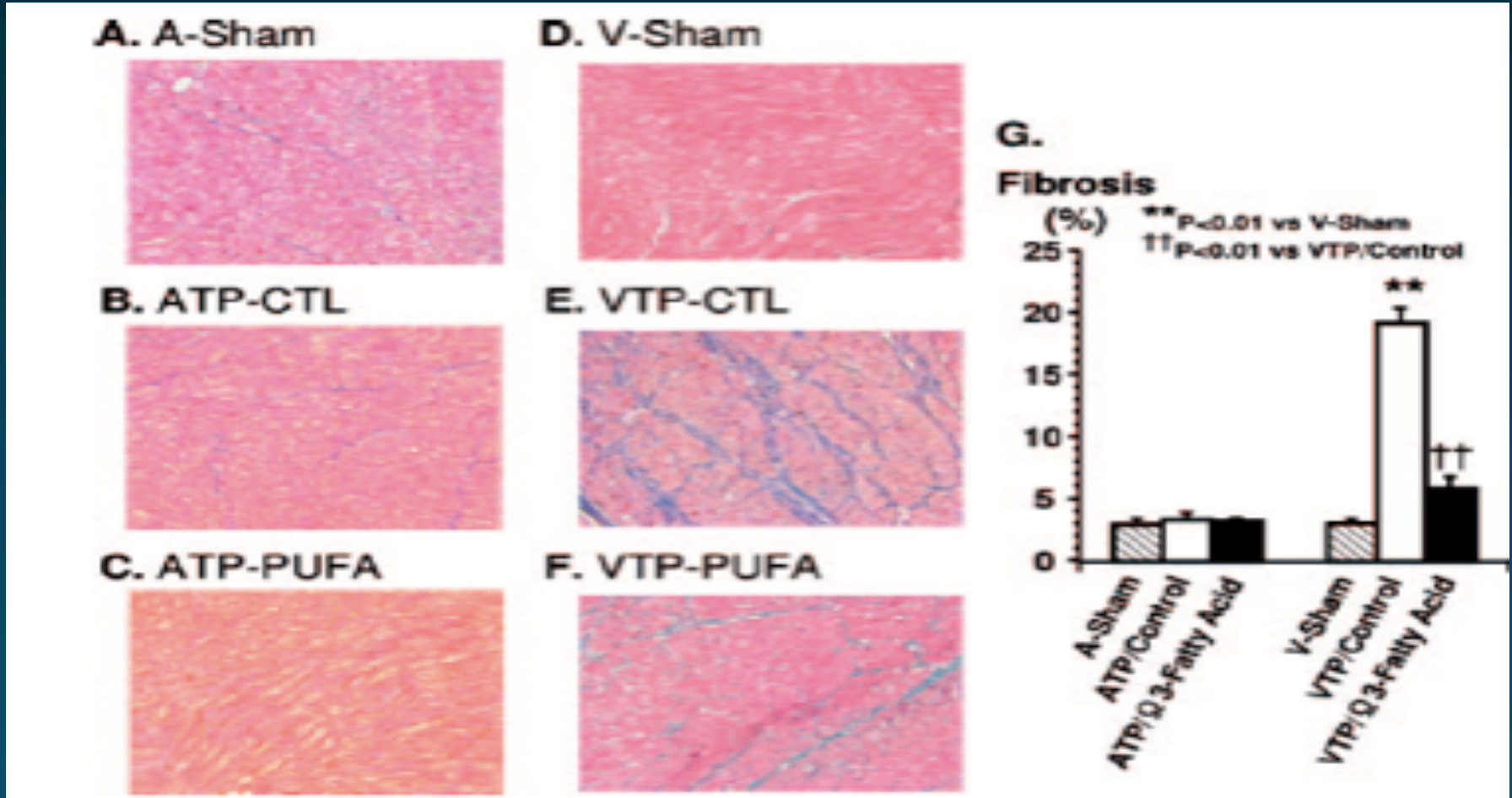
*NSR*



*Atrial Pacing*



# Omega 3-PUFA Prevent AF Associated with Heart Failure But Not Atrial Tachycardia Remodeling



“The beneficial effects of PUFAs on structural remodeling, possibly related to prevention of mitogen-activated protein kinase activation, may contribute to their clinical anti-AF potential.”



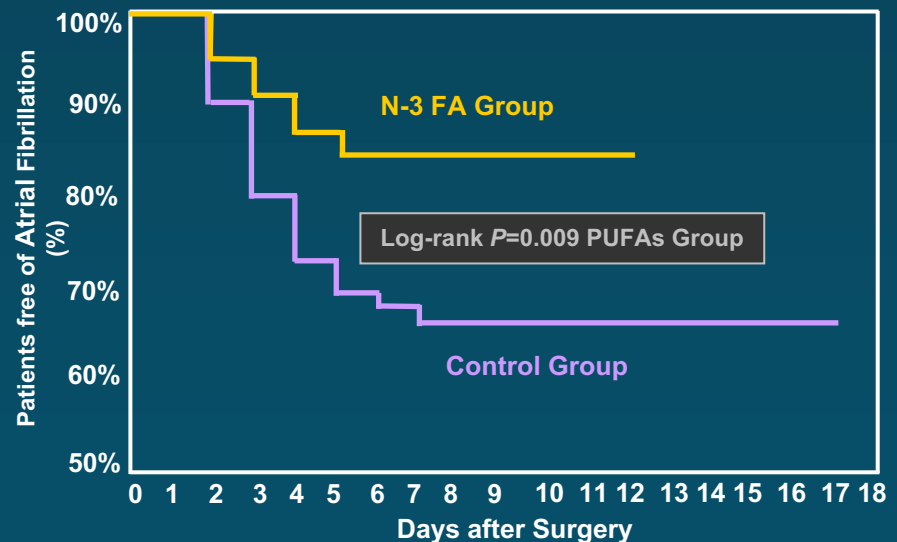
# Reduced Incidence of Vagally Induced AF and Expression of Connexins by n-3-PUFAs in Dogs

- Atrial tissue n-3 PUFA levels increased in oral treatment dogs ( $p < .0001$ )
- Incidence of AF inducibility decreased from 48.9% in controls to 10.5% in treated dogs using the extrastimulus technique ( $p < .003$ )
- Both Cx40 and Cx43 levels (primary components of the atrial GAP junctions) were lower in treated dogs ( $p = .02$ )
- Conclusion: Oral Rx with fish oils increased atrial n-3 PUFA levels and reduced vulnerability to AF induction. Modulation of cardiac connexin expression by n-3 PUFAs may contribute to the antiarrhythmic effect of fish oils

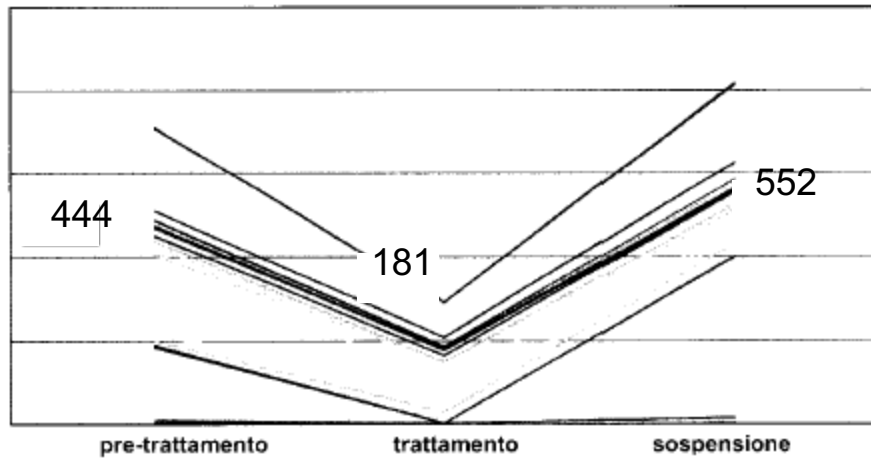
# Omega-3 for Prevention of Atrial Fibrillation Post-CABG

- 160 patients awaiting CABG
- Randomized to usual care or EPA+DHA (1.7 g/d)
- From 5 days pre-surgery through hospitalization
- Endpoint was AF detected by ECG during hospitalization. AF >5 min or requiring intervention

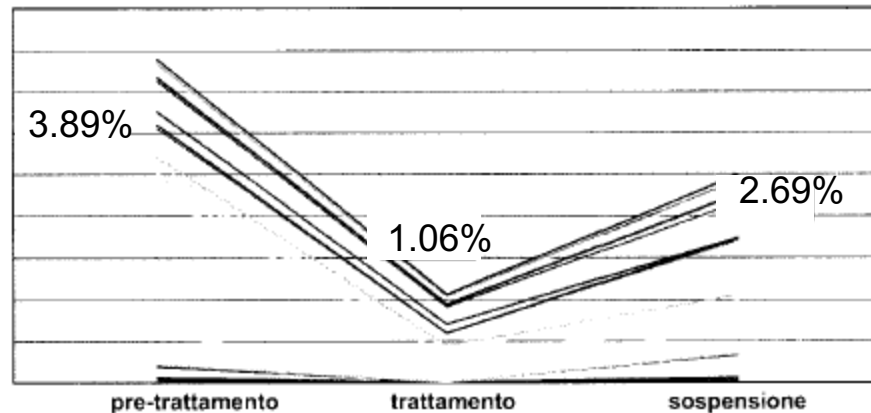
	Control (n=81)	N-3 FA (n=79)	<i>P</i>
Post CABG AF	33%	15%	0.013
Hours of AF	24	16	0.12
Length of Stay	8.2 days	7.3 days	0.017



# Reduction in AF in Patients with Dual Chamber Pacemakers I



Number of atrial tachyarrhythmia episodes



Atrial tachyarrhythmia burden

- N=46 (6 not analyzed) with dual chamber PM
- Design—OLX, received 1gm N-3 or nothing for treatment periods of 4 months
- Results
  - 59% reduction in AFib episodes ( $P=0.037$ ); 67% reduction in AFib burden ( $P=0.029$ )
  - $P=0.065$  and  $0.003$  for increase in AFib episodes and AFib burden following cessation of therapy
  - For patients with sustained AFib there were similar significant reductions in AFib episodes and AFib burden.

# Benefit of Fish Oils in Suppressing Atrial Fibrillation: Possible Mechanisms

- Direct electrophysiologic effects
- Anti-inflammatory effects
- Slow progression of CAD
- Structural
- Metabolic
- Autonomic



# OM8

- Randomized, double blind, placebo controlled, parallel-group trial to assess the efficacy and safety of Lovaza for the prevention of recurrent, symptomatic AF
- **Primary Objective:** assess the effect of Lovaza® on time to the first symptomatic recurrence of AF
  - Time will be measured as event-free days from the end of the loading period (Week 1)
- **Inclusion criteria:**  $\geq 18$  years old, electrocardiographic evidence of symptomatic paroxysmal AF, no current anti-arrhythmic therapy.
  - Rate control and/or anticoagulation therapy or no therapy is permitted.
  - Approximately 550 subjects (275 per treatment group) will be recruited

# What Is The Mechanism Of Down Stream Therapies in Preventing AF?

- **ACEI, ARB** – Angiotensin II blockade; decreasing stretch activated channels; slowing progression of atrial remodeling
- **Statins** – Anti-inflammatory; decrease lipids
- **PUFA** – Anti-inflammatory; decreasing triglycerides; direct channel EP effects

# Conclusion: Upstream Drugs For AF

- Encouraging data for a number of new concepts
- Proof of efficacy/safety will come from well-done, adequately powered, controlled, randomized trials
  - Which patient groups will benefit?
- Added benefits of beta-blockers, ACEI, ARB, statins, Omega-3 fish oil will have to be considered as part of any new drug's efficacy in specific patient populations
- Future possibility of drug combinations with antiarrhythmic agents