## **EUROPEAN COMMISSION**

DIRECTORATE-GENERAL FOR HEALTH AND FOOD SAFETY

Health systems, medical products and innovation **Medical products: quality, safety, innovation** 

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## TEMPLATE FOR THE QUALIFIED PERSON'S DECLARATION EQUIVALENCE TO EU GMP FOR INVESTIGATIONAL MEDICINAL PRODUCTS MANUFACTURED IN THIRD COUNTRIES

This document provides the template for the certification by the qualified person in the Union that the manufacturing of an investigational medicinal product (IMP) outside of the EU/EEA complies with GMP at least equivalent to the GMP in the Union, as described in the Clinical Trials Regulation 536/2014<sup>1</sup>

The aim is to harmonise this template and hence the information submitted with a request for authorisation of a clinical trial.

Document history:				
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Regulation (EU) No 536/2014 of the European Parliament and of the Council of 16 April 2014 on Clinical Trials On Medicinal Products For Human Use, And Repealing Directive 2001/20/EC

## QUALIFIED PERSON'S DECLARATION EQUIVALENCE TO EU GMP FOR IMP MANUFACTURED IN THIRD COUNTRIES<sup>2</sup> (ARTICLE 63 AND ANNEX I (F) (33) (b) OF REGULATION (EC) 536/2014)

EUCT number(s)	Name of the IMP(s)				
			`	. 2	
Manufacturing and/o	·	norisation (IVII <i>P</i>	() num	ber under w	/nich thi
declaration is made:_					
Part A					
Name of the IMP(s)	Manufacturing site(s)		Activity(-ies) performed at this site		rmed
(Name and	(Name and addres	ss where the is (are)		ding packa	aina.
performed)		()	labelling, storage, testing and release)		
Part B					
☐I confirm that I am	a QP and am author	ised to make th	is decla	ration.	
☐I declare that composite on the basis of:	pliance with GMP at	least equivaler	nt to EU	GMP has bee	en verifie
(i) Audit					
Manufacturing site	e(s)	Auditing part	:y	Date of last	audit
(Name and address	Name and address as in part A)			(completion)	
		<u> </u>			

<sup>&</sup>lt;sup>2</sup> Countries other than EU Member States or contracting states of the European Economic Area (EEA).

<sup>&</sup>lt;sup>3</sup> If no number is issued please state the name of the authorisation holder.

	erformed, please provide a brief justification. at standards at least equivalent to EU GMP
Manufacturing site(s)	Justification
(Name and address as in part A)	
This declaration is submitted by:	
Signature	Date
Name and role	

E.g. assessment of documentation provided by the manufacturer, valid GMP certificate (EudraGMP), etc.