



Clinical Trials of Texas, Inc.

Dedicated to the advancement of new medical therapies.

Curriculum Vitae

Cesar Reyes, MD

7940 Floyd Curl Drive, Suite 700
San Antonio, Texas 78229

7940 Floyd Curl Drive, Suite 900
San Antonio, Texas 78229

EDUCATION

- 1977-1980 Obstetrics and Gynecology Residence, St. Louis University, St. Louis, MO
1976-1977 General Surgery Residence, PGY II, St. Luke's Hospital, St. Louis, MO
1975-1976 Categorical Internship Surgery, St. Luke's Hospital St. Louis, MO
 • 8 months – General Surgery
 • 4 months – Internal Medicine
1973-1975 Rotating Internship, Clinical Londres, Mexico City
1968-1973 Doctor of Medicine, National University of Mexico School of Medicine, Mexico City

LICENSURE

Texas Medical Board, # G5998

PROFESSIONAL AFFILIATIONS

American Board of Obstetrics and Gynecology
Fellow of the American College of OB-GYN
San Antonio OB/GYN Society
Bexar County Medical Society
Mexican American Physician's Association
Texas Medical Association
The American Fertility Society

RESEARCH AFFILIATION

2006-Present Investigator, Clinical Trials of Texas, Inc., San Antonio, Texas

PROFESSIONAL EXPERIENCE

- 1996-Present Private Practice (Founding Partner), Institute For Women's Health, San Antonio, Texas
1985-1996 Private Practice, Live Oak OB-GYN, San Antonio, Texas
1984 Private Practice, Health America HMO, San Antonio, Texas
1980-1983 Clinical Instructor and Director of the OB/GYN clinics, Medical Education Department
 at Deaconess Hospital, St. Louis, MO

RESEARCH EXPERIENCE

2008-Present: A Multi-Center, Randomized, Controlled Study To Investigate The Safety And Tolerability Of A Single Dose Of "Study Drug" vs. Standard Medical Care In Treating Iron Deficiency Anemia In Subjects Who Are Not Dialysis Dependent

2008-Present: A Multicenter, Randomized, Controlled Study To Investigate The Safety And Tolerability Of "Study Drug" vs. Standard Medical Care In Treating Iron Deficiency Anemia

2008-Present: A Phase III, Three-Arm, Parallel Design, Placebo-Controlled, Randomized, Double-Blind, Multicenter Study Evaluating The Safety And Efficacy Of "Study Drug" In The Treatment Of Premenopausal Women With Symptomatic Uterine Fibroids

2008-Present: Efficacy And Safety Of "Study Drug" In The Treatment Of Moderate To Severe Vaginal Dryness And Vaginal Pain Associated With Sexual Activity, Symptoms Of Vulvar And Vaginal Atrophy (VVA), Associated With Menopause: A 12-Week, Randomized, Double-Blind, Placebo-Controlled, Parallel-Group Study Comparing Oral "Study Drug" 60 MG Daily Dose With Placebo In Postmenopausal Women

2008-Present: A Multi-Center, Placebo Controlled, Safety And Efficacy Study Of "Study Drug" In Anemic, Pre-Menopausal Women With Symptomatic Uterine Fibroids Requiring Hysterectomy

2008-Present: A Multicenter, Randomized, Double-Blind, Parallel Group Study To Evaluate The Efficacy And Safety Of Two Doses Of "Study Drug" Versus Placebo In Women With Overactive Bladder

2007-2008: Long-Term Safety Of 30 MG And 60 MG Oral Daily Doses Of "Study Drug" In The Treatment Of Vulvar And Vaginal Atrophy (VVA) In Postmenopausal Women With An Intact Uterus: A 40-Week Randomized, Double-Blind, Placebo-Controlled, Follow-Up To "Protocol"

2007-Present: A Phase II, Three-Arm, Parallel Design, Dose-Ranging Placebo-Controlled, Randomized, Double-Blind, Multicenter Study Evaluating The Safety And Efficacy Of "Study Drug" In The Treatment Of Premenopausal Women With Symptomatic Endometriosis

2007-Present: A Multi-Center, Randomized, Controlled Study To Investigate The Safety And Tolerability Of "Study Drug" vs. Standard Medical Care In Treating Iron Deficiency Anemia In Heavy Uterine Bleeding And Postpartum Patients

2007-2008: A Randomized, Double-Blind, Placebo-Controlled, Dose-Ranging Multicenter Evaluation Of The Use Of Topically Administered "Study Drug" Versus Placebo In Subjects With Pain Associated With Fibrocystic Breast Disease

2007-Present: An Open Label Study Of The Contraceptive Efficacy Of An Extended Regimen Of "Study Drug" And "Study Drug"

2006-2008: A Randomized, Placebo-Controlled Phase II Study Of Multiple Dosing Regimens Of Intravaginally Administered "Study Drug" Gel For The Treatment Of Cervical High Risk HPV Infection

2006-2008: Efficacy And Safety Of "Study Drug" In The Treatment Of Vulvar And Vaginal Atrophy (VVA) In Postmenopausal Women: A 12-Week, Randomized, Double-Blind, Placebo-Controlled, Parallel-Group Study Comparing Oral "Study Drug" 30 mg And 60 mg Daily Doses With Placebo

 Reyes 1/30/09