Product name:

Valid from: HSA/LC 14-01-2021 Replaces: HSA/LC N/A

CVR NO.: DK15517085 VAT NO. EXPORT: DK29127204

Batch Release Certificate

Dextran 500 Ultra

Specification No.: Batch No.: Manufacturing sites: DKMA* No.: GMP certificate No.: FDA establishment No.: FDA facility classification: EDQM* certificate No.:		50014			
		XXXXXX			
		Pharmacosmos A/S, Roervangsvej 30, DK-4300 Holbaek, Denmark 254629 N/A N/A N/A N/A			
Method:	Parameter:		Results of analysis:	Limits:	
_	Description:		Complies	White or almost white powder	
DF001	Loss on drying (105°C, 5h), % w/w:		X.X	** ≤7.0	
DF030	Absorbance (at 375 nm, 10% sol., 1 cm)		X.XX	≤0.12	
DF009	Specific rotation, (+/-) °:		+X	+195 – +201	
DF019	Nitrogen, ppm:		X	<110	
DF046	Average molecular mass (laser), Mw:		X,XXX	450,000 – 550,000	
DF046	Mw/Mn (laser):		X.X	<3.0	
DF007	Sulphated ash, % w/w:		X.X	≤0.3	
DF022		amination, cfu/g:	Complies	≤100	
		ation is authentic and accurate		dextran has been manufactured, including	
Date (dd.mm.yyyy):		Heidi Skjødt Andersen, M.Sc. Pharm, Quality Control			
		r	isiai Snjeut Aliusisell, IVI.SC. Plidi	iiii, Quality Control	

Page 1 of 1

PHARMACOSMOS A/S ROERVANGSVEJ 30 DK-4300 HOLBAEK DENMARK TELEPHONE: +45 59 48 59 59
TELEFAX: +45 59 48 59 60
E-MAIL: info@pharmacosmos.com
WEB-SITE: www.pharmacosmos.com