

Background

Methods

after a single tablet of TNX-102 SL 2.8 mg or CBP IR 5 mg,

Results

TNX-102 SL is a eutectic CBP formulation which contains

potassium phosphate dibasic as a basifying agent that

in an un-ionized state at the mucosal membrane, thus rapidly

driving CBP across the mucosa into the bloodstream. For

TNX-102 SL 2.8 mg v. ingested CBP IR 5 mg, plasma CBP levels were: at 10 min 338 pg/ml v. below limit of detection

(BLD); at 20 min 739 pg/mL v. BLD; at 30 min 988 pg/mL v.

BLD; at 45 min 1209 v. 280 pg/mL (p=0.001); at 60 min 1545

v. 913 pg/mL (p=0.062); and at 120 min 2296 v. 1737 pg/mL

(p=0.043). For TNX-102 SL 2.8 mg v. CBP IR 5 mg tablets. the mean exposure was 338% (p=0.009) higher at 1h, and

83% (p=0.034) higher at 2h. TNX-102 SL 2.8 mg had Cmay = 3.4 ng/mL and AUC₀₋₈ = 79 ng hr/mL while CBP IR 5 mg had $C_{max} = 4.3$ ng/mL and AUC₀₋₈= 92 ng hr/mL showing more

efficient dose-adjusted absorption for TNX-102 SL. The

plasma levels of norcyclobenzaprine (nCBP), the major

metabolite of CBP, were lower with TNX-102 SL consistent

with bypassing first pass hepatic metabolism. TNX-102 SL was well tolerated and side effects were similar to those of

oral CBP although some subjects experienced numbness in

Conclusions

TNX-102 SL delivers CBP rapidly across the sublingual

mucosal membrane into plasma resulting in 12 times faster

onset of absorption relative to oral CBP IR, and provides

hours. The relative bioavailability was 154% when compared to the CBP IR tablet. The SL formulation had no effect on

Tmax. Sublingual administration of CBP via TNX-102 SL bypasses "first-pass" metabolism reducing Cmax and AUC to

nCBP, the active metabolite. The pharmacokinetic properties

of TNX-102 SL appear to be well suited for its development

as a potential bedtime medication for FM in a long-term

the mouth that was transient and self-limited.

transmucosal absorption.

and PK parameters were calculated.

Rapid Sublingual Absorption of Cyclobenzaprine (CBP) with Basifying Agents: Prospect for Bedtime Treatment of Fibromyalgia Syndrome (FM)

Bruce Daugherty, Nunzia Ceppi Monti, Valentina Panzeri, Roberto Marelli, Enrico Magnocavallo, Giorgio Reiner and Seth Lederman Tonix Pharmaceuticals, Inc., New York, NY and APR Applied Pharma Research s.a., Balerna, Switzerland

0-2 hr

Time after Dose (hours)

2.0

1.0

1.5

0.0

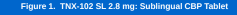
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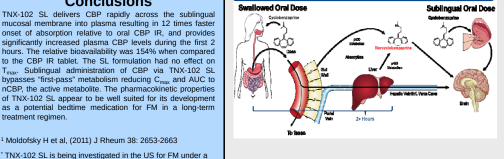
Funded by Tonix Pharmaceuticals. Inc.

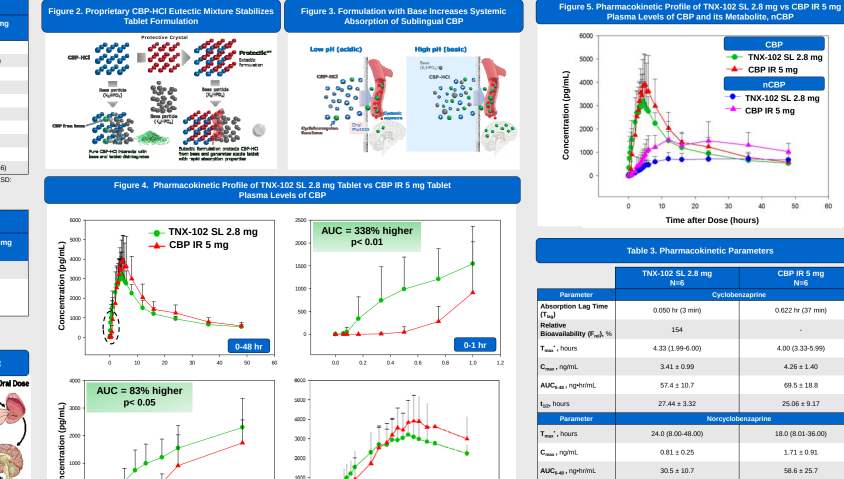
Table 1. Demographic Data Cyclobenzaprine (CBP) exposure during sleep improves TNX-102 SL 2.8 mg CBP IR 5 mg daytime fibromyalgia (FM) symptoms and sleep guality¹. CBP Category N=6 N=6 absorption into plasma is delayed after ingesting immediate release (IR) tablets. To speed absorption, TNX-102 SL*, a Age (years) sublingual (SL) formulation of 2.8 mg CBP was developed for Mean (SD 36.7 (15.0) 37.3 (15.4) Gender, N (%) Female 4 (66.7%) 3 (50.0%) Plasma CBP was measured in healthy subjects (N=6/group) Male 2 (33.3%) 3 (50.0%) Ethnicity, N (%) Not Hispanic 5 (83.3%) 5 (83.3%) Hispanic 1 (16.7%) 1 (16.7%) BMI (kg/m²) disintegrates in saliva and rapidly dissolves. The addition of a Mean (SD) 26,160 (2.821) 25.873 (2.456) basifying agent results in a higher pH, thereby rendering CBP

Abbreviations: SL: sublingual; IR: immediate release; CBP: cyclobenzaprine; SD: standard deviation; BMI: body-mass index

Table 2. Treatment-Emergent Adverse Events			
System Organ Class/ Preferred Term	TNX-102 SL 2.8 mg N (%)	CBP IR 5 mg N (%)	
Gastrointestinal disorders	3 (50.0)	0	
Hypoaesthesia oral	2 (33.3)	0	
Oral mucosal erythema	1 (16.7)	0	
Abbreviations: SL: sublingual; IR: immediate-release; CBP: cyclobenzaprine			







0-8 hr

Tiles is defined at the first nominal sampling time after administration from which onward the CBP concentrations consistently exceed the limit of quantitation; Relative Bioavailability (Free) was calculated using the formula, Frei= 100 x [Dose (IR) x AUC (SL)/Dose (SL) x AUC (IR)]; Mean ± SD; *Median (Min-

Presentation Number LB-026

CBP

nCBP

TNX-102 SL 2.8 mg CBP IR 5 mg

Time after Dose (hours)

Table 3. Pharmacokinetic Parameters			
	TNX-102 SL 2.8 mg N=6	CBP IR 5 mg N=6	
Parameter	Cyclobenzaprine		
bsorption Lag Time T _{lag})	0.050 hr (3 min)	0.622 hr (37 min)	
elative lioavailability (F _{rel}), %	154	-	
max*, hours	4.33 (1.99-6.00)	4.00 (3.33-5.99)	
m _{ax} , ng/mL	3.41 ± 0.99	4.26 ± 1.40	
UC ₀₋₄₈ , ng•hr/mL	57.4 ± 10.7	69.5 ± 18.8	
_{//2} , hours	27.44 ± 3.32	25.06 ± 9.17	
Parameter	Norcyclobenzaprine		
max*, hours	24.0 (8.00-48.00)	18.0 (8.01-36.00)	
m _{ax} , ng/mL	0.81 ± 0.25	1.71 ± 0.91	
UC ₀₋₄₈ , ng•hr/mL	30.5 ± 10.7	58.6 ± 25.7	
_{//2} , hours	71.95 ± 30.97	66.70 ± 35.11	

treatment regimen

1 Moldofsky H et al. (2011) J Rheum 38: 2653-2663

* TNX-102 SL is being investigated in the US for FM under a US IND and is not approved for any indication