



Certificate of registration / Registreringsbevis

Manufacturer or Authorised Representative / Tillverkare eller Auktoriserad Representant

Organisation:	Absorbest AB
Organisation no/Organisationsnr:	556547-1413
Postal address/Postadress:	Klintvägen 1, 590 40 KISA

This certificate is valid until / Detta registreringsbevis gäller till: **2005-09-04**

Medical Products Agency hereby confirms that the above mentioned manufacturer/authorised representative has registered the listed medical devices in accordance with the regulations issued by Medical Products Agency

- LVFS 2003:11 on medical devices
- LVFS 2001:7 on medical devices for in vitro diagnostics respectively.

The manufacturer or his representative in the EEA is responsible for the products listed in this certificate and must ensure that they are in accordance with the legislation in force (the Act (1993:584) on medical devices and the regulations issued by Medical Products Agency, LVFS 2003:11 and LVFS 2001:7 respectively)

Läkemedelsverket bekräftar härmed att ovan angivna tillverkare/auktoriserade representant fullgjort sin skyldighet att registrera angivna medicintekniska produkter i enlighet med kraven i Läkemedelsverkets föreskrifter

- LVFS 2003:11, om medicintekniska produkter
- LVFS 2001:7, om medicintekniska produkter för in vitro diagnostik

Tillverkaren eller hans representant inom EES, svarar för att de produkter som ingår i detta bevis uppfyller kraven i gällande författningar (lag (1993:584) om medicintekniska produkter samt Läkemedelsverkets föreskrifter LVFS 2003:11 respektive LVFS 2001:7).

Products / Produkter

PID	Product/Produkt	Article No/Artikelnr	Product class /Produktklass	GMDN
28600	DryMax 2,4		MD, Klass I	35008
30369	DryMax neo		MD, Klass I	35008
32384	DryMax XL		MD, Klass I	35008