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APPLICATION NUMBER:

204592Orig1s000

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505(b)(2) ASSESSMENT

Application Information						
NDA # 204592	NDA Supplement #:	S-	Efficacy Supplement Type SE-			
Proprietary Name: Zory	volex					
Established/Proper Nam	e: diclofenac					
Dosage Form: Capsule	5					
Strengths: 18 mg and 3	5 mg					
Applicant: Iroko Phari	Applicant: Iroko Pharmaceuticals					
Date of Receipt: December 20, 2012						
PDUFA Goal Date: Oct	ober 20, 2013	Action	Goal Date (if different):			
October 18, 2013						
RPM: Swati Patwardhan						
Proposed Indication(s): treatment of acute pain of mild to moderate (b) (4) in adults						

GENERAL INFORMATION

1) Is this application for a recombinant or biologically-derived product and/or protein or peptide product *OR* is the applicant relying on a recombinant or biologically-derived product and/or protein or peptide product to support approval of the proposed product?

YES	NO	\times

If "YES "contact the (b)(2) review staff in the Immediate Office, Office of New Drugs.

INFORMATION PROVIDED VIA RELIANCE (LISTED DRUG OR LITERATURE)

2) 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 by reliance on published literature, or by reliance on a final OTC monograph. *(If not clearly identified by the applicant, this information can usually be derived from annotated labeling.)*

Source of information* (e.g., published literature, name of listed drug(s), OTC final drug monograph)	Information relied-upon (e.g., specific sections of the application or labeling)
NDA 020142 Cataflam 50 mg	FDA's previous finding of safety and efficacy

*each source of information should be listed on separate rows, however individual literature articles should not be listed separately

3) 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)

Two Phase 1 studies were conducted.

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Study DICI-12-07, a pivotal study, was a randomized, single-dose, five-way crossover, relative bioavailability study of ZorvolexTM (diclofenac ^{(b) (4)}) capsules 18 mg and 35 mg and cataflam® 50 mg tablets, in healthy subjects under fed and fasting conditions, conducted with commercial scale formulation.

RELIANCE ON PUBLISHED LITERATURE

4) (a) Regardless of whether the applicant has explicitly stated a reliance on published literature to support their application, is reliance on published literature necessary to support the approval of the proposed drug product (i.e., the application *cannot* be approved without the published literature)?

YES		NO	\boxtimes
If "NO," proce	ed to	question	#5.

(b) Does any of the published literature necessary to support approval identify a specific (e.g., brand name) *listed* drug product?

YES	NO	\boxtimes
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If "NO", proceed to question #5.

If "YES", list the listed drug(s) identified by name and answer question #4(c).

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

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 #5-9 accordingly.

5) Regardless of whether the applicant has explicitly cited reliance on listed drug(s), does the application **rely** on the finding of safety and effectiveness for one or more listed drugs (approved drugs) to support the approval of the proposed drug product (i.e., the application cannot be approved without this reliance)?

Y	ES	\boxtimes	NO	
If "NO, "	proce	ed to	question	#10.

6) Name of listed drug(s) relied upon, and the NDA #(s). Please indicate if the applicant explicitly identified the product as being relied upon (see note below):

Name of Listed Drug	NDA #	Did applicant specify reliance on the product? (Y/N)
Cataflam	020142	Y

Applicants should specify reliance on the 356h, in the cover letter, and/or with their patent certification/statement. If you believe there is reliance on a listed product that has not been explicitly identified as such by the applicant, please contact the (b)(2) review staff in the Immediate Office, Office of New Drugs.

7) If this is a (b)(2) supplement to an original (b)(2) application, does the supplement rely upon the same listed drug(s) as the original (b)(2) application?

	N/A	\bowtie	YES		NO	
If this application is a $(b)(2)$ supplement to an original	al (b)(1)	applica	tion or	not a s	uppleme	ntal
		a	pplicat	ion, an	swer "N	/ A" .
If "NO", please contact the $(b)(2)$ review staff in the	he Imme	ediate O	ffice, O	ffice of	New Dr	ugs.

- 8) Were any of the listed drug(s) relied upon for this application:
 - a) Approved in a 505(b)(2) application?

	11	20		110	
If "Y	'ES ", pl	lease	list w	hich druz	g(s).
Name of drug(s) approved in a 505(b)(2) application	on:				

b) Approved by the DESI process?



NO

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VEC

Name of drug(s) approved via the DESI process:

c) Described in a final OTC drug monograph?

YES		NO	\boxtimes
If "YES", please	list w	hich drug	(s).

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Name of drug(s) described in a final OTC drug monograph:

d) Discontinued from marketing?

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YES NO X If "**YES**", please list which drug(s) and answer question d) i. below. If "**NO**", proceed to question #9. Name of drug(s) discontinued from marketing:

i) Were the products discontinued for reasons related to safety or effectiveness? YES NO (Information regarding whether a drug has been discontinued from marketing for reasons of safety or effectiveness may be available in the Orange Book. Refer to section 1.11 for an explanation, and section 6.1 for the list of discontinued drugs. If a determination of the reason for discontinuation has not been published in the Federal Register (and noted in the Orange Book), you will need to research the archive file and/or consult with the review team. Do not rely solely on any statements made by the sponsor.)

9) Describe the change from the listed drug(s) relied upon to support this (b)(2) application (for example, "This application provides for a new indication, otitis media" or "This application provides for a change in dosage form, from capsule to solution").

The proposed drug product is a reformulation of diclofenac with reduced particle size and is the free acid and not a salt. Cataflam is the potassium diclofenac salt. The proposed drug product is 20% lower in strength compared to the listed drug, Cataflam. The application also provides for a change in dosage from a tablet to a capsule.

The purpose of the following two questions is to determine if there is an approved drug product that is equivalent or very similar to the product proposed for approval that should be referenced as a listed drug in the pending application.

The assessment of pharmaceutical equivalence for a recombinant or biologically-derived product and/or protein or peptide product is complex. If you answered **YES to question #1**, proceed to question #12; if you answered **NO to question #1**, proceed to question #10 below.

10) (a) Is there a pharmaceutical equivalent(s) to the product proposed in the 505(b)(2) application that is already approved (via an NDA or ANDA)?

(**Pharmaceutical equivalents** are drug products in identical dosage forms intended for the same route of administration that: (1) contain identical amounts of the identical active drug ingredient, i.e., the same salt or ester of the same therapeutic moiety, or, in the case of modified release dosage forms that require a reservoir or overage or such forms as prefilled syringes where residual volume may vary, that deliver identical amounts of the active drug ingredient over the identical dosing period; (2) do not necessarily contain the same inactive ingredients; <u>and</u> (3) meet the identical compendial or other applicable standard of identity, strength, quality, and purity, including potency and, where applicable, content uniformity, disintegration times, and/or dissolution rates. (21 CFR 320.1(c), FDA's "Approved Drug Products with Therapeutic Equivalence Evaluations" (the Orange Book)).

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