

Declaration of Conformity



AGFA NV

Septestraat 27, 2640 Mortsel, Belgium

declares that the product

Category: X-ray film, medical, screen

Name/Type:

DRYSTAR DT 1 B	DRYSTAR DT 1 C
DRYSTAR DT 2 B	DRYSTAR DT 2 C
DRYSTAR DT 5 B	DRYSTAR DT 5 C
DRYSTAR DT 10 B	DRYSTAR DT 10 C
DRYSTAR DT 1.000 B	DRYSTAR DT 1.000 C
DRYSTAR DT 5.000 B	DRYSTAR DT 5.000 C
DRYSTAR DT 5.000I B	DRYSTAR DT 5.000I C
DRYSTAR DT 2 B XL	DRYSTAR DT 2 C XL

Application: General Radiology

**complies with the requirements of the 93/42/EEC Directive (Medical Device).
For this Class I device the procedures of Annex VII have been applied in order to
mark the device with the CE-label.**

In case of product changes not accepted in writing by AGFA this declaration will expire.
This declaration is valid maximum for 5 years after the signature date.

Position, Signature & Date

01-07-2018

Paul Merckx
Head of Quality Assurance & Regulatory Affairs
Agfa NV