



## Declaration of Conformity

Manufacturer **Actim Oy**  
Klovinpellontie 3, FI-02180, Espoo  
Finland  
IVD product **Actim® 1ngeni** (product code 19100AC)

We hereby declare that the above mentioned device complies with the requirements of

98/79/EC *In Vitro* Diagnostic Medical Devices Directive (IVD Directive)  
Finnish National Act 629/2010  
2014/35/EC The Low Voltage Directive (LVD)  
20114/30/EC The Electromagnetic Combability Directive (EMC)  
2011/65/EU + 2015/863/EU + 2017/2102/EU The Restriction of Hazardous Substances Directive (RoHS)

and to the following standards

Standards **SFS-EN ISO 13485:2016, AC 2016, AC 2018**  
Medical devices - Quality Management Systems - Requirement for the regulatory purposes.  
**SFS-EN ISO 14971: 2019**  
Medical Devices. Application of risk management to medical devices.  
**ISO 15223-1:2016**  
Medical devices - Symbols to be used with medical device labels, labelling and information to be supplied Part 1: general requirements.  
**ISO 15223-2:2010**  
Medical devices -- Symbols to be used with medical device labels, labelling, and information to be supplied -- Part 2: Symbol development, selection and validation.  
**SFS-EN ISO 18113-1:2012**  
*In vitro* diagnostic medical devices - Information supplied by the manufacturer (labelling) - Part 1: Terms, definitions and general requirements.  
**SFS-EN ISO 18113-3:2012**  
*In vitro* diagnostic medical devices - Information supplied by the manufacturer (labelling). Part 3: *In vitro* diagnostic instruments for professional use  
**SFS-EN 61010-1:2011**  
Safety requirements for electrical equipment for measurement, control, and laboratory use - Part 1: General requirements  
**SFS-EN 61010-2-101:2017**  
Safety requirements for electrical equipment for measurement, control, and laboratory use - Particular requirements for *in vitro* diagnostic medical equipment  
**SFS-EN 61326-2-6:2013**  
Electrical equipment for measurement, control and laboratory use - EMC requirements - Particular requirements - *In vitro* diagnostic medical equipment  
**SFS-EN 62366-1:2015/A1:2020:en**  
Medical devices - Part 1: Application of usability engineering to medical devices

IVDD  
classification

**General class**

In Espoo 6<sup>th</sup> of May 2021

  
**Magnus Pålsson**  
Managing Director