## CENTER FOR DRUG EVALUATION AND RESEARCH

APPLICATION NUMBER: 022328Orig1s000

**OTHER REVIEW(S)** 



### 505(b)(2) ASSESSMENT

Application Information				
NDA # 22328	NDA Supplement #:S-	Efficacy Supplement Type SE-		
Proprietary Name: Inter				
Established/Proper Name: zolpidem tartrate SL				
Dosage Form: SL tablets				
Strengths: 1.75 mg and 3.5 mg oral SL tablets				
Applicant: Transcept Pharma, Inc.				
Date of Receipt: original submission 9/30/08; re-sub 1/14/11; re-sub 9/27/11				
PDUFA Goal Date: 11/27/11 Action Goal Date (if different):				
11/23/11				
Proposed Indication(s): as needed, middle-of-the- night (MOTN) insomnia				

	GENERAL INFORMATION			
1.	Is this application for a drug that is an "old" antibiotic as described in the Guidance to Industry, Repeal of Section 507 of the Federal Food, Drug and Cosmetic Act? (Certain antibiotics are not entitled to Hatch-Waxman patent listing and exclusivity benefits.)			
	YES		NO	X
	If "YES," proc	ceed to	question	#3.
2.	2. Is this application for a recombinant or biologically-derived product and/or protein or peptide product?			
	YES		NO	X
If "YES" contact the $(b)(2)$ review staff in the Immediate Office, Office of New Drugs.				



## INFORMATION PROVIDED VIA RELIANCE (LISTED DRUG OR LITERATURE)

3. List the information essential to the approval of the proposed drug that is provided by reliance on our previous finding of safety and efficacy for a listed drug or by reliance on published literature. (If not clearly identified by the applicant, this information can usually be derived from annotated labeling.)

Source of information (e.g.,	Information provided (e.g.,	
published literature, name of	pharmacokinetic data, or specific	
referenced product)	sections of labeling)	
NDA 19908 Ambien (zolpidem	Three Biopharm studies; specific sections	
tartrate)	PI changed	
	Five clinical studies; specific sections PI	
	changed	

4. Reliance on information regarding another product (whether a previously approved product or from published literature) must be scientifically appropriate. An applicant needs to provide a scientific "bridge" to demonstrate the relationship of the referenced and proposed products. Describe how the applicant bridged the proposed product to the referenced product(s). (Example: BA/BE studies)

This NDA comprises of the following 3 single-dose pharmacokinetic (PK)/ bioequivalence (BE) bridging studies in healthy adult and elderly subjects. Study ZI-15, provides comparative bioavailability information relative to reference Ambien®. Study ZI-14 includes comparative bioavailability of Intermezzo® 1.75 mg and 3.5 mg in elderly and adult cohorts. Study ZI-13 provides a bridging link between IND formulation and final commercial formulation used in different studies. Final commercial formulation was used in most of the studies including pivotal BE, pharmacodynamic, and efficacy studies.

#### RELIANCE ON PUBLISHED LITERATURE

5.	proposed drug product (i.e., the application <i>cannot</i> be approved without the published				
	literature)?	YES	X	NO	
	If ".	<b>NO</b> ," prod	ceed to	questior	ı #6.
	(b) Does any of the published literature necessary to support	approval	identify	a speci	fic
	(e.g., brand name) <i>listed</i> drug product?  Ambien (zolpidem tartrate)		x	NO	
YES	Amoren (zorpidem tartrate)		Λ	110	
-	If "YES", list the listed drug(s) identified by nar	' <b>NO"</b> , pro ne and an		•	



(c) Are the drug product(s) listed in (b) identified by the applicant as t	he liste	ed drug(s)	?
YES	X	NO	

### RELIANCE ON LISTED DRUG(S)

Reliance on published literature which identifies a specific approved (listed) drug constitutes reliance on that listed drug. Please answer questions #6-10 accordingly.

application <b>r</b> (approved dr	gardless of whether the applicant has explicitly referenced the listed drug(s), does the plication <b>rely</b> on the finding of safety and effectiveness for one or more listed drugs proved drugs) to support the approval of the proposed drug product (i.e., the plication cannot be approved without this reliance)?			
		YE	S x NO	
		If " <b>NO</b> ," pr	roceed to question #11.	
	ed drug(s) relied upon, and the plicitly identified the product			
Na	me of Drug	NDA/ANDA #	Did applicant specify reliance on the product? (Y/N)	
Ambien		19908	yes	
certification/s	hould specify reliance on the statement. If you believe ther identified as such by the appo	e is reliance on a listed pro licant, please contact the (l	oduct that has not been	
	pplement, does the suppleme 2) application? N/A		_	
If " <b>NO</b> ", pleas	se contact the (b)(2) review si	YE taff in the Immediate Office	- <u>-</u>	
-	the listed drug(s) relied upon proved in a 505(b)(2) applica		S NO X	
Naı	me of drug(s) approved in a 5	If "YES", ple	ease list which drug(s).	
b. App	proved by the DESI process?	YE		
Nai	me of drug(s) approved via th		ease list which drug(s).	
c. Des	scribed in a monograph?	YE		
Nai	me of drug(s) described in a r		ease list which drug(s).	



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