

EC DECLARATION OF CONFORMITY

(Medical Device Directive 93/42/EEC Annex VII)

Orkla Care AB

Hereby declare that the Medical Device products listed below conform to the relevant provisions of the Swedish Law regarding Medical Devices (1993:584) and the current version of LVFS 2003:11, which is the Swedish implementation of the Medical Device Directive 93/42/EEC including amendments to date.

For devices class IIa as verified by our Notified Body, # 0413 according to Medical Device Directive 93/42/EEC, Annex II, EC Certificate A2 41315275-06.

REF	Name of product		Medical Device Class	GMDN
901900	Cederroth	Burn Gel Dressing	IIa	47693
51618575	Cederroth	Burn Gel Dressing	IIa	47693

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