CENTER FOR DRUG EVALUATION AND RESEARCH

APPLICATION NUMBER: 22-304

ADMINISTRATIVE and CORRESPONDENCE DOCUMENTS



Department of Health and Human Services Food and Drug Administration

PATENT INFORMATION SUBMITTED WITH THE FILING OF AN NDA, AMENDMENT, OR SUPPLEMENT

For Each Patent That Claims a Drug Substance

Form Approved: OMB No. 0910-0513 Expiration Date: 04/30/10 See OMB Statement on Page 3.

NDA NUMBER 22-304 NAME OF APPLICANT / NDA HOLDER

(Active Ingredient), Drug Product (Formulation and Composition) and/or Method of Use				Ontho-McNeil-Janssen Pharmaceuticals, Inc.			
The following is provided in accordance with	Section 505	(b) and (c) of the Fe	ederal Fo	od, Drug, a	nd Cosmetic Act.		
TRADE NAME (OR PROPOSED TRADE NAME) TBD	<u></u>						
ACTIVE INGREDIENT(S) Tapentadol HCL		STRENGTH(S) 50mg, 75mg, 100mg					
DOSAGE FORM Tablets		· · · · · · · · · · · · · · · · · · ·					
This patent declaration form is required to be submamendment, or supplement as required by 21 CFR 314.53 Within thirtly (30) days after approval of an NDA or su declaration must be submitted pursuant to 21 CFR 3 or supplement. The information submitted in the declar upon by FDA for listing a patent in the Orange Book.	at the addres pplement, or 14.53(c)(2)(ii)	s provided in 21 CFR (within thirty (30) day with all of the requ	314.53(d) /s of issuited info	(4). Jance of a m Mation base	ew patent, a new patent		
For hand-written or typewriter versions (only) of t that does not require a "Yes" or "No" response), please	attach an ad	ditional page reference	cing the c	uestion num	ber.		
FDA will not list patent information if you file all patent is not eligible for listing.							
For each patent submitted for the pending NDA, information described below. If you are not sub- complete above section and sections 5 and 6.	amendmen mitting any	t, or supplement re patents for this p	erence ending	d above, yo NDA, amen	ou must submit all the dment, or supplement,		
1. GENERAL		<u></u>					
a. United States Patent Number RE 39,593E (Reissue of US 6,248,737)	b. Issue Da April 24 (US 6,24			c. Expiration June 19, 20	Date of Patent 018		
d. Name of Patent Owner Grünenthal GmbH	Address (of Zieglerstr. 6	Patent Owner) , 52078					
	City/State Aachen						
	ZIP Code Germany 52	078		if available) 55			
	Telephone N 49 241 569			E-Mail Addres patents@grun	• •		
 Name of agent or representative who resides or maintains a place of business within the United States authorized to receive notice of patent certification under section 505(b)(3) and (j)(2)(8) of the Federal Food, Drug, and 	Address (of agent or representative named in 1.e.) Crowell & Morning, P.O. Box 14300						
Cosmetic Act and 21 CFR 314.52 and 314 95 (il patent owner or NDA applicant/holder does not reside or have a place of business within the United States)	City/State Washington, D.C.						
Joseph D. Evans	ZIP Code 20044-4300	FAX Number (if available) 202-628-8844					
	Telephone N 202-624-250						
 Is the patent referenced above a patent that has been submapproved NDA or supplement referenced above? 	itted previous	y for the] Yes	⊠ No		
g. If the patent referenced above has been submitted previous date a new expiration date?	ly for listing, is	the expiration		Yes	□No		

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PSC Onspice (301) 443-1099 RF



For the patent referenced above, provide the following information on the drug substance, drug product and/or method of use that is the subject of the pending NDA, amendment, or supplement.							
2. Drug Substance (Active Ingredient)							
	Does the patent claim the drug substance that is the active ingredient in the drug product described in the pending NDA, amendment, or supplement?			· · · · · · · · · · · · · · · · · · ·	⊠ Yes	□No	
2.2 Does	the patent claim a dn	ing substance that is a different polymorph of the active			note)	Yes	_
2.3 If the	2.3 If the answer to question 2.2 is "Yes," do you certify that, as of the date of this declaration, you have test data						[2] I40
		product containing the po type of test data require			roduct	Yes	□ No
2.4 Spec	ify the polymorphic for	m(s) claimed by the pate	ent for which you hav	e the test results descri	bed in 2.3.		
					•		
2.5 Does	the patent claim only	a metabolite of the activ	e ingredient gending	in the NDA or sunnleme	ant?		
(Com		n section 4 below if the p				Yes	⊠No
	the patent claim only					Yes	⊠ No
		2.1 is a product-by-proce s required only if the pate				Yes	□ No
	Product (Composit	<u> </u>					
3.1 Does the patent claim the drug product, as defined in 21 CFR 314.3, in the pending NDA, amendment, or supplement?					⊠ Yes	□ No	
3.2 Does the patent claim only an intermediate?				·····		-	
3.3 If the patent referenced in 3.1 is a product-by-process patent, is the product claimed in the					Yes	⊠ No	
patent novel? (An answer is required only if the patent is a product-by-process patent.)					Yes	□No	
	4. Method of Use						
Sponsors must submit the information in section 4 for each method of using the pending drug product for which approval is being sought that is claimed by the patent. For each pending method of use claimed by the patent, provide the following information:							
	the patent claim one of ending NDA, amendm	or more methods of use ent, or supplement?	for which approval is	being sought in		⊠ Yes	□ No
	nt Claim Number(s) (a			nt claim(s) referenced in			
112,114,1	1,93,94,95,96,98,100,1 17,136,137,138,140		in the pending NDA	use for which approval i amendment, or supple	ment?	⊠ Yes	□ No
"Yes, ficity t ence	answer to 4.2 is identify with speci- the use with refer- to the proposed ng for the drug ict.			formation as identified s of moderate to severe a		the approved lab	eling.)
5. No Relevant Patents							
For this pending NDA, amendment, or supplement, there are no relevant patents that claim the drug substance (active ingredient), drug product (formulation or composition) or method(s) of use, for which the applicant is seeking approval and with respect to which a claim of patent infringement could reasonably be asserted if a person not licensed by the owner of the patent engaged in the manufacture, use, or sale of the drug product.							

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6. Declaration Certification						
6.1 The undersigned declares that this is an accurate and complete submission of patent information for the NDA, amendment, or supplement pending under section 505 of the Federal Food, Drug, and Cosmetic Act. This timesensitive patent information is submitted pursuant to 21 CFR 314.53. Lattest that Lam familiar with 21 CFR 314.53 and this submission complies with the requirements of the regulation. I verify under penalty of perjury that the foregoing is true and correct.						
6.2	Warning: A willfully and knowingly false statement is a criminal offense under 18 U.S.C. 1001. 6.2 Authorized Signature of NDA Applicant/Holder or Patent Owner (Attorney, Agent, Representative or other Authorized Official) (Provide Information below) Date Signed					
	Eller Colets	r	1-15-08			
NOT hold	NOTE: Only an NDA applicant/holder may submit this declaration directly to the FDA. A patent owner who is not the NDA applicant/holder is authorized to sign the declaration but may not submit it directly to FDA. 21 CFR 314.53(c)(4) and (d)(4).					
Chec	k applicable box and provide information below.					
	☐ NDA Applicant/Holder	⊠ N A	DA Applicant's/Holder's Attorney, Agent (Representative) or other uthorized Official			
	Patent Owner		ntent Owner's Attorney, Agent (Representative) or Other Authorized ficial			
	Name Ellen Coletti		-			
	Address Johnson & Johnson One Johnson & Johnson Plaza		City/State New Brunswick, NJ			
	ZIP Code 08933		Telephone Number 732-524-2359			
	FAX Number (if available) 732-524-5889		E-Mail Address (if available) ecoletti@corus.jnj.com			
The public reporting burden for this collection of information has been estimated to average 20 hours per response, including the time for reviewing instructions, searching existing data sources, gathering and maintaining the data needed, and completing and reviewing the collection of information. Send comments regarding this						
burden estimate or any other aspect of this collection of information, including suggestions for reducing this burden to: Food and Drug Administration CDER (HFD-007) 5600 Fishers Lane Rockville, MD 20857						
An agency may not conduct or sponsor, and a person is not required to respond to, a collection of information unless it displays a currently valid OMB control number.						
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FORM FDA 3542a (7/07)

INFORMATION AND INSTRUCTIONS FOR FORM 3542a

PATENT INFORMATION SUBMITTED WITH THE FILING OF AN NDA, AMENDMENT OR SUPPLEMENT

General Information

- To submit patent information to the agency the appropriate patent declaration form must be used. Two forms are available for patent submissions. The approval status of your New Drug Application will determine which form you should use.
- Form 3542a should be used when submitting patent information with original NDA submissions, NDA amendments and NDA supplements prior to approval.
- Form 3542 should be used after NDA or supplemental approval. This form is to be submitted within 30 days after approval of an application. This form should also be used to submit patent information relating to an approved supplement under 21 CFR 314.53(d) to change the formulation, add a new indication or other condition of use, change the strength, or to make any other patented change regarding the drug, drug product, or any method of use.
- Form 3542 is also to be used for patents issued after drug approval. Patents issued after drug approval are required to be submitted within 30 days of patent issuance for the patent to be considered "timely filed."
- Only information from form 3542 will be used for Orange Book publication purposes.
- Forms should be submitted as described in 21 CFR 314.53.
 Sending an additional copy of form 3542 to the Orange Book Staff will expedite patent publication in the Orange Book. The Orange Book Staff address (as of April 2007) is: Orange Book Staff, Office of Generic Drugs OGD/HFD-610, 7500 Standish Place, Rockville, MD 20855.
- The receipt date is the date that the patent information is date stamped in the central document room. Patents are considered listed on the date received.

Additional copies of these forms may be downloaded from the Internet at: http://www.fda.gov/opacom/morechoices/fdaforms/fdaforms.html.

First Section

Complete all items in this section,

1. General Section

Complete all items in this section with reference to the patent itself.

- 1c) Include patent expiration date, including any Hatch-Waxman patent extension already granted. Do not include any applicable pediatric exclusivity. The agency will include pediatric exclusivities where applicable upon publication.
- Id) Include full address of patent owner. If patent owner resides outside the U.S. indicate the country in the zip code block.

1e) Answer this question if applicable. If patent owner and NDA applicant/holder reside in the United States, leave space blank.

2. Drug Substance (Active Ingredient)

Complete all items in this section if the patent claims the drug substance that is the subject of the pending NDA, amendment, or supplement.

- 2.4) Name the polymorphic form of the drug identified by the parent.
- 2.5) A patent for a metabolite of the approved active ingredient may not be submitted. If the patent claims an approved method of using the approved drug product to administer the metabolite, the patent may be submitted as a method of use patent depending on the responses to section 4 of this form.
- Answer this question only if the patent is a product-byprocess patent.

3. Drug Product (Composition/Formulation)

Complete all items in this section if the patent claims the drug product that is the subject of the pending NDA, amendment, or supplement.

3.3) An answer to this question is required only if the referenced patent is a product-by-process patent.

4. Method of Use

Complete all items in this section if the patent claims a method of use of the drug product that is the subject of the pending NDA, amendment, or supplement (pending method of use).

- 4.2) For each pending method of use claimed by the patent, identify by number the claim(s) in the patent that claim the pending use of the drug. An applicant may list together multiple patent claim numbers and information for each pending method of use, if applicable. However, each pending method of use must be separately listed within this section of the form.
- 4.2a) Specify the part of the proposed drug labeling that is claimed by the patent.

5. No Relevant Patents

Complete this section only if applicable.

6. Declaration Certification

Complete all items in this section.

6.2) Authorized signature. Check one of the four boxes that best describes the authorized signature.

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DOCKET

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