


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	

To whom it may concern,

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

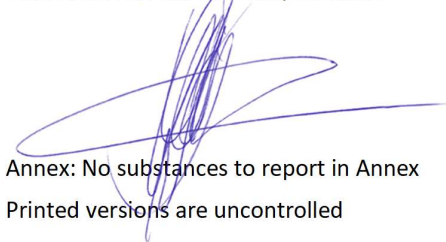
Legislation	
<i>In case reportable hazardous substances are present, those are listed in Annex I of this document</i>	
REACH Regulation (EC) No. 1907/2006 on Registration, Evaluation, Authorization and Restriction of Chemical substances (REACH), Candidate list substances, latest list update date: 19 th January 2021	<input checked="" type="checkbox"/> Not present <input type="checkbox"/> Present and authorization does/does not apply
RoHS Directive 2011/65/EU (including delegated directive 2015/863/EU) of the European Parliament and of the Council of June 8, 2011 on the restriction of the use of certain Hazardous Substances in electrical and electronic equipment (EEE).	<input checked="" type="checkbox"/> Not present <input type="checkbox"/> Present and exemptions apply
MDR Substances, marked as Carcinogenic, Mutagenic, Reprotoxic (CMR) 1a/1b under the CLP - Classification, Labelling and Packaging (CLP) Regulation ((EC) No 1272/2008) and human Endocrine Disrupting Chemicals (EDC), classified under EU REACH Candidate list, latest update yyyy-mm-dd: The article is not manufactured utilizing: <ul style="list-style-type: none"> • 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. <i>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.</i>	<input type="checkbox"/> N/A, Not for medical purpose <input checked="" type="checkbox"/> Not present <input type="checkbox"/> Present
Conflict minerals Regulation (EU) 2017/821 of the European Parliament and of the Council of 17 May 2017 laying down supply chain due diligence obligations for Union importers of tin, tantalum and tungsten, their ores, and gold originating from conflict-affected and high-risk areas	<input type="checkbox"/> Not applicable <input checked="" type="checkbox"/> Not present <input type="checkbox"/> Present, doesn't origin from conflict or high risk 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.

With Kind regards,

Sentron B.V. QA-Officer Wendy Goedhart

Date: 2021-05-20



Annex: No substances to report in Annex
Printed versions are uncontrolled

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

Annex 1: Reportable Hazardous Substances

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

* Relevant for medical customers