SYNOPSIS

NAME OF SPONSOR/COMPANY:	INDIVIDUAL STUDY TABLE	(FOR NATIONAL
The R.W. Johnson Pharmaceutical Research	REFERRING TO PART	AUTHORITY USE ONLY)
Institute	OF THE DOSSIER	
NAME OF FINISHED PRODUCT:	@@7000fad8004b3c3	
TOPAMAX [®] (topiramate) tablet		
NAME OF ACTIVE INGREDIENT(S):		
$2,3:4,5$ -bis- O -(1-methylethylidene)- β -D-		
fructopyranose sulfamate		
Protocol No.: CR005455		·
Title of Study: Topiramate (RWJ-17021-00	0) Clinical Trial in Primary Generalized	Tonic-Clonic Seizures
Investigators: 18 investigators		
Study Centres: 18 study centers		
Publication (Reference): None		
Studied Period (years): 5 May 1994 - 5 July	1996	Phase of development: 3
Objectives: This trial was designed to evalu	ate the safety and efficacy of oral to	piramate as adjunctive therapy in
subjects with uncontrolled primary generaliz	red tonic-clonic (PGTC) seizures (i.e.	, tonic-clonic seizures considered
to be generalized from the onset) with or wit	hout other generalized seizure subtype	es.
Methodology: This was a multicenter, rand	omized, double-blind, placebo-control	led trial that evaluated topiramate
total daily (target) dosages of 175, 225, or	400 mg/day based on subject's wei	ght to approximate 6 mg/kg/day
(theoretical range: ≤9.3 mg/kg/day) as adju	nctive therapy in subjects with PGT	C seizures with or without other
generalized seizure subtypes. The trial inclu	ded a baseline phase (approximately 5	6 days in duration) and a double-
blind phase (approximately 140 days in dura	tion). During the baseline phase, the	number and type of seizures that
occurred were monitored while subjects recei	ved a constant dosage of one or two a	ntiepileptic drugs (AEDs). Those
subjects who were eligible for the double-blin	d phase of the trial were randomized in	equal proportions at each center to
receive either placebo or topiramate while c	ontinuing on their background AED	regimen. Efficacy was evaluated
based on the reduction from baseline in avera	ge monthly PGIC seizure rate, the pri	mary efficacy variable, and on the
reduction from baseline in average monthly seizure rate based on all seizures. Efficacy was also evaluated by the		
percent of treatment responders (subjects with a \geq 50% reduction in average monthly seizure rate) based on PGTC asize and based on PGTC.		
seizures and based on all seizures, and subject's global evaluation of improvement in seizure severity. Safety was		
electrocardiograms (ECGs) physical examin	ations and neurologic examinations	Subjects (or their parents/legal
guardians) completed evaluations on ment	ations, and neurologic examinations.	level of interaction with their
environment ability to perform activities of	daily living and responsiveness to ver	bal requests In addition plasma
AED concentrations were measured at periodic intervals to assess potential effects of topiramate on background AEDs		
Number of Subjects (planned and analyzed): Eighty subjects were enrolled: 41 were randomly assigned to receive		
placebo and 39 were randomly assigned to	receive topiramate. All 80 subjects w	vere included in the intent-to-treat
analyses of efficacy and safety.	i i i i i i i i i i i i i i i i i i i	
Diagnosis and Main Criteria for Inclusion	: Subjects (≥ 4 years of age; ≥ 25 kg	g) enrolled in the trial had PGTC
seizures with or without other generalized seiz	ure subtypes. Subjects were to have th	ree or more PGTC seizures during
the 56-day baseline phase (with at least one du	ring each 28-day period) while on a sta	ble regimen of one or two AEDs.
Test Product, Dose and Mode of Administr	ation, Batch No.: Topiramate was sur	oplied as white 25 mg (R5489) and
vellow 100 mg (R5509) tablets. Maximum dosages of topiramate based on subjects' weight were 175 mg/day (25 to		
33.9 kg), 225 mg/day (34 to 42.9 kg), 400 mg/	day (≥43 kg).	
Duration of Treatment : The total duration of	of double-blind therapy was 140 days,	including a 56-day titration period
and an 84-day stabilization period.		
Reference Therapy, Dose and Mode of Administration, Batch No.:		
Placebo was supplied as "25 mg" (R5721) and	"100 mg" (R4567) tablets to match top	piramate tablets.
Criteria for Evaluation:		
Efficacy: The efficacy of topiramate in the treatment of PGTC seizures was based on a statistically significant		
between-group difference with respect to per	cent reduction in average monthly PG	TC seizure rate.
Safety: Safety was evaluated by reported	adverse events, clinical laboratory to	ests, vital sign and body weight

measurements, ECGs, physical and neurologic examinations, and evaluations of the subject's mental status.

SYNOPSIS (Continued)

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fructopyranose sulfamate		

Statistical Methods: A two-way (with treatment and center as factors) analysis of variance on ranks was used to analyze treatment group differences in percent reduction from baseline seizure rate for both PGTC seizures and all seizures. An additional efficacy assessment compared treatment groups with respect to percent of PGTC responders and responders based on all seizures, stratified by center, using the Cochran-Mantel-Haenszel method. The global evaluation of improvement in seizure severity was analyzed by exact Wilcoxon rank-sum tests stratified by center and unstratified.

SUMMARY - CONCLUSIONS

<u>EFFICACY RESULTS</u>: Topiramate was statistically superior to placebo with respect to percent reduction in average monthly seizure rate and percent treatment responders for both PGTC seizures and all seizures combined (Table 1). The percent reduction from baseline in average monthly seizure rate numerically favored topiramate over placebo for absence (53% vs. 4%), myoclonic (52% vs. an increase of 401%), and tonic (16% vs. an increase of 1%) seizures.

Table 1:	Summary of the Efficacy Results for the Double-Blind Phase
	(All Randomized Subjects: Protocol CR005455)

Efficacy Assessment	Placebo	Topiramate	p-value
Primary Variable Percent reduction from baseline in average monthly seizure rate for PGTC seizures	9.0	56.7	0.019 ^b
Secondary Variables Percent reduction from baseline in average monthly seizure rate for all seizures	0.9	42.1	0.003 ^b
Percent treatment responders ^a : PGTC seizures All seizures	20 17	56 46	0.001 ^c 0.003 ^c
Subject's global evaluation of improvement in seizure severity ^d	56	62	$0.490^{\rm e}$ $0.388^{\rm f}$

^a A treatment responder is defined as subject whose seizure rate was reduced 50% or more during the double-blind phase.

^b Topiramate vs. placebo; two factor (treatment and center) ANOVA on ranks.

^c Topiramate vs. placebo; Cochran-Mantel-Haenszel test.

^d Percent of subjects who had minimal, moderate, or marked improvement in seizure severity.

^e Topiramate vs. placebo; Wilcoxon rank-sum test stratified by center.

^f Topiramate vs. placebo; Wilcoxon rank-sum test unstratified.

When treatment responder was defined more rigorously as \geq 75% seizure rate reduction for both PGTC and all seizures, the difference between topiramate (33% and 26%) and placebo (13% and 7%) was statistically significant (p \leq 0.037). The percentage of subjects who were seizure-free numerically favored topiramate (13% and 5%) over placebo (5% and 0%) for both PGTC seizures and all seizures, respectively. Sixty-two percent and 56% of topiramate and placebo subjects, respectively, reported improvement in seizure severity; the between-group difference in improvement in seizure severity was not statistically significant (p=0.490). Because plasma concentrations of concomitant AEDs were generally comparable over time between topiramate- and placebo-treated subjects, the topiramate effects observed in this study were not mediated through changes in plasma concentrations of concomitant AEDs. There was no consistent relationship between efficacy and plasma topiramate concentration.

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<u>SAFETY RESULTS</u>: Seasonal illnesses and associated symptoms were the most commonly reported treatmentemergent adverse events in both treatment groups in this trial. Neuropsychiatric adverse events were reported in both topiramate and placebo treatment groups in this trial. Anorexia, difficulty with memory, fatigue, nervousness, psychomotor slowing, somnolence, and speech disorders and related speech problems were reported more frequently for topiramate than for placebo. In comparison, insomnia, dizziness, headache, and personality disorder were reported more frequently for placebo than for topiramate (Table 2). In addition to these neuropsychiatric adverse events, differences between placebo and topiramate were also found for weight decrease (2%, placebo; 15%, topiramate) and injury (20%, placebo; 8%, topiramate). Overall, the neuropsychiatric treatment-emergent adverse event profile was milder compared to that previously reported for topiramate-treated adult subjects with partial onset seizures.

	Placebo (N=41)		Topiramate (N=39)	
Preferred Term	No.	%	No.	%
Somnolence	6	15	10	26
Fatigue	3	7	7	18
Anorexia	3	7	6	15
Difficulty with memory	0	0	5	13
Headache	8	20	5	13
Dizziness	6	15	4	10
Nervousness	0	0	4	10
Psychomotor slowing	1	2	4	10
Speech disorders and related speech problems	1	2	4	10
Insomnia	5	12	1	3
Personality disorder	4	10	0	0

Table 2:	Incidence of Common ^a Treatment-Emergent Neuropsychiatric Adverse Events
	(All Randomized Subjects: Protocol CR005455)

^a Treatment-emergent neuropsychiatric adverse events reported by 10% or more subjects in either treatment group

No subject died during the trial or within 30 days of completion of the double-blind phase. Two subjects (one topiramate-treated and one placebo-treated) prematurely discontinued study medication and an additional six subjects (two placebo-treated and four topiramate-treated) had serious treatment-emergent adverse events.

Compared with placebo, topiramate had no effect on the subjects' mental status in terms of improvement in alertness, level of interaction with the environment, ability to perform the activities of daily living, and responsiveness to verbal requests.

No noteworthy hematologic, renal, or liver toxicity was observed (most of the few observed abnormalities were sporadic, transient, and did not lead to alteration in treatment). Except for mild changes in body weight, there were no noteworthy treatment-related changes in vital signs, ECGs, neurologic or physical examinations.

<u>CONCLUSION</u>: Topiramate was well-tolerated in this trial when administered to subjects with PGTC seizures. Topiramate was effective in reducing the rate of occurrence of PGTC seizures. Topiramate was also effective in reducing the rate of occurrence of all seizures.

Date of the report: 24 June 1997

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