## CENTER FOR DRUG EVALUATION AND RESEARCH

**APPLICATION NUMBER:** 

208026Orig1s000

# ADMINISTRATIVE and CORRESPONDENCE DOCUMENTS



### **EXCLUSIVITY SUMMARY**

NDA # 208026

Trade Name Jentadueto XR

Generic Name (linagliptin and metformin hydrochloride extended-release) tablets

Applicant Name Boehringer Ingelheim Pharmaceuticals, Inc.

Approval Date, If Known May 27, 2016

### PART I IS AN EXCLUSIVITY DETERMINATION NEEDED?

a) Is it a 505(b)(1), 505(b)(2) or efficacy supplement?

1.	An	exclu	sivity	determination	will	be	made	for	all	original	applications,	and	all	efficacy
sup	plen	nents.	Comp	olete PARTS II	and	III	of this	Exc	lusi	vity Sum	mary only if	you a	ınsw	er "yes"
to o	one c	or mor	e of th	e following qu	estion	is a	bout th	ie su	bm	ission.				

	YES 🖂	NO 📙
If yes, what type? Specify 505(b)(1), 505(b)(2), SE1, SE2, SE3,SI	E4, SE5, SE	6, SE7, SE8
505(b)(1)		
b) Did it require the review of clinical data other than to s in labeling related to safety? (If it required review himsen in the control of	1.1	,
bioequivalence data, answer "no.")	YES 🗌	NO 🖂

If your answer is "no" because you believe the study is a bioavailability study and, therefore, not eligible for exclusivity, EXPLAIN why it is a bioavailability study, including your reasons for disagreeing with any arguments made by the applicant that the study was not simply a bioavailability study.

Three clinical studies were completed by the applicant, one study for bioavailability (Study 1288.8) and two studies for bioequivalence (Studies 1288.9 and 1288.11) were completed. Per the note above, BA and BE studies do not qualify. The applicant acknowledges only BA and BE studies were completed for this application, they refer to NDA 201280 and have right of reference to NDA 021748 for safety and efficacy.

If it is a supplement requiring the review of clinical data but it is not an effectiveness supplement, describe the change or claim that is supported by the clinical data:



N/A

c) Did the applicant request exclusivity?	YES 🗌	NO 🖂
If the answer to (c) is "yes," how many years of exclusivity	did the applica	ant request?
N/A		
d) Has pediatric exclusivity been granted for this Active Mo	oiety? YES	NO 🖂
<u>If the answer to the above question in YES</u> , is this approval a in response to the Pediatric Written Request?	result of the st	udies submitted
N/A		
IF YOU HAVE ANSWERED "NO" TO <u>ALL</u> OF THE ABOVE OF TO THE SIGNATURE BLOCKS AT THE END OF THIS DOCU		GO DIRECTLY
2. Is this drug product or indication a DESI upgrade?	YES 🗌	NO 🖂
IF THE ANSWER TO QUESTION 2 IS "YES," GO DIRECT BLOCKS ON PAGE 8 (even if a study was required for the upgraded)		E SIGNATURE
PART II FIVE-YEAR EXCLUSIVITY FOR NEW CHEM (Answer either #1 or #2 as appropriate)	IICAL ENTI	ΓIES
1. Single active ingredient product.		
Has FDA previously approved under section 505 of the Act any same active moiety as the drug under consideration? Answe (including other esterified forms, salts, complexes, chelates or c approved, but this particular form of the active moiety, e.g., this particular salts with hydrogen or coordination bonding) or other non-cocomplex, chelate, or clathrate) has not been approved. Answer metabolic conversion (other than deesterification of an esterified for already approved active moiety.	er "yes" if the lathrates) has articular ester of ovalent derivation if the control of the contr	e active moiety been previously r salt (including tive (such as a appound requires
	YES 🗌	NO 🗌



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

NDA#

NDA#

NDA#

### 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 <u>any one</u> 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# NDA 201280 Tradjenta (linagliptin) tablets

NDA# NDA 021748 Glumetza (metformin extended-release) tablets

NDA# NDA 201281 Jentadueto (linagliptin and metformin) tablets

IF THE ANSWER TO QUESTION 1 OR 2 UNDER PART II IS "NO," GO DIRECTLY TO THE SIGNATURE BLOCKS ON PAGE 8. (Caution: The questions in part II of the summary should only be answered "NO" for original approvals of new molecular entities.) IF "YES," GO TO PART III.

## PART III THREE-YEAR EXCLUSIVITY FOR NDAs 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 NO
IF "NO," GO DIRECTLY TO THE SIGNATURE BLOCKS ON PAGE 8.
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 available data that independently would have been sufficient to support approval of the application, without reference to the clinical investigation submitted in the application.
(a) In light of previously approved applications, is a clinical investigation (either conducted by the applicant or available from some other source, including the published literature) necessary to support approval of the application or supplement?  YES \( \subseteq \text{NO} \subseteq \)
If "no," state the basis for your conclusion that a clinical trial is not necessary for approval AND GO DIRECTLY TO SIGNATURE BLOCK ON PAGE 8:
(b) Did the applicant submit a list of published studies relevant to the safety and effectiveness of this drug product and a statement that the publicly available data would not independently support approval of the application?  YES NO
(1) If the answer to 2(b) is "yes," do you personally know of any reason to disagree with the applicant's conclusion? If not applicable, answer NO.
YES NO NO
If yes, explain:



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