

CE DECLARATION OF CONFORMITY

Plum QuickFix Plaster

Plaster for first aid

We, the manufacturer, of the products covered by this declaration, hereby declare, that the plasters listed below, meet the requirements referred to in Annex I of Regulation (EU) 2017/745 on medical devices and its relevant transposition into national laws of the member states into which we place the device. Plum Safety ApS is responsible for the Declaration of Conformity concerning the medical devices listed below:

Art No	Art Name	UDI-DI
5511	QuickFix Water resistant	05715205000100
5512	QuickFix Elastic	05715205000209
5508	QuickFix Elastic Long	05715205000308
5504	QuickFix Elastic Mini	05715205000407
5518	QuickFix Elastic Micro	05715205000506
5513	QuickFix Detectable	05715205000605
5509	QuickFix Detectable long	05715205000704
5515	QuickFix Alu	05715205000803
5519	QuickFix Alu Micro	05715205000902

With the intended purpose:

Plum QuickFix plasters are used as a first aid product as mechanical barrier and for absorption of exudates.

We guarantee and declare that:

1. The device is a Class I device according to Annex VIII of Regulation (EU) 2017/745, rule 4.
2. The planning and manufacture procedures are conducted in conformity with the Company Quality System, in compliance with the provisions of Annex IX of Regulation (EU) 2017/745.

Assens, 2021-05-18



Regulatory Affairs and Quality Assurance Manager