DoC. date: 2021-06-11	Declaration of Compliance	
Template: NLDoc21-004.01	For Sentron products: E2310004 - C2.1 ISFET chip E2310006 – C5.0 ISFET chip E2310400 - C3.0 ISFET chip	Sentron

To whom it may concern,

Sentron B.V. confirms with this Declaration of Compliance that the (semi-) finished end products meet the requirements of:

requirements of:		
Legislation		
In case reportable hazardous substances are present, those are listed in Annex I of this document		
REACH	☑ Not present	
Regulation (EC) No. 1907/2006 on Registration, Evaluation, Authorization and	☐ Present and	
Restriction of Chemical substances (REACH), Candidate list substances, latest list	authorization does/does	
update date: 19 th January 2021	not apply	
RoHS	☑ Not present	
Directive 2011/65/EU (including delegated directive 2015/863/EU) of the European	☐ Present and	
Parliament and of the Council of June 8, 2011 on the restriction of the use of certain	exemptions apply	
Hazardous Substances in electrical and electronic equipment (EEE).		
MDR	☐ N/A, Not for medical	
Substances, marked as Carcinogenic, Mutagenic, Reprotoxic (CMR) 1a/1b under the	purpose	
CLP - Classification, Labelling and Packaging (CLP) Regulation ((EC) No 1272/2008) and	☑ Not present	
human Endocrine Disrupting Chemicals (EDC), classified under EU REACH Candidate	☐ Present	
list, latest update yyyy-mm-dd:		
 The article is not manufactured utilizing: Nanomaterials, meaning a natural, incidental or manufactured material containing particles in an unbound state or as an aggregate or as an agglomerate and where, for 50 % or more of the particles in the number size distribution, one or more external dimensions is in the size range 1-100 nm; Derivatives of tissues or cells of human origin which are non-viable or are rendered non-viable; Tissues or cells of animal origin, or their derivatives, which are non-viable or rendered non-viable. Note: Regulation (EU) 2017/745 on medical devices (MDR), Annex I "General Safety and Performance Requirements", Chapter II "Requirements regarding design and manufacture", point 10.4 deals with the presence of substances that may be released from a medical device. 		
Conflict minerals	☐ Not applicable	
Regulation (EU) 2017/821 of the European Parliament and of the Council of 17 May	☑ Not present	
2017 laying down supply chain due diligence obligations for Union importers of tin,	☐ Present, doesn't origin	
tantalum and tungsten, their ores, and gold originating from conflict-affected and	from conflict or high risk	
high-risk areas	area's	

Unless otherwise stated in Annex I, no other substances, impacted by the applicable legislations mentioned above are present in the listed substance, mixtures and/or articles.

Date: 2021-05-20

With Kind regards,

Sentron B.V. QA-Officer Wendy Goedhart

Annex: No substances to report in Annex

Printed versions are uncontrolled

DoC. date: 2021-06-11	Declaration of Compliance	
Template: NLDoc21-004.01	For Sentron products: E2310004 - C2.1 ISFET chip E2310006 – C5.0 ISFET chip E2310400 - C3.0 ISFET chip	Sentron

Annex 1: Reportable Hazardous Substances

Substance name	CAS No.	MDR*	w/w%	Located where	Measures for safe use:	REACH	RoHS
		(substance marked				authorization ID	exemption ID
		as CMR1a/1b or				(when	(when applicable).
		human EDC)				applicable).	
N/A	N/A	N/A	N/A	N/A	N/A	N/A	N/A

^{*} Relevant for medical customers