CENTER FOR DRUG EVALUATION AND RESEARCH

APPLICATION NUMBER: 21-450

ADMINISTRATIVE and CORRESPONDENCE DOCUMENTS



EXCLUSI	VITY SUM	MARY for ND	A # <u>21-</u>	450	SUPPL	#
Trade N	ame Zomic	J Nasal Spr	ay	Generic	Name Zolmi	triptan
	nt Name 1 1 Date	AstraZeneca 0/26/03	<u> </u>	HFD- 1	20	
PART I:	IS AN E	CLUSIVITY	DETERMIN	ATION N	EEDED?	
appl: Parts answe	ications, s II and	but only : III of this to one or :	for certa Exclusi	ain supp vity Su	nde for all plements. (nmmary only owing quest	Complete
a)	Is it an	original 1	NDA?		YES/_x/	NO /_/
b)	Is it an	effective	ness supp	olement?	YES //	NO / x /
	If yes,	what type	(SE1, SE2	2, etc.)	?	
c)	support safety?	a safety c	laim or o quired re	change i eview or	n labeling lly of bioa	ther than to related to vailability
					YES / <u>x</u> /	NO //
	bioavail exclusiv includin made by	ability str ity, EXPLA: g your reas	idy and, IN why it sons for ant that	therefo is a b disagre	ore, not el: pioavailabil	lity study, any argument
					ne review o ss supplemen	f clinical nt, describe
	the chan data:	ge or clai	n that is	s suppor	rted by the	clinical
d)	Did the	applicant :	request e	exclusiv	rity?	
					YES /	_/ NO / <u>x</u> /
		an manage of	Page	1		

If the answer to (d) is "yes," how many years of exclusivity did the applicant request?

e) Has pediatric exclusivity been granted for this Active Moiety?
YES // NO / <u>x</u> /
IF YOU HAVE ANSWERED "NO" TO ALL OF THE ABOVE QUESTIONS, GO DIRECTLY TO THE SIGNATURE BLOCKS ON Page 9.
2. Has a product with the same active ingredient(s), dosage for strength, route of administration, and dosing schedule previously been approved by FDA for the same use? (Rx to OTC) Switches should be answered No - Please indicate as such).
YES // NO /_x_/
If yes, NDA # Drug Name
IF THE ANSWER TO QUESTION 2 IS "YES," GO DIRECTLY TO THE SIGNATURE BLOCKS ON Page 9.
3. Is this drug product or indication a DESI upgrade?
YES // NO /_x_/
IF THE ANSWER TO QUESTION 3 IS "YES," GO DIRECTLY TO THE SIGNATURE BLOCKS ON Page 9 (even if a study was required for thupgrade).

PART II: FIVE-YEAR EXCLUSIVITY FOR NEW CHEMICAL ENTITIES





(Answer either #1 or #2, as appropriate)

1. Single active ingredient product.

Has FDA previously approved under section 505 of the Act any drug product containing the same active moiety as the drug under consideration? Answer "yes" if the active moiety (including other esterified forms, salts, complexes, chelates or clathrates) has been previously approved, but this particular form of the active moiety, e.g., this particular ester or salt (including salts with hydrogen or coordination bonding) or other non-covalent derivative (such as a complex, chelate, or clathrate) has not been approved. Answer "no" if the compound requires metabolic conversion (other than deesterification of an esterified form of the drug) to produce an already approved active moiety.

YES / x / NO / ___/

If "yes," identify the approved drug product(s) containing the active moiety, and, if known, the NDA #(s).

NDA # 20-768 Zomig tablets

NDA # 21-231 Zomig-ZMT

NDA #

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2. Combination product.

If the product contains more than one active moiety (as defined in Part II, #1), has FDA previously approved an application under section 505 containing any one of the active moieties in the drug product? If, for example, the combination contains one never-before-approved active moiety and one previously approved active moiety, answer "yes." (An active moiety that is marketed under an OTC monograph, but that was never approved under an NDA, is considered not previously approved.)

YES /__/ NO /___,
If "yes," identify the approved drug product(s) containing the active moiety, and, if known, the NDA #(s).

NDA #

Page 3



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ON ORIGINAL

NDA #

NDA #

IF THE ANSWER TO QUESTION 1 OR 2 UNDER PART II IS "NO," GO DIRECTLY TO THE SIGNATURE BLOCKS ON Page 9. IF "YES," GO TO PART III.

PART III: THREE-YEAR EXCLUSIVITY FOR NDA'S AND SUPPLEMENTS

To qualify for three years of exclusivity, an application or supplement must contain "reports of new clinical investigations (other than bioavailability studies) essential to the approval of the application and conducted or sponsored by the applicant." This section should be completed only if the answer to PART II, Question 1 or 2, was "yes."

1. Does the application contain reports of clinical investigations? (The Agency interprets "clinical investigations" to mean investigations conducted on humans other than bioavailability studies.) If the application contains clinical investigations only by virtue of a right of reference to clinical investigations in another application, answer "yes," then skip to question 3(a). If the answer to 3(a) is "yes" for any investigation referred to in another application, do not complete remainder of summary for that investigation.

YES /<u>X</u>/ NO /_/

IF "NO," GO DIRECTLY TO THE SIGNATURE BLOCKS ON Page 9.

2. A clinical investigation is "essential to the approval" if the Agency could not have approved the application or supplement without relying on that investigation. Thus, the investigation is not essential to the approval if 1) no clinical investigation is necessary to support the supplement or application in light of previously approved applications (i.e., information other than clinical trials, such as bioavailability data, would be sufficient to provide a basis for approval as an ANDA or 505(b)(2) application because of what is already known about a previously approved product), or 2) there are published reports of studies (other than those conducted or sponsored by the applicant) or other publicly





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