

EU DECLARATION OF CONFORMITY

MANUFACTURER PEROXYMED s.r.l.
REGISTERED OFFICE ADDRESS Via Brusuglio 52, 20161, Milano (MI) - Italy

DECLARES UNDER ITS SOLE LIABILITY THAT THE DEVICE:

BD/RDM REGISTRATION 2159868
COMMERCIAL NAME VIRO CLEAN EASY
UDI-DI Base 805750611MEDICALEQB
CND V07 - PRODUCTS FOR CLEANING MEDICAL DEVICES NOT INCLUDED IN OTHER CLASSES

INTENDED USE The devices of the VIRO CLEAN SYSTEM series consist of atomising machines for the diffusion of disinfectant liquids based on hydrogen peroxide of the NTOXY line developed by PEROXYMED s.r.l. These are devices for the atomized sanitization of surfaces and devices in hospitals using "dry fog" technology.

DESCRIPTION The VIRO CLEAN EASY device is an atomizer for chemical products which consists of:
• 1 bottle
• 1 very high speed electric turbine
• 1 union / diffuser
The exit speed of the air from the nozzle that is produced by the electric turbine causes the production of a dry fog whose particles are less than 5 microns.
The dry fog generated in this way is uniformly distributed within the environment to be treated. This process saturates the atmosphere homogeneously with the chemical product used, ensuring optimal application on surfaces, without creating deposits and humidity.
The VIRO CLEAN EASY device has been designed for the use, in the form of dry atomization, of disinfectant chemical products (such as hydrogen peroxide) for the treatment of confined spaces, in the absence of people (after checking the technical data sheet and safety of the product used).

IS ACCORDING TO:

REGULATION (EU) 2017/745

CLASSIFICATION Class I MEDICAL DEVICE (Ann. VIII regola 13)
CONFORMITY ASSESSMENT
PROCEDURE Ann. II e III

AND TO THE FOLLOWING LEGISLATIVE ACTS WHICH PROVIDE FOR THE RELEASE OF AN EU DECLARATION OF CONFORMITY:

Directive 2012/19 / EU on waste electrical and electronic equipment (WEEE / RAEE)

RoHS Directive 2011/65 / EU (See Annex I)

Date and place of issue of the
declaration of conformity

Milan, July 19th 2021

Legal Representative

Angelo Ernesto Rinaldi



Annex I

Declaration of Conformity to EU RoHS

Products are in compliance with Directive 2011/65/EU of the European Parliament and of the Council of 8 June 2011 on the restriction of the use of certain hazardous substances in electrical and electronic equipment (also known as "RoHS Recast"). In addition, this declaration of conformity issued under the sole responsibility of the manufacturer, specifically, products manufactured do not contain the substance is listed in the table below in concentrations greater than the listed maximum value.

This declaration also implements the COMMISSION DELEGATED DIRECTIVE (EU) 2015/863 of 31 March 2015 amending Annex II to Directive 2011/65/EU.

Substance	Maximum Limit %
Lead	0,1
Mercury	0,1
Cadmium	0,1
Hexavalentchromium	0,1
Polybrominatedbiphenyls (PBB)	0,1
Polybrominateddiphenylethers (PBDE)	0,1
Bis(2-ethylhexyl) phthalate (DEHP)	0,1
Butylbenzylphthalate (BBP)	0,1
Dibutylphthalate (DBP)	0,1
Diisobutylphthalate (DIBP)	0,1