



DECLARATION OF CONFORMITY

The manufacturer: FRANKLAB
Z.A. De L'Observatoire
3 Avenue de Frênes
78180 Montigny-Le-Bretonneux
FRANCE

Hereby declares that, the following product:

RINCE L7

Class I Medical device according to rule 1 of annex VIII of the European Medical Device Regulation 2017/745.

Is manufactured and delivered in accordance with the following regulations:

European Medical Device Regulation 2017/745 (April 5th 2017)
Public Health Code: Part 5 Book II

This statement of conformity is based on Technical File (DT RINCE L7) constituted according to the annex II of the European Medical Device Regulation 2017/745.

The product is placed on the market with following packaging:

RINCE L7 (Basic UDI-ID: 3701387510608P)

- | | |
|-----------------------------|-----------------|
| • 5L Can | Ref. : 1032905 |
| • Low 5L Can | Ref. : 1032905B |
| • 10L Can | Ref. : 1032911 |
| • 20L Can | Ref. : 1032921 |
| • 200L Drum | Ref. : 1032989 |
| • 200L Drum with drumtainer | Ref. : 10329128 |

26th September 2022,

FRANKLAB SAS
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Nicolas VARAY
President

