

#### November 2015

# **HERZUMA**

Ministry of Food and Drug Safety

#### **APPROVED**

	PART A - ADMINIS	TRATIVE INFORMATION			
Entered by:	<b>Biosimilar Product Information</b>				
MAH	Name of the biosimilar medicinal product	Herzuma			
MAH	МАН	Celltrion Inc. 13-6 Songdo-dong, Yeonsu-gu, Incheon City, Republic of Korea			
NRA	Authorisation / Licence number	Celltrion / 6 Celltrion / 7			
MAH / NRA	API manufacturing facilities and batch release site for the finished product (if applicable)	Confidential – Not Released			
MAH	Name of the active substance	Trastuzumab (INN)			
MAH	Pharmaco-therapeutic group	ATC code: L01XC03			
MAH	Substance category	Monoclonal antibody			
MAH	Pharmaceutical form	White to pale yellow lyophilized powder / Clear to slightly opalescent/ Colourless to pale yellow solution			
MAH	Quantitative composition	150 mg/vial 440mg/vial			
MAH	Route of administration	IV (Intravenous, Infusion)			
MAH	Packaging/material	Glass vial			
MAH	Package size(s)	1 vial/pack			
MAH	Local legal basis	Pharmaceutical Affairs Act article 31 and Enforcement for drug safety article 4			
MAH	Local biosimilar guidelines	"Guideline on Evaluation of Biosimilar Products (MFDS 2009)"			
MAH	Date of authorisation/licensing of biosimilar	15 January 2014			
	Reference Bioth	erapeutic Product (RBP) Information			
MAH	Name of the RBP	Herceptin			
MAH	Authorised indications for RBP	Metastatic Breast Cancer			



		Early Breast Cancer
		Metastatic Gastric Cancer
MAH	Pharmaceutical form	Powder for concentrate for solution for infusion. White
		to pale yellow lyophilised powder
MAH	Quantitative composition	150 mg/vial
		440mg/vial
MAH	Route of administration	IV(Intravenous, Infusion)
MAH	Packaging/material	Glass vial
MAH	Package size(s)	1 vial/pack
MAH /	Availability of the RBP	Metastatic Breast Cancer, Early Breast Cancer and
NRA	assessment report	Metastatic Gastric Cancer
	(language)/link	http://www.mfds.go.kr/index.do?searchkey=product_nm∣=11
		6&searchword=허셉틴&cd=191&pageNo=1&seq=14215&cmd=v
		Summary of outcomes
MAH	Comparability exercise to	Physicochemical and biological, in vitro and in vivo
	demonstrate similarity to RBP	functional study
		Toxicological study
		PK/PD study
		Efficacy study (safety and efficacy)
NRA	Availability of full assessment	http://www.mfds.go.kr/index.do?x=0&searchkey=prod
	report (language)/link	uct_nm∣=1176&searchword=허쥬마&cd=191&y
		=0&pageNo=1&seq=19725&cmd=v
MAH	Indications applied for (if	The indications applied for were all authorized for
	different to RBP)	RBP (see section Authorised indications for RBP)
NRA	Authorised indications for	Metastatic Breast Cancer
	biosimilar	Early Breast Cancer
		Metastatic Gastric Cancer

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MAH (Marketing Authorisation Holder) NRA (National Regulatory Authority)



MAH	PART B - SUBMITTED DATA AND REVIEWER SUMMARY           Quality data. Composition of the biosimilar product(s)				
	Trastuzumab 150 mg, Trastuzumab 440	) mg			
MAH	Quality data. State-of-the-art method	ls			
	<ul> <li>Physicochemical Test Methods         <ol> <li>Primary structure : Amino Acid Analysis, Molar Absorptivity, Peptide Mapping(LC-MS, HPLC), N-terminal Sequencing, C-terminal Sequencing, Reduced Mass / Intact Mass</li> <li>High order structure : Disulphide Bonds, Free Thiol Analysis, FTIR, CD, DSC</li> <li>Micro-heterogeneity and Post-translational Forms : IEF, IEC-HPLC, Oligosaccharide Profiling(HPLC), N-linked Glycan Analysis, Monosaccharide Analysis, Sialic Acid Analysis</li> </ol> </li> </ul>				
	<ul> <li>Biological Activity</li> <li>1. In vitro Bioactivity(Anti-proliferation assay)</li> <li>2. HER2 Binding Affinity (ELISA)</li> <li>3. Cell Based Binding Affinity</li> <li>4. C1q Binding Affinity (ELISA)</li> <li>5. FcγRI Binding Affinity (ELISA)</li> <li>6. FcγRIIa Binding Affinity (SPR)</li> <li>7. FcγRIIIa Binding Affinity (SPR)</li> <li>8. FcRn Binding Affinity (SPR)</li> <li>9. ADCC</li> </ul>				
	<ol> <li>FcγRIIIa Binding Affinity (SP.</li> <li>FcRn Binding Affinity (SPR)</li> <li>ADCC</li> </ol>				
NRA	<ol> <li>FcγRIIIa Binding Affinity (SP</li> <li>FcRn Binding Affinity (SPR)</li> </ol>				
NRA	<ul> <li>7. FcγRIIIa Binding Affinity (SP.</li> <li>8. FcRn Binding Affinity (SPR)</li> <li>9. ADCC</li> <li>Quality data assessment outcome</li> <li>Attributes</li> </ul>		Remarks		
NRA	7. FcγRIIIa Binding Affinity (SP.         8. FcRn Binding Affinity (SPR)         9. ADCC         Quality data assessment outcome         Attributes         Primary Structure	R) Comparability	Remarks		
NRA	7. FcγRIIIa Binding Affinity (SP.         8. FcRn Binding Affinity (SPR)         9. ADCC         Quality data assessment outcome         Attributes         Primary Structure         Amino acid analysis	R) Comparability Comparable	Remarks		
NRA	7. FcγRIIIa Binding Affinity (SP.         8. FcRn Binding Affinity (SPR)         9. ADCC         Quality data assessment outcome         Attributes         Primary Structure         Amino acid analysis         N/C-terminal sequence	R) Comparability Comparable Comparable	Remarks		
NRA	7. FcγRIIIa Binding Affinity (SP.         8. FcRn Binding Affinity (SPR)         9. ADCC         Quality data assessment outcome         Attributes         Primary Structure         Amino acid analysis	R) Comparability Comparable	Remarks		
NRA	7. FcγRIIIa Binding Affinity (SP.         8. FcRn Binding Affinity (SPR)         9. ADCC         Quality data assessment outcome         Attributes         Primary Structure         Amino acid analysis         N/C-terminal sequence         Peptide mapping(LC-MS, HPLC)         Molecular weight(LC-MS); Intact,	R) Comparability Comparable Comparable Comparable	Remarks		
NRA	7. FcγRIIIa Binding Affinity (SP.         8. FcRn Binding Affinity (SPR)         9. ADCC         Quality data assessment outcome         Attributes         Primary Structure         Amino acid analysis         N/C-terminal sequence         Peptide mapping(LC-MS, HPLC)         Molecular weight(LC-MS); Intact, reduced	R) Comparability Comparable Comparable Comparable	Remarks		
NRA	7. FcγRIIIa Binding Affinity (SP.         8. FcRn Binding Affinity (SPR)         9. ADCC         Quality data assessment outcome         Attributes         Primary Structure         Amino acid analysis         N/C-terminal sequence         Peptide mapping(LC-MS, HPLC)         Molecular weight(LC-MS); Intact, reduced         Molecular absorptivity	R) Comparability Comparable Comparable Comparable	Remarks		
NRA	7. FcγRIIIa Binding Affinity (SP.         8. FcRn Binding Affinity (SPR)         9. ADCC         Quality data assessment outcome         Attributes         Primary Structure         Amino acid analysis         N/C-terminal sequence         Peptide mapping(LC-MS, HPLC)         Molecular weight(LC-MS); Intact, reduced         Molecular absorptivity         Higher-order structure	R) Comparability Comparable Comparable Comparable Comparable	Remarks		
NRA	<ul> <li>7. FcγRIIIa Binding Affinity (SP. 8. FcRn Binding Affinity (SPR)</li> <li>9. ADCC</li> <li>Quality data assessment outcome</li> <li>Attributes</li> <li>Primary Structure</li> <li>Amino acid analysis</li> <li>N/C-terminal sequence</li> <li>Peptide mapping(LC-MS, HPLC)</li> <li>Molecular weight(LC-MS); Intact, reduced</li> <li>Molecular absorptivity</li> <li>Higher-order structure</li> <li>FTIR, CD, DSC</li> </ul>	R) Comparability Comparable Comparable Comparable Comparable Comparable Comparable Comparable	Remarks		
NRA	7. FcγRIIIa Binding Affinity (SP.         8. FcRn Binding Affinity (SPR)         9. ADCC         Quality data assessment outcome         Attributes         Primary Structure         Amino acid analysis         N/C-terminal sequence         Peptide mapping(LC-MS, HPLC)         Molecular weight(LC-MS); Intact, reduced         Molecular absorptivity         Higher-order structure         FTIR, CD, DSC         Disulfide bond	R) Comparability Comparable Comparable Comparable Comparable Comparable Comparable Comparable Comparable Comparable	Remarks		



NOVEILIDE			IFRE DIUSIIIIIdis W		
	CE-SDS (subunits)	Comparable			
	IEF(isomers)	Comparable			
	IE-HPLC(charge variants)	(Minor)Difference	No effect on biological activity		
	Protein content	Comparable			
	Forced degradation				
	Glycosylation analysis				
	Monosaccharide	Comparable			
	Sialic acid content	Comparable			
	Oligosaccharide profile (HPLC)	Difference	No effect on biological activity		
	N-linked Glycan Analysis	Comparable			
	Biological activity				
	In vitro Bioactivity(Anti-proliferation assay)	Comparable			
	HER2 Binding Affinity (ELISA)	Comparable			
	Cell Based Binding Affinity	Comparable			
	C1q Binding Affinity (ELISA)	Minor difference	Few outlier batches exist		
	FcγRI Binding Affinity (ELISA)	Comparable			
	FcγRIIa Binding Affinity (SPR)	Minor difference	Few outlier batches exist		
	FcγRIIIa Binding Affinity (SPR)	Difference	Comparable in ADCC		
	FcRn Binding Affinity (SPR)	Comparable			
	ADCC(Effector cells: PBMC)	Comparable			
MAH	Mechanism of actionHerzuma (Trastuzumab) is a humanized monoclonal antibody that binds with high affir and specificity to the extracellular domain of HER2.				
MAH	Nonclinical data. In vitro studies				
	<ol> <li>Inhibition of proliferation of I</li> <li>Evaluation of ADCC and CD</li> <li>Comparative analysis for cell</li> <li>HER2 binding affinity to imn</li> <li>Cell based HER2 binding affi</li> <li>Inhibition of growth of HER2</li> <li>FcγRI binding affinity</li> <li>FcγRIIa binding affinity</li> <li>FcγRIIa binding affinity</li> <li>FcRn binding affinity</li> <li>FcRn binding affinity</li> <li>Clq binding affinity</li> <li>Clq binding affinity</li> <li>Tissue cross-reactivity</li> </ol>	C cycle profile and ce nobilised target by H inity	ell cycle controlling proteins		
MAH	Nonclinical data. <i>In vivo</i> studies				
	In vivo pharmacological study Inhibition of tumour xenograft growth	in nude mice			



Novem	per 2015 IPRF Biosimilars WC
	<b>Pharmacokinetics</b> The Pharmacokinetics of Herzuma and Herceptin <sup>®</sup> in the female cynomolgus monkey following single and 4-week repeat dose intravenous administration.
	<ul> <li>Toxicity Study (including TK)</li> <li>Repeat dose toxicity(Comparative design) <ol> <li>Two-week pilot study to compare two intravenous dosing regimens of CT-P6 in the Cynomolgus Monkey</li> <li>Four-week Intravenous Repeat-Dose Toxicity Study in the Cynomolgus Monkey (GLP)</li> <li>13-Week Intravenous Repeat Dose Toxicity Study in the Cynomolgus Monkey (GLP)</li> </ol> </li> </ul>
NRA	Nonclinical data assessment outcome
	1. In vitro studies
	See Quality assessment data outcome.
	In tissue cross reactivity, Herzuma and Herceptin showed same results.
	2. In vivo studies
	<ul> <li>In vivo pharmacological study (Inhibition of tumour xenograft growth in nude mid showed similar result.</li> <li>In repeat dose toxicity, both Herzuma and Herceptin showed similar responses. In 13-week repeat study, reduced heartbeats were observed in both groups at 6 and 1 weeks. No immunogenicity was observed in both groups.</li> </ul>
	- In ADME studies, single and 4 week repeat dose IV studies in monkey showed similar PK profile.
	<ul> <li>CLINICAL STUDIES <ul> <li>include relevant study data from the following (not all may be required) which have been included to demonstrate biosimilarity.</li> <li>Pharmacokinetic, PK</li> <li>Pharmacodynamic, PD</li> <li>Efficacy,</li> <li>Safety,</li> <li>Immunogenicity.</li> </ul> </li> </ul>
MAH	Clinical data. PK studies
	Study Number: CT-P6 1.1Summary of design: Comparative PK study for Double-blind, randomized, Parallel groupphase I/IIb trial Population: Metastatic Breast Cancer patient with active disease(Randomized 170: Herceptin: 85 and Herzuma: 85) Objective and primary endpoint:Demonstration of equivalence PK in terms of area under the curve at steady state (AUCss)between Herzuma and Herceptin in patients with metastatic breast cancer.Dose used: Initial dose of 8 mg/kg followed by 6 mg/kg every 3 weeks, plus paclitaxel (1mg/m²) in 3 week cycles, over 1 year. Length of the Study: Until disease progression, death, discontinuation
	<b>Study Number: CT-P6 1.2</b> Summary of design: Initial PK study for CT-P6 in combination with paclitaxel, phase I trip Population: 8 patients with active disease Metastatic Breast Cancer



	r 2015			IPRF	Biosimilars WG	
	Objective and prim metastatic breast ca mg/kg every 3 wee Length of the study	ancer. Dose used: ks. All patients al	CT-P6 at an initia so received paclita	l dose of 8 mg/kg axel (175 mg/m <sup>2</sup> )	, then at a dose of 6	
	<b>Study Number: C</b> Summary of design		ve PK study for C	T-P6 and Hercep	tin with double-	
	blind, randomized,	•	•	I		
	Population: Active Objective: Demons concentration (C <sub>trot</sub> Herceptin, in patien mg/kg) of CT-P6 o weeks. All the treat	stration of compar $g_{gh}$ ) prior to the sec nts with metastatic r Herceptin and the ted patients also re-	able pharmacokin cond dose, betwee breast cancer (M len a dose of 6 mg eceived Paclitaxel	etics (PK), in term n CT-P6 and the of BC). Dose used: t/kg of CT-P6 or I (175 mg/m <sup>2</sup> ) in 3	ns of trough comparator, an initial dose (8 Herceptin every 3 -week cycles.	
	Length of the study		•	r discontinuation.		
NRA	Clinical data. PK	data assessment o	outcome			
	The primary PK endpoint, the geometric mean of AUC at steady state (AUC <sub>SS</sub> ) at cycle 8 was comparable in the CT-P6 and Herceptin. The 90% CI of geometric mean of AUC <sub>SS</sub> was 93.6% ~ 116.8%, which are within the limit of the acceptance margin ( $80\%$ ~125%). The 90% CI of geometric mean in antibody-negative subset patient was also within the limit of margin.					
MAH	Clinical data. PD studies					
	No specific PD study was conducted due to no relevant biomarker of therapeutic activity.					
NRA	Clinical data. PD data assessment outcome No specific PD study was conducted due to no relevant biomarker of therapeutic activity.					
MAH	Clinical data. Efficacy studies					
	Study Number: CT-P6 3.1					
	Summary of design: Efficacy and safety study of CT-P6 and Herceptin with double-blind, randomized, parallel group, phase 3 trial.					
	Population: Metastatic Breast Cancer patient with active disease (Randomized 366:					
	Herzuma: 244, Herceptin 231 patients) Objective: Demonstration of equivalence of					
	Herzuma and Herceptin, both given in combination with paclitaxel, in terms of efficacy					
	determined by overall response rate (ORR). Equivalence margin was $\pm 15\%$					
	Dose used: CT-P6 or Herceptin at an initial dose of 8 mg/kg, then at a dose of 6 mg/kg					
	every three weeks. All patients also received paclitaxel ( $175 \text{ mg/m}^2$ ) in 3-week cycles.					
	Length of the study					
NRA	Clinical data. Efficacy data assessment outcome					
	The results of the primary endpoints met the equivalence margin both in the full analysis population and per protocol population. The primary endpoint which is overall response rat at 6 month confirmed by ITRC(Independent Tumour Review Committee) was met the equivalence criteria( $\pm 15\%$ )					
		( <u>+</u> 15%)				
		· · ·	S set	P	P set	
		· · ·	S set Herceptin(231)	P CT-P6(236)	P set Herceptin(228)	



Novem	per 2015		essment Summary into		Biosimilars WG
	95% CI	[-0.143:0.036]	[-0.141:	0.041]	
	progression or de similar between	eath due to disease a Herzuma and Herce	e to progression, time to rest t one year, best changes in t otin. Other secondary endporter re not reported at the time o	total targ pints whi	et lesion size were ich are progression
MAH		afety/ Immunogeni mparability margins	city studies (specify popula	ition, dos	se used, length of
	Safety and immunogenicity data were collected from all clinical study; CT-P6 1.1, 1.2, 1.3 and 3.1.				
<u>NRA</u>	Clinical data. Safety/ Immunogenicity data assessment outcome         1. Safety:         The overall adverse event profile collected from all clinical studies was similar for both         Herzuma and Herceptin groups. The percentage of TEAE in Herzuma and Herceptin were         89.3% and 90.9% respectively.         2. Immunogenicity         Immunogenicities of Herzuman and Herceptin from all clinical studies were very low         (<1%).				
MAH	Interchangeschility with the DDD				
	Interchangeability with the RBP           No additional data were provided				
MAH	Additional infor the comparabili	rmation about	As appropriate, if not previ	ously in	cluded.
MAH	Post-authorization measuresRe-examination study in Korea- Period: 2014. 1.15~2018. 1.14				
NRA	Post-authorizati	ion measures assess	sment outcome.		
MAH	- Availability of a relevant inform		As required / appropriate		



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# PART C - REVIEWER CONCLUSIONS

# NRA Conclusions on biosimilarity, approval, interchangeability

The data provided by the Applicant were in line with the local legislation, guidelines and international guidelines.

Quality

All major physicochemical characteristics and biological activities of Herzuma were comparable to those of the reference biotherapeutic product Herceptin.

**Nonclinical** 

No major differences in nonclinical data were observed for Herzuma compared to the reference biotherapeutic product Herceptin .

**Clinical Studies** 

The PK / PD / efficacy studies to demonstrate biosimilarity conducted in metastatic breast cancer patients provided robust evidence of therapeutic equivalence between Herzuma and the reference biotherapeutic product Herceptin

Safety: The ADRs observed with Herzuma were in the same range as the ADRs observed with the reference biotherapeutic product Herceptin.

Immunogenicity: The proportions of patients who developed anti-drug antibodies (ADA) with Herzuma and the reference biotherapeutic product Herceptin were very low (less than 1%).

Risk Management

The risk management plan (or equivalent) was considered to be acceptable.

**Overall Conclusion** 

Satisfactory assurance of biosimilarity was demonstrated using an appropriate comparability exercise.

The biosimilar product Herzuma was considered approvable.