CENTER FOR DRUG EVALUATION AND RESEARCH

APPLICATION NUMBER: 21-926

ADMINISTRATIVE and CORRESPONDENCE DOCUMENTS



TreximaTM

NDA 21-926

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 (Active Ingredient), Drug Product (Formulation and Composition) and/or Method of Use

Form Approved: OMB No. 0910-0513
Expiration Date: 07/31/06
See OMB Statement on Page 3.

NDA NUMBER	
21-926	
21-720	
NAME OF APPLICANT /	NDA HOLDER
POZEN Inc.	
1 0	

The following is provided in accordance with Section 505(b) and (c) of the Federal Food, Drug, and Cosmetic Act.			
TRADE NAME (OR PROPOSED TRADE NAME) TREXIMA			
ACTIVE INGREDIENT(S) sumatriptan succinate and naproxen sodium		STRENGTH(S) sumatriptan 85 mg (as the succinate) and naproxen sodium 500 mg	
DOSAGE FORM Tablet (oral)			
This patent declaration form is required to be submit amendment, or supplement as required by 21 CFR 314.53 at Within thirty (30) days after approval of an NDA or supplement. The information submitted in the declaration by FDA for listing a patent in the Orange Book.	at the addres oplement, or 4.53(c)(2)(ii)	s provided in 21 CFR 314.53(d within thirty (30) days of iss with all of the required info)(4). suance of a new patent, a new patent ormation based on the approved NDA
For hand-written or typewriter versions (only) of the that does not require a "Yes" or "No" response), please a			
FDA will not list patent information if you file an patent is not eligible for listing.	incomplet	te patent declaration or th	ne patent declaration indicates the
For each patent submitted for the pending NDA, amendment, or supplement referenced above, you must submit all the information described below. If you are not submitting any patents for this pending NDA, amendment, or supplement, complete above section and sections 5 and 6.			
1. GENERAL			
a. United States Patent Number 4,816,470	b. Issue Dat 3/28/1989	e of Patent	c. Expiration Date of Patent 12/28/2006
d. Name of Patent Owner Glaxo Group Limited	Address (of Patent Owner) Glaxo Wellcome House, Berkeley Avenue City/State Greenford, England		
	ZIP Code UB6 0NN		FAX Number (if available) 919-483-7988
	Telephone N 919-483-69	1	E-Mail Address (if available)
e. 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)(B) of the Federal Food, Drug, and	Address (of agent or representative named in 1.e.) One Franklin Plaza		
Cosmetic Act and 21 CFR 314.52 and 314.95 (if patent owner or NDA applicant/holder does not reside or have a place of business within the United States)	City/State Philadelphia, PA		
SmithKline Beecham Corp.	ZIP Code 19101		FAX Number (if available) 919-483-7988
	Telephone N 919-483-6	983	E-Mail Address (if available)
f. Is the patent referenced above a patent that has been submitted previously for the approved NDA or supplement referenced above? Yes No			
g. If the patent referenced above has been submitted previously for listing, is the expiration			



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?		
 Does the patent claim a drug substance the ingredient described in the pending NDA, 	amendment, or supplement?	Yes	⊠ No
demonstrating that a drug product contain	ou certify that, as of the date of this declaration, you have test dat ing the polymorph will perform the same as the drug product ta required is described at 21 CFR 314.53(b).	ta Yes	□No
2.4 Specify the polymorphic form(s) claimed b	y the patent for which you have the test results described in 2.3.		
	f the active ingredient pending in the NDA or supplement? ow if the patent claims a pending method of using the pending)	Yes	⊠ No
2.6 Does the patent claim only an intermediat	e?	Yes	⊠ No
	-by-process patent, is the product claimed in the if the patent is a product-by-process patent.)	Yes	☐ No
3. Drug Product (Composition/Formula	tion)		
3.1 Does the patent claim the drug product, a amendment, or supplement?	s defined in 21 CFR 314.3, in the pending NDA,	∑ Yes	☐ No
3.2 Does the patent claim only an intermediat	e?	Yes	⊠ No
	-by-process patent, is the product claimed in the if the patent is a product-by-process patent.)	Yes	☐ No
4. Method of Use			
	n section 4 separately for each patent claim claiming a . For each method of use claim referenced, provide the follow		
4.1 Does the patent claim one or more metho the pending NDA, amendment, or suppler		⊠ Yes	☐ No
4.2 Patent Claim Number (as listed in the pat Claim 16	of use for which approval is being sought in the pending	NDA,	
4.2a If the answer to 4.2 is "Yes," identify with specificity the use with reference to the proposed labeling for the drug product. Use: (Submit migraine)	amendment, or supplement? indication or method of use information as identified specifically in	Yes n the approved l	No abeling.)
5. No Relevant Patents			
drug product (formulation or composition) or m	nent, there are no relevant patents that claim the drug substance (ethod(s) of use, for which the applicant is seeking approval and wisonably be asserted if a person not licensed by the owner of the particle.	ith respect to	



	Frexima TM			NDA 21-926
6. D	6. Declaration Certification			
6.1	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. I attest that I am 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.			
	Warning: A willfully and knowingly false statem			1001.
6.2	Authorized Signature of NDA Applicant/Holder or Patent Cother Authorized Official) (Provide Information, below)	Owner <i>(Attorney, .</i>	Agent, Representative or	Date Signed 7/14/05
	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	ck applicable box and provide information below.			
	NDA Applicant/Holder		A Applicant's/Holder's Attorney, horized Official	Agent (Representative) or other
	Patent Owner		ent Owner's Attorney, Agent (Ro	epresentative) or Other Authorized
	Name POZEN Inc.		·	
	Address 1414 Raleigh Road, Suite 400		City/State Chapel Hill, NC	
	ZIP Code 27517		Telephone Number 919-913-1030	
	FAX Number (if available) 919-913-1039		E-Mail Address (if available)	

The public reporting burden for this collection of information has been estimated to average 9 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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4. Method of Use (continue	ed)		
Sponsors must submit the information in section 4 separately for each patent claim claiming a method of using the pending drug product for which approval is being sought. For each method of use claim referenced, provide the following information:			
4.1 Does the patent claim one obeing sought in the pending N	or more methods of use	for which approval is	⊠ Yes □ No
4.2 Patent Claim Number (as I:		Does the patent claim referenced in 4.2 claim a pending method of use for which approval is being sought in	
4.2a If the answer to 4.2 is "Yes," identify the use with specific reference to the proposed labeling for the drug product	Use (Submit indication o migraine	the pending NDA, amendment or supplement? or method of use information as identified specifically in the approve	⊠ Yes □ No ed labeling.)



DOCKET

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