

 Manufacturer

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 European Authorized Representative

UAB «V&D group»
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Medical device type
Intended purpose
device

IN VITRO diagnostic medical device: Kits for liquid-based cytology "NOVAPREP"
 The kits are intended for use mainly in multiprocessor systems or manually for the preparation of human cellular biomaterial (gynecological and non-gynecological) by liquid-based cytology for further staining with chemical reagents and microscopy, in order to diagnose oncological diseases, including screening for cervical cancer. The intended users are professional personnel of clinical laboratories.

Product name

Kits for liquid-based cytology "NOVAPREP"

Product name	Catalog number
«NOVAPREP NOVASTICK»	NOV016-30

We, the undersigned, hereby declare that the described products meet the essential requirements listed in Directive 98/79/EC of the European Parliament and of the Council dated October 27, 1998. The Technical File required by this Directive is maintained at the Manufacturer's Authorized Representative: UAB "V&D group" 51/412 Paneryu St., LT 03160, Vilnius, Lithuania.

Standards Applied

EN ISO 18113-1:2011; EN ISO 18113-2:2011 (In vitro diagnostic medical devices. Information supplied by the manufacturer (labelling). Terms, definitions and general requirements. In vitro diagnostic reagents for professional use); EN ISO 15223-1:2021 (Symbols to be used with medical device labels, labelling and information to be supplied); EN ISO 13485:2016 +AC:2016 (Medical devices. Quality management systems. Requirements for regulatory purposes); EN 13612:2002 (Performance evaluation of in vitro diagnostic medical devices); EN 23640:2015 (In vitro diagnostic medical devices. Evaluation of stability of in vitro diagnostic reagents); EN ISO 14971:2019 (Medical devices. Application of risk management to medical devices).

Licence No.

LT/CA01/IVD/011/22

Licensee

UAB «V&D group», 51/412 Paneryu St., Vilnius

Start of CE-Marking

11.04.2022

Classification

Other diagnostic medical devices (All devices except Annex II and shelf-testing devices)

Conformity

Assessment Procedure

Annex III of the Directive 98/79/EC (not including section 6)

Director
 Algimed Techno LLC
 (Minsk, Belarus)



Vladislav Liutynski

Date: 11.04.2022