



THIEVON-WRIGHT

CONSULTING GROUP®

Thievon-Wright Consulting Group

12 New Providence Road, Watchung, NJ 07069

www.cnssites.com

Office: (908) 756-4411 • Fax: (908) 756-1122

Email: info@cnssites.com

# Schizophrenia Case Study 6

**An Open-Label, Randomized, Two Treatment, Multi-Site, Multiple Dose, Steady State, Three-Way, Reference-Replicated Crossover, Pharmacokinetic Study To Determine The In-Vivo Bioequivalence Between XXXXX XXmg Sublingual Tablet And XXXX XXmg Sublingual Tablet**

## Key Elements:

- Rescue Study
- Ultra Rapid Start with high retention

**Jim Aukstuolis, MD**

**Phase:** 1

**Indication:** Schizophrenia (Stable) or Bipolar (stable) subjects

**End Date:** August, 2013

**Rapid Study Start-up:** Contracting, regulatory, and site initiation completed within 48 hours

**# of Subjects:** 11 (study average was 5-6 completed subjects per site)

**In-patient Period:** 6 full bed days with an additional 11.5 days of non-treatment, confinement days per subject

**Completion Rate:** ~91% compared to a study average of 75%

**Other Study Information:** Woodland completed ~26% of study enrollment out of 8 total sites.

## Objectives:

### Primary

Establish the pharmacokinetic bioequivalence between XXXXX sublingual tablets and XXXXX sublingual tablets.  
ECGs—5 per subject

### Secondary

Compare the safety profiles of the test and reference products by examining the adverse events profiles of the two products.

**PK Lab Draws**—9 per subject

Day 5, Day 6, Day 12, Day 13, Day 19 and Day 20. PK draws performed on days 7, 14 and 21 at hour 0 (pre-dose), 0.25, 0.5, 1.0, 1.5, 2.0, 2.5, 3.0, 3.5, 4.0, 5.0, 6.0, 8.0, 10.0, 12.0. All pre-dose draws were performed within 5 minutes prior to dosing.

**ECGs**—5 per subject

Day -7

Day 1

Day 8

Day 15

Day 24

- Standard 12-lead ECGs

**Vitals**—14 per subject

Daily

- Assessed within 15 minutes before and within 30 minutes after the morning dose

**Hematology / Chemistry**—6 per subject

Day -7

Day 1

Day 7

Day 14

Day 21

Day 24