## CENTER FOR DRUG EVALUATION AND RESEARCH

APPLICATION NUMBER: 22-228

# ADMINISTRATIVE and CORRESPONDENCE DOCUMENTS



Department of Health and Human Services Food and Drug Administration

### PATENT INFORMATION SUBMITTED WITH THE FILING

Form Approved: OMB No. 0910-0513 Expiration Date: 7/31/10 See OMB Statement on Page 3.

NDA NUMBER
22-288

OF AN NDA, AMENDMENT, OR SUPPLEMENT			22-288			
For Each Patent That Claims a Drug Substance (Active Ingredient), Drug Product (Formulation and Composition) and/or Method of Use			NAME OF APPLICANT/NDA HOLDER			
			ISTA Pharmaceuticals <sup>®</sup> , Inc.			
The following is provided in accordance with	Section 50	5(b) and (c) of th	e Federal F	ood, Dr	rug, and Cosmetic Act.	
TRADE NAME (OR PROPOSED TRADE NAME) Bepreve <sup>TM</sup>						
ACTIVE INGREDIENT(S)		STRENGTH(S)	<del>~~</del>			
Bepotastine Besilate		1.5%				
DOSAGE FORM		L				
Ophthalmic solution	<u> </u>				<u></u>	
This patent declaration form is required to be submamendment, or supplement as required by 21 CFR 314 Within thirty (30) days after approval of an NDA or supplement must be submitted pursuant to 21 CFR 314 supplement. The information submitted in the declaration by FDA for listing a patent in the Orange Book.	4.53 at the ac pplement, or 4.53(c)(2)(ii)	ddress provided in r within thirty (30) with all of the req	n 21 CFR 31 days of issu juired informa	4.53(d) uance of ation ba	(4). f a new patent, a new patent ised on the approved NDA or	
For hand-written or typewriter versions (only) of thi does not require a "Yes" or "No" response), please atta	is report: If a	additional space i onal page referen	is required fo cing the que	or any na estion nu	arrative answer (i.e., one that	
FDA will not list patent information if you submit a patent is not eligible for listing.	an incomple	ete patent decla	ration or th	e pater	nt declaration indicates the	
For each patent submitted for the pending NDA, a information described below. If you are not submodule to above section and sections 5 and 6.	amendment nitting any	t, or supplement patents for this	t referenced pending N	d above IDA, an	e, you must submit all the nendment, or supplement,	
AKGENERAL TELEVISION OF THE PROPERTY OF THE PR			in periodici di carini di carini di preside di carini di carini di carini periodici di carini	An order species. An order species and species species.		
a. United States Patent Number 6,780,877	b. Issue Da 8/24/04	ate of Patent	C	2. Expirat 12/25/	tion Date of Patent /17	
d. Name of Patent Owner		f Patent Owner)	·			
(1) Ube Industries, Ltd.		Nishihonmachi 1-Cl osho-Machi 3-Chor			ch;	
	City/State	USIIO ITILOII. 2	the, Chico,	Course	MI	
(2) Tanabe Seiyaku Co. Ltd.		uchi 755-863, Japan				
	ZIP Code		FAX	Number	(if available)	
	Telephone N (908) 607-1	1950		E-Mail Address (if available)		
<ul> <li>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 Cosmetic Act and 21 CFR 314.52 and 314.95 (if patent owner or NDA</li> </ul>	Address (of agent or representative named in 1.e.) ISTA Pharmaceuticals, Inc. 15295 Alton Parkway City/State					
applicant/holder does not reside or have a place of business within the United States)	Irvine, CA		* = :			
Mary Garrett	ZIP Code 92618			FAX Number (if available) (949) 727-0833		
Vice President Regulatory Affairs, Quality Assurance, and Compliance	Telephone N (949) 788-5	303		es <i>(if available)</i> avision.com		
f. Is the patent referenced above a patent that has been submapproved NDA or supplement referenced above?	nitted previous	ly for the	Y€	es	✓ No	
g. If the patent referenced above has been submitted previous date a new expiration date?	ly for listing, is	s the expiration	Y€	<b>9</b> 8	✓ No	

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Page 1

PSC Graphics (301) 443-1090 EF



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)					
2.1 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 drug substance that is a different polymorph of the active ingredient described in the pending NDA, amendment, or supplement?	☐ Yes	☑ No			
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 demonstrating that a drug product containing the polymorph will perform the same as the drug product described in the NDA? The type of test data required is described at 21 CFR 314.53(b).	Yes	□ No			
2.4 Specify the polymorphic form(s) claimed by the patent for which you have the test results described in 2.3.					
·					
2.5 Does the patent claim only a metabolite of the active ingredient pending in the NDA or supplement?		· · · · · · · · · · · · · · · · · · ·			
(Complete the information in section 4 below if the patent claims a pending method of using the pending drug product to administer the metabolite.)	☐ Yes	✓ No			
2.6 Does the patent claim only an intermediate?					
	Yes	☑ No			
2.7 If the patent referenced in 2.1 is a product-by-process patent, is the product claimed in the patent novel? (An answer is required only if the patent is a product-by-process patent.)	Yes	✓ No			
3. Drug Product (Composition/Formulation)	i (Colordonia dell'illa dell' Regionalità dell'illa dell'illa Regionalità dell'illa				
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?	☐ Yes	☑ No			
3.3 If the patent referenced in 3.1 is a product-by-process patent, is the product claimed in the 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 produ sought that is claimed by the patent. For each pending method of use claimed by the patent, provide the fo	ct for which ap llowing informa	proval is being tion:			
4.1 Does the patent claim one or more methods of use for which approval is being sought in the pending NDA, amendment, or supplement?	☐ Yes	☑ No			
4.2 Patent Claim Number(s) (as listed in the patent)  Does (Do) the patent claim(s) referenced in 4.2 claim a pending method of use for which approval is being sought in the pending NDA, amendment, or supplement?	Yes	☐ No			
4.2a If the answer to 4.2 is Use: (Submit indication or method of use information as identified specifically in	the proposed lab	eling.)			
"Yes," identify with speci- ficity the use with refer-					
ence to the proposed labeling for the drug					
product.	,				
	Tie (Digniya digniya maka da an	STATE DAMES THE SALE AND STATE OF THE SALES			
5. No Relevant Patents	Celerina da del Personal del Colo de despois de despois de del	ing by the second			
For this pending NDA, amendment, or supplement, there are no relevant patents that claim the drug substance (activity product (formulation or composition) or method(s) of use, for which the applicant is seeking approval and with a claim of patent infringement could reasonably be asserted if a person not licensed by the owner of the patent engagement could reasonably be asserted if a person not licensed by the owner of the patent engagement could reasonably be asserted if a person not licensed by the owner of the patent engagement could reasonably be asserted if a person not licensed by the owner of the patent engagement could reasonably be asserted if a person not licensed by the owner of the patent engagement could reasonably be asserted if a person not licensed by the owner of the patent engagement could reasonably be asserted if a person not licensed by the owner of the patent engagement could reasonably be asserted if a person not licensed by the owner of the patent engagement could reasonably be asserted if a person not licensed by the owner of the patent engagement could reasonably be asserted if a person not licensed by the owner of the patent engagement could reasonably be asserted if a person not licensed by the owner of the patent engagement could reasonably be asserted if a person not licensed by the owner of the patent engagement could be asserted in the patent engagement could be asserted in the patent engagement of the patent engagement of the patent engagement of the patent engagement engagement of the patent engagement e	espect to which	☐ Yes			
FORM FDA 3542a (7/07)		Page 2			



Page 2

Declaration Certification	day fine transportation with			Parte to the Control of the Control		
1 The undersigned declares that this is amendment, or supplement pending usensitive patent information is submit this submission complies with the rectrue and correct.  Warning: A willfully and knowingly follows:	under section 50 Ited pursuant to puirements of th	05 of the 1 21 CFR in 1990 of	Federal Food, Drug, and 314.53. I attest that I am ion. I verify under pena	d Cosmetic Act. This time- n familiar with 21 CFR 314.53 and lity of perjury that the foregoing		
Warning: A willfully and knowingly fal			· · · · · · · · · · · · · · · · · · ·			
2 Authorized Signature of NDA Applicant/Holde other Authorized Official) (Provide Information		(Attorney, )	Agent, Representative or	Date Signed		
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Marvin & lan	o H			03 POTOBEN 20		
OTE: Only an NDA applicant/holder may su older is authorized to sign the declaration but						
neck applicable box and provide information	below.					
✓ NDA Applicant/Holder		NDA Applicant's/Holder's Attorne		ey, Agent (Representative) or other		
☐ Patent Owner		Patent Owner's Attorney, Agent (R		lepresentative) or Other Authorized		
Name	·					
ISTA Pharmaceuticals®, Inc.						
Address 15295 Alton Parkway		1	City/State			
13293 Alton Parkway		'	rvine, CA			
ZIP Code			Telephone Number			
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### **EXCLUSIVITY SUMMARY**

NDA # 22-288	SUPPL#	HF	D # 520			
Trade Name Bepreve						
Generic Name bepota	stine besilate ophthalmic solution 1.5	5%				
Applicant Name Ista I	harmaceuticals, Inc.					
Approval Date, If Know	vn					
PART I IS AN E	EXCLUSIVITY DETERMINATIO	ON NEEDED?				
1. An exclusivity determination will be made for all original applications, and all efficacy supplements. Complete PARTS II and III of this Exclusivity Summary only if you answer "yes" to one or more of the following questions about the submission.						
a) Is it a 505(b)	(1), 505(b)(2) or efficacy supplement	nt? YES ⊠	NO 🗌			
If yes, what type? Speci	ify 505(b)(1), 505(b)(2), SE1, SE2, S	SE3,SE4, SE5, SE6	, SE7, SE8			
505(b)(1)						
c) Did it require the review of clinical data other than to support a safety claim or change in labeling related to safety? (If it required review only of bioavailability or bioequivalence						
data, answer "no	).")	YES 🖂	NO 🗌			
not eligible for	s "no" because you believe the study is exclusivity, EXPLAIN why it is a greeing with any arguments made b ilability study.	bioavailability stu	dy, including your			
	ement requiring the review of clinic scribe the change or claim that is sup-					



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