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## **DECLARATION OF CONFORMITY**

We, undersigned GIMA S.p.A., with operational headquarters in Gessate (MI), Via Marconi 1, and registered office in Milano, Via Tommaso Grossi 2, acting as manufacturer of the medical device:

GIMA Single Registration Number (SRN):

Medical Device ( Trade Name and description)	Code	Basic UDI-DI code
WINTER ROOT ELEVATOR -	60204	802327900L1490020000000UL

Risk class I (Not sterile), according to the Rule 5 Annex VIII of Regulation (EU) 2017/745 (MDR), declares, under its own responsibility, that this medical device:

- comply with essential requirements and dispositions of Regulation (EU) 2017/745 (MDR), as from the Technical File filed at the company;
- common Specifications have not been used for the compliance of the above medical device;

Gessate, 5/28/2021

GIMA S.p.A.
The legal Representative
(Nicola Manzoni)