

# Curriculum Vitae

## Scott A. Wright, MD, FACC

### Practice

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- 2012-Present      Center for Medical Weight Loss of East Texas, Medical Director  
1761 Troup Hwy, Tyler, Texas
- 2009-Present      Tyler Cardiac and Endovascular Center  
1769 Troup Hwy, Tyler, Texas
- 2003-Present      Cardiovascular Associates of East Texas, P.A., Tyler, Texas

### Affiliation

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- 2003-Present      CAET - Research  
1761 Troup Hwy, Tyler, Texas
- 1/1/13-4/26/13      TAD Clinical Research  
1741 Troup Hwy, Tyler, Texas

### Education

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- 1993-1997      University of Texas Medical Branch at Galveston  
Galveston, Texas  
Doctorate of Medicine
- 1991-1993      University of Texas at Arlington  
Arlington, Texas  
Master of Computer Science
- 1984-1989      Texas A&M University  
College Station, Texas  
Bachelor of Science (Major: Bioengineering)

### Post-Graduate Training

Cardiology Fellowship

- 2000-2003      University of Texas Medical Branch at Galveston  
Galveston, Texas

Internal Medicine Residency

- 1997-2000      University of Texas Medical Branch at Galveston  
Galveston, Texas

## **Board Certification**

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2004 American Board of Internal Medicine – Cardiovascular Disease

2004 Board Certified in Echocardiography

2004 Board Certified in Nuclear Cardiology

## **Licensure**

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2003 Nuclear Radiation Certification State of Texas #L04800

2000 Texas State Board of Medical Examiners #K9950

## **Hospital Appointments**

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Trinity Mother Frances Hospital, Tyler, Texas

East Texas Medical Center, Tyler, Texas

East Texas Medical Center, Athens, Texas

East Texas Medical Center, Carthage, Texas

East Texas Medical Center, Jacksonville, Texas

Titus Regional Medical Center, Mount Pleasant, Texas

Texas Spine and Joint, Tyler, Texas

## **Professional Societies/Memberships**

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1992 Tau Beta Pi Engineering Honors Society

## **Research**

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Principle Investigator – CAMELLIA - A Randomized, Double-blind, Placebo-controlled, Parallel-group Study to Evaluate the Effect of Long-term Treatment with BELVIQ (lorcaserin HCl) on the Incidence of Major Adverse Cardiovascular Events and Conversion to Type 2 Diabetes Mellitus in Obese and Overweight Subjects with Cardiovascular Disease or Multiple Cardiovascular Risk Factors.

Principle Investigator - ODYSSEY - A Randomized, Double-Blind, Placebo-Controlled, Parallel-Group Study to Evaluate the Effect of SAR236553/REGN727 on the Occurrence of Cardiovascular Events in Patients Who Have Recently Experienced an Acute Coronary Syndrome.

Principle Investigator - SUMMIT - HZC113782 A Clinical Outcomes Study to compare the effect of Fluticasone Furoate/Vilanterol Inhalation Powder 100/25mcg with placebo on Survival in Subjects with moderate Chronic Obstructive Pulmonary Disease (COPD) and a history of or at increased risk for cardiovascular disease

Co-Investigator – DECLARE - A Multicenter, Randomized, Double-Blind, Placebo-Controlled Trial to Evaluate the Effect of Dapagliflozin 10mg Once Daily on the Incidence of Cardiovascular Death, Myocardial Infarction or Ischemic Stroke in Patients with Type 2 Diabetes.

Co-Investigator – ABSORB III - Randomized Controlled Trial. A Clinical Evaluation of Absorb™ BVS, the Everolimus Eluting Bioresorbable Vascular Scaffold in the treatment of Subjects with de novo Native coronary Artery Lesions.

Co-Investigator – AD HOC PCI - A randomized, open-label, multiple-center, parallel group, study to compare the platelet inhibition with Verify Now™ assay of ticagrelor vs. clopidogrel in troponin negative Acute Coronary Syndrome (ACS) subjects undergoing Ad Hoc percutaneous coronary intervention (PCI)

Co-Investigator – EXCITE - Excimer laser randomized Controlled study for treatment of femoropopliteal In-Stent Restenosis.

Co-Investigator – EUCLID -A randomized, double-blind, parallel group, multicenter phase IIIb study to compare ticagrelor with clopidogrel treatment on the risk of cardiovascular death, myocardial infarction and ischemic stroke in patients with established Peripheral Artery Disease (EUCLID – Examining Use of tiCagrelor In paD)

Co-Investigator – PARADIGM - A multicenter, randomized, double-blind, parallel group, active-controlled study to evaluate the efficacy and safety of LCZ696 compared to enalapril on morbidity and mortality in patients with chronic heart failure and reduced ejection fraction.

Co-Investigator - XIENCE™ V: Everolimus Eluting Coronary Stent System (EECSS) USA Post-Approval Study sponsored by Abbott Vascular Inc.

Investigator – MUSIC – Multi-Sensor Monitoring in Congestive Heart Failure Study.

Investigator – RECORD AF – **RE**gistry on **C**ardiac rhythm dis**ORD**ers: an international, observational, prospective survey assessing the control of **A**trial **F**ibrillation.

Co-Investigator – RE-LY - Randomized Evaluation of Long term anticoagulant therapy (RE-LY) comparing the efficacy and safety of two blinded doses of dabigatran etexilate with open label warfarin for the prevention of stroke and systemic embolism in patients with non-valvular atrial fibrillation: prospective, multi-centre, parallel-group, non-inferiority trial.

Co-Investigator – ENDEAVOR IV – A Randomized, Controlled Trial of the Medtronic Endeavor Drug Eluting Coronary Stent System versus the Taxus Paclitaxel-Eluting Coronary Stent System in De Novo Native Coronary Artery Lesions.

Co-Investigator – A-HeFT - A Placebo-Controlled Trial of Bidil Added to Standard Therapy In African-American Patients with Heart Failure.

Co-Investigator - ACCOMPLISH- A prospective, multicenter, double-blind, randomized, active-controlled trial to compare the effects of Lotrel (amlodipine/benazepril to benazepril and hydrochlorothiazide combined on the reduction of cardiovascular morbidity and mortality in patients with high risk hypertension.

Co-Investigator - AMIHOT II – A prospective, Multi-center, randomized study of Aqueous Oxygen Therapy for 90 minutes in Anterior Acute MI patients with successful PCI/stenting presenting within 6 hours from time of symptom onset until time of reperfusion.

Co-Investigator – TIMI38/TRITON – A Comparison of CS-747 and Clopidogrel in Acute Coronary Syndrome Patients who are to Undergo Percutaneous Coronary Intervention; a Phase III, multicenter, randomized, parallel-group, double-dummy, active controlled trial in patients with acute coronary syndrome (ACS), who are to undergo percutaneous coronary intervention (PCI).

Co-Investigator – VISION 305 Study - “Vasodilator Induced Stress In Concordance with Adenosine” A multicenter, risk-stratified, randomized, double-blind, double-dummy study drug administration, active-controlled, complete two-arm crossover study with a reference third arm.

Co-Investigator - APEX-AMI Study - “A Multicenter, Randomized, Double-Blind, Parallel-Group, Placebo-Controlled Study of Pexelizumab in Patients with Acute Myocardial Infarction Undergoing Primary Percutaneous Coronary Intervention.” (Assessment of PEXelizumab in Acute Myocardial Infarction)-APEX-AMI.

Co-Investigator - XIENCE™ V: Everolimus Eluting Coronary Stent System (EECSS) USA Post-Approval Study sponsored by Abbott Vascular Inc.

High Frequency Electrocardiograms versus Standard Electrocardiograms in Acute Coronary Syndromes: assessing improved sensitivity of HF-EKG in multiple clinical scenarios including adenosine stress testing, percutaneous interventions, risk stratification in emergency departments. Study done in conjunction with NASA.

Retrospective Analysis of Adenosine Perfusion Studies in Prediction of Coronary Artery Disease: Comparing results of adenosine perfusion studies with documented coronary anatomy by cardiac catheterization.

Masters Thesis, Roger S. Walker, Ph.D. University of Texas at Arlington, Department of Computer Science Developed high resolution laser measurement system for Texas Highway Department Graduate Research Assistant, Roger S. Walker, Ph.D. University of Texas at Arlington, Department of Computer Science Studied feasibility and software modeling of laser systems.

Co-Investigator – SYNERGY - A Prospective, Randomized, Open-Label, Multicenter Study in Patients Presenting with Acute Coronary Syndromes Aventix Pharmaceuticals Protocol Number ENO.GMA.301. IND# 31532

Co-Investigator – A Double-Blind Comparison Of The Incidence Of Hypotension With Two Formulations Of Intravenous Amiodarone: Cordarone• I.V. vs Amiodarone Aqueous I.V. Injection - Protocol No. 058K1-312-US

Co-Investigator – Otsuka Protocol 21-98-214-01 CASTLE “ A Randomized, Double-Blind, Placebo-Controlled, Multicenter, Parallel-Arm, Study to Assess The Long-Term Effects of Pletal® (Cilostazol) Versus Placebo Administered Orally to Patients with Intermittent Claudication Secondary to Peripheral Arterial Disease

Co-Investigator – AMIHOT – TherOx Aqueous Oxygen System – Acute Myocardial Infarction with Hyperoxemic Therapy “AMIHOT” Phase II clinical trial. A Randomized, Controlled, Multicenter Trial of Aqueous Oxygen Infusion for 90 Minutes Post-Primary PTCA/Stent Intervention in Acute Myocardial Infarction Patients.

Co-Investigator – A Multicenter, Randomized, Double-blind, Placebo Controlled Study to Evaluate the Efficacy and Safety of Ad5FGF-4 in Patients with Stable Angina.

Co-Investigator – Extra Point. Protocol # S1710202. Cardiovascular Safety Study of Nicotine Transdermal System.