

Tamara L. Tulowitzky-Ryan

EDUCATION:

Certified Clinical Research Coordinator (ACRP) 2018

CITI Training 2017

IATA 2014

BLS for Healthcare Providers (CPR & AED) 2017

NIH GCP Certification 2007

Nursing Assistant Program 1996-96

Medical Assistant Program, Bryman College 1995-95

Graduated Desert Winds High School, Lancaster California 1989-1993

EXPERIENCE:

Chesapeake Research Group, CCRC. Night Shift Manager, May 2014-present

Review and become familiar with sponsor protocols. Recruit, consent, and screen subjects. Conduct study visits per protocol. Complete various protocol driven activities to include; pain assessments, VS, phlebotomy, EKG's, study drug administration, and source documentation for subjects enrolled in clinical trials.

Horton Pediatrics and Associates 2009-2014

Provided assistance to the Doctor, triaged patients, vitals, administered vaccinations, phlebotomy, catheterizations, hearing and vision screenings. Responsible for obtaining cultures, urine and blood specimens.

Chesapeake Research Group, CRC. July 2007-2009

Complete various protocol driven activities to include; pain assessments, VS, phlebotomy, EKG's, and source documentation for subjects enrolled in clinical trials.

Chesapeake Foot & Ankle Center, Medical Assistant 2007 to 2009.

Assist doctors in procedures, apply dressings, suture removal, phlebotomy, EKG's, circulate & assist in surgeries, xrays and preparing charts.

Marta Markman, MD & Associates, P.A. 2000-2006.

Provided assistance to the Doctor, triaged patients, vitals, administered vaccinations, phlebotomy, catheterizations, hearing and vision screenings. Responsible for obtaining cultures, urine and blood specimens.

Antelope Valley Foundation For The Developed Mentally Disabled 1997-2000.

One on one instruction, implementing behavior modification with emphasis on positive reinforcement.

CLINICAL TRIAL EXPERIENCE:

Protocol VX18-150-104 (2018)

A Phase 2B Randomized, Double-blind, Placebo-controlled, Dose-ranging, Parallel-design Study of the Efficacy and Safety of (study drug) for Acute Pain Following Bunionectomy

Protocol CA-PS-201 (2018)

A Phase 2, Randomized, Double-blind, Placebo-controlled Safety, Pharmacokinetics and Efficacy Study of (study drug) in Patients Undergoing Bunionectomy

HTX-011-301 (2017)

A Phase 3, Randomized, Double-Blind, Saline Placebo- and Active-Controlled, Multicenter Study of (the study drug) via Local Administration for Postoperative Analgesia and Decreased Opioid Use Following Unilateral Simple Bunionectomy

AVE-901-102 (2017)

A Phase 3, Multicenter, Randomized, Double Blind, Three-Arm Study to Evaluate the Efficacy and Safety of (the study drug) versus Placebo in the Management of Postoperative Pain Following Bunionectomy

CLCT-018 (2017)

A Randomized, Double-Blind, Placebo- and Active-Controlled Study to Determine the Efficacy and Safety of (the study drug) as a Treatment for Moderate-to-Severe Acute Pain and the Prevention of Opioid-Induced Nausea and Vomiting (OINV) Following Orthopedic Surgery

ESTEVE-SUSA-301 (2017)

A Randomized, Double-blind, Active- and Placebo-controlled, Parallel Groups, Phase 3 Clinical Trial to Establish the Efficacy of (the study drug) for the Management of Moderate to Severe Post-surgical Pain after Bunionectomy

AFT-MXIV-07 (2016)

A Phase 3, Randomized, Double-Blind, Multiple-Dose, Parallel-Group and Placebo-Controlled Study of IV (study drug), IV acetaminophen and IV ibuprofen for the Treatment of Acute Postoperative Pain after Bunionectomy

CP130-3002 (2016)

A Phase 3, Multicenter, Randomized, Double-Blind, Placebo- and Active-Controlled Study of (the study drug) for the Treatment of Moderate to Severe Acute Pain After Abdominoplasty

CP130-3001 (2016)

A Phase 3, Multicenter, Randomized, Double-Blind, Placebo- and Active-Controlled Study of (the study drug) for the Treatment of Moderate to Severe Acute Pain After Bunionectomy

HTX-011-C2015-203 (2016)

A Phase 2, Randomized, Controlled Evaluation of the Efficacy and Safety of (the study drug) for Post-Operative Analgesia Following Abdominoplasty Surgery

HTX-011-C2016-208 (2016)

A Phase 2, Randomized, Controlled, Multicenter, Evaluation of the Efficacy and Safety of Locally Administered (study drug) for Postoperative Analgesia Following Bunionectomy

REC-15-016 (2016)

A Phase 3, Multicenter, Randomized, Double-Blind, Placebo-Controlled, Evaluation of the Efficacy and Safety of (the study drug) Following Bunionectomy

INS005-16-062 (2016)

A Phase 3, Randomized, Double Blind, Multiple Dose, Parallel Group, Placebo Controlled Study of (the study drug) for the Treatment of Moderate to Severe Pain

4975-MN-203 (2015)

An Open-Label, Multiple-Dose Extension Study to Evaluate the Safety and Efficacy of (the study drug) in Subjects with Painful Intermetatarsal Neuroma (Morton's Neuroma)

TV46763-CNS-30031 (2015)

A Multicenter, Randomized, Double-blind, Placebo-controlled, Phase 3 Study to Evaluate the Analgesic Efficacy and Safety of (study drug) Every 4 to 6 Hours in Patients with Moderate to Severe Pain Following Bunionectomy

REC-15-014 (2015)

A Phase 2, Single-Center, Randomized, Double-Blind, Placebo-Controlled, Evaluation of the Safety, Efficacy, and Pharmacokinetics of (the study drug) Following Bunionectomy

ELI-200-003-2014 (2015)

A Multi-Center, Randomized, Multiple-Dose, Double-Blind, Placebo-Controlled, Parallel Group, Phase 3 Study to Evaluate the Efficacy and Safety of (the study drug) for the Treatment of Adults with Moderate to Severe Pain Following Bunionectomy Surgery

4975-MN-202 (2014)

A Randomized, Double-Blind, Placebo-Controlled, Dose-Ranging Study to Evaluate the Efficacy and Safety of (the study drug) in Subjects with Painful Intermetatarsal Neuroma (Morton's Neuroma)

ELI-200-003-2014 (2015)

A Multi-Center, Randomized, Multiple-Dose, Double-Blind, Placebo-Controlled, Parallel Group, Phase 3 Study to Evaluate the Efficacy and Safety of (the study drug) for the Treatment of Adults with Moderate to Severe Pain Following Bunionectomy Surgery

INS-14-026 (2015)

A Phase 3, Randomized, Double-Blind, Multiple-Dose, Parallel-Group, Placebo-Controlled Study of (the study drug) for the Treatment of Moderate to Severe Pain

4975-MN-202 (2014)

A Randomized, Double-Blind, Placebo-Controlled, Dose-Ranging Study to Evaluate the Efficacy and Safety of (the study drug) in Subjects with Painful Intermetatarsal Neuroma (Morton's Neuroma)

CP130-2001 (2014)

A Phase 2, Multicenter, Randomized, Double-Blind, Multiple-Dose, Adaptive, Placebo-and Active-controlled Study of (the study drug) for the Treatment of Acute Post-Operative Pain After Bunionectomy

MN-201 (2014)

An Open Label, Ascending Dose Study to Evaluate the Safety and tolerability of (the study drug) in Subjects with Painful Intermetatarsal Neuroma (Morton's Neuroma)

TCO2-2012-01 (2012)

A Prospective, Randomized, Double-Blind Multicenter Study Comparing (the study device) standard Moist Wound Therapy (MWT) in the Treatment of Diabetic Foot Ulcers

R331333PAI3018 (2008)

A Randomized, Double-Blind, Active- and Placebo-Controlled, Parallel-Group, Multicenter Study to Evaluate the Efficacy and Safety of (the study drug) in the Treatment of Acute Pain From Bunionectomy

SKY2028-1-003 (2008)

A Randomized, Double-Blind, Placebo- and Active-Controlled, Parallel Group, 6-Week Study to Evaluate the Effect of Multiple Doses of (the study drug) twice daily, Prednisone and Placebo on the Hypothalamic-Pituitary-Adrenal Axis in Adult Subjects with Mild to Moderate Asthma

Q8003-007 (2007)

A Placebo-Controlled, Randomized, Double-Blind Study of the Safety and Efficacy of Q8003 in the Management of Post-Bunionectomy Pain

KF5503/37 (2007)

A Phase 3, Randomized, Double-Blind, Parallel-Group, Multi-Center, Active and Placebo-Controlled Trial to Evaluate the Analgesic Efficacy and Safety of Multiple Doses of (the study drug) for Postoperative Pain Following Bunionectomy