

# DECLARATION OF CONFORMITY

## N° 008/2022

### Manufacturer

Algimed Techno LLC  
 22/1 Logoyskiy trakt, room 309, 220090,  
 Minsk, Belarus  
 Tel.: +375 (29) 893-14-44  
 E-mail: techno@algimed.com

### European Authorized Representative

UAB «V&D group»  
 51/412 Paneryu St., LT 03160,  
 Vilnius, Lithuania  
 Tel.: + 370 (60) 65-31-86  
 E-mail: logistic@vd-group.lt

### 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 Sample Preparation Kit»	NOV020

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/008/22

### 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