

EU DECLARATION OF CONFORMITY

REGULATION ON MEDICAL DEVICES (EU) 2017/745, ANNEX IV

ORKLA WOUND CARE AB

SRN: SE-MF-000021041

This EU Declaration of Conformity is issued under the sole responsibility of Orkla Wound Care AB.

We hereby declare that the Medical Device products listed below conform to the Regulation on Medical Devices (EU) 2017/745.

Basic UDI-DI: 7310610105003G

EMDN (CND) code: R03010199

GMDN code: 61326

Intended purpose: The device is intended to be used when giving mouth-to-mouth resuscitation. It functions as a barrier and limits the contact between the resuscitator and the injured person.

REF	Name of product		Risk class
1921	Cederroth	Breathing mask	I
1925	Cederroth	Breathing mask in Key fob	I

Common Specifications applied: No applicable CS available

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