



Distributeur France Tel: +33 5 46 97 60 00 @:contact@wesper.com



DECLARATION OF CONFORMITY

The Manufacturer ReSPR Technologies EUROPE declares under its own responsibility that the device

Model and Article No.	Denomination	CND Classification	No Technical File
ReSPR 50 EU40469b	System for treatment and sanitization of air ducts, surfaces, medical apparatus in general, sanitization of disposable devices for general purpose and specialty, clothing, medical disposable and reusable.	V07	NW990-0006

CLASSIFICATION I

Satisfies all applicable dispositions and the essential requirements (Annexe 1) of Directive 93/42CEE on Medical Devices, modified by the Directive 2007/47/CE.

The Medical Device is manufactured also in conformity with the following Technical standards:

CEI EN 60601-1 (CEI 62.5) for the applicable points

Moreover ReSPR Technologies EUROPE is committed to:

- Keep the technical documentation specified at point 3 of Annex VII of Directive 93/42/CEE at the disposal of the Notified Body for a period of five years from the date of manufacture of the product. The aforesaid documentation supports this declaration of conformity;
- Maintain an appropriate system for the monitoring of the device, in the phase successive to that of production, and to apply eventual necessary corrective measures, as prescribed in Annex VII.

It is therefore declared that the above-named device will be put on the market with the Class 1 CE mark, according to the dispositions of Article 17 of Directive 93/42/CEE, modified by the Directive 2007/47/CE.

Date 09 June 2018

Certified by:







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DECLARATION OF CONFORMITY

The Manufacturer ReSPR Technologies Europe declares under its own responsibility that the device

Model and Article No.	Denomination	CND Classification	No Technical File
ReSPR 200 EU40309	System for treatment and sanitization of air ducts, surfaces, medical apparatus in general, sanitization of disposable devices for general purpose and specialty, clothing, medical disposable and reusable.	V07	NW990-0001

CLASSIFICATION I

Satisfies all applicable dispositions and the essential requirements (Annexe 1) of Directive 93/42CEE on Medical Devices, modified by the Directive 2007/47/CE.

The Medical Device is manufactured also in conformity with the following Technical standards:

CEI EN 60601-1 (CEI 62.5) for the applicable points

Moreover ReSPR Technologies Europe is committed to:

- Keep the technical documentation specified at point 3 of Annex VII of Directive 93/42/CEE at the disposal of the Notified Body for a period of five years from the date of manufacture of the product. The aforesaid documentation supports this declaration of conformity;
- Maintain an appropriate system for the monitoring of the device, in the phase successive to that of production, and to apply eventual necessary corrective measures, as prescribed in Annex VII.

It is therefore declared that the above-named device will be put on the market with the Class 1 CE mark, according to the dispositions of Article 17 of Directive 93/42/CEE, modified by the Directive 2007/47/CE.

Date 23 June 2018

Certified by:







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DECLARATION OF CONFORMITY

The Manufacturer ReSPR Technologies Europe declares under its own responsibility that the device

Model and Article No.	Denomination	CND Classification	No Technical File
ReSPR 400 EU40659	System for treatment and sanitization of air ducts, surfaces, medical apparatus in general, sanitization of disposable devices for general purpose and specialty, clothing, medical disposable and reusable.	V07	NW990-0002

CLASSIFICATION I

Satisfies all applicable dispositions and the essential requirements (Annexe 1) of Directive 93/42CEE on Medical Devices, modified by the Directive 2007/47/CE.

The Medical Device is manufactured also in conformity with the following Technical standards:

CEI EN 60601-1 (CEI 62.5) for the applicable points

Moreover ReSPR Technologies Europe is committed to:

- Keep the technical documentation specified at point 3 of Annex VII of Directive 93/42/CEE at the disposal of the Notified Body for a period of five years from the date of manufacture of the product. The aforesaid documentation supports this declaration of conformity;
- Maintain an appropriate system for the monitoring of the device, in the phase successive to that of production, and to apply eventual necessary corrective measures, as prescribed in Annex VII.

It is therefore declared that the above-named device will be put on the market with the Class 1 CE mark, according to the dispositions of Article 17 of Directive 93/42/CEE, modified by the Directive 2007/47/CE.

Date 09 June 2018

Certified by:

Member of CISQ Federation

RINA

ISO 9001:2008

Sistema Qualità Cerifficato





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DECLARATION OF CONFORMITY

The Manufacturer ReSPR Technologies EUROPE declares under its own responsibility that the device

Model and Article No.	Denomination	CND Classification	No Technical File
ReSPR 1000 EU40427	System for treatment and sanitization of air ducts, surfaces, medical apparatus in general, sanitization of disposable devices for general purpose and specialty, clothing, medical disposable and reusable.	V07	NW990-0003

CLASSIFICATION I

Satisfies all applicable dispositions and the essential requirements (Annexe 1) of Directive 93/42CEE on Medical Devices, modified by the Directive 2007/47/CE.

The Medical Device is manufactured also in conformity with the following Technical standards:

CEI EN 60601-1 (CEI 62.5) for the applicable points

Moreover ReSPR Technologies EUROPE is committed to:

- Keep the technical documentation specified at point 3 of Annex VII of Directive 93/42/CEE at the disposal of the Notified Body for a period of five years from the date of manufacture of the product. The aforesaid documentation supports this declaration of conformity;
- Maintain an appropriate system for the monitoring of the device, in the phase successive to that of production, and to apply eventual necessary corrective measures, as prescribed in Annex VII.

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Date 09 June 2018

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DECLARATION OF CONFORMITY

The Manufacturer ReSPR Technologies EUROPE declares under its own responsibility that the device

Model and Article No.	Denomination	CND Classification	No Technical File
ReSPR 2500 EU40425	System for treatment and sanitization of air ducts, surfaces, medical apparatus in general, sanitization of disposable devices for general purpose and specialty, clothing, medical disposable and reusable.	V07	NW990-0004

CLASSIFICATION I

Satisfies all applicable dispositions and the essential requirements (Annexe 1) of Directive 93/42CEE on Medical Devices, modified by the Directive 2007/47/CE.

The Medical Device is manufactured also in conformity with the following Technical standards:

CEI EN 60601-1 (CEI 62.5) for the applicable points

Moreover ReSPR Technologies EUROPE is committed to:

- Keep the technical documentation specified at point 3 of Annex VII of Directive 93/42/CEE at the disposal of the Notified Body for a period of five years from the date of manufacture of the product. The aforesaid documentation supports this declaration of conformity;
- Maintain an appropriate system for the monitoring of the device, in the phase successive to that of production, and to apply eventual necessary corrective measures, as prescribed in Annex VII.

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DECLARATION OF CONFORMITY

The Manufacturer ReSPR Technologies EUROPE declares under its own responsibility that the device

Model and Article No.	Denomination	CND Classification	No Technical File
ReSPR 3001 EU40135g	System for treatment and sanitization of air ducts, surfaces, medical apparatus in general, sanitization of disposable devices for general purpose and specialty, clothing, medical disposable and reusable.	V07	NW990-0007

CLASSIFICATION I

Satisfies all applicable dispositions and the essential requirements (Annexe 1) of Directive 93/42CEE on Medical Devices, modified by the Directive 2007/47/CE.

The Medical Device is manufactured also in conformity with the following Technical standards:

CEI EN 60601-1 (CEI 62.5) for the applicable points

Moreover ReSPR Technologies EUROPE is committed to:

- Keep the technical documentation specified at point 3 of Annex VII of Directive 93/42/CEE at the disposal of the Notified Body for a period of five years from the date of manufacture of the product. The aforesaid documentation supports this declaration of conformity;
- Maintain an appropriate system for the monitoring of the device, in the phase successive to that of production, and to apply eventual necessary corrective measures, as prescribed in Annex VII.

It is therefore declared that the above-named device will be put on the market with the Class 1 CE mark, according to the dispositions of Article 17 of Directive 93/42/CEE, modified by the Directive 2007/47/CE.

Date 09 June 2018

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Sistema Qualità Certificato





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DECLARATION OF CONFORMITY

The Manufacturer ReSPR Technologies EUROPE declares under its own responsibility that the device

Model and Article No.	Denomination	CND Classification	No Technical File
ReSPR 5.000 EU40426	System for treatment and sanitization of air ducts, surfaces, medical apparatus in general, sanitization of disposable devices for general purpose and specialty, clothing, medical disposable and reusable.	V07	NW990-0005

CLASSIFICATION I

Satisfies all applicable dispositions and the essential requirements (Annexe 1) of Directive 93/42CEE on Medical Devices, modified by the Directive 2007/47/CE.

The Medical Device is manufactured also in conformity with the following Technical standards:

CEI EN 60601-1 (CEI 62.5) for the applicable points

Moreover ReSPR Technologies EUROPE is committed to:

- Keep the technical documentation specified at point 3 of Annex VII of Directive 93/42/CEE at the disposal of the Notified Body for a period of five years from the date of manufacture of the product. The aforesaid documentation supports this declaration of conformity;
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