Valid from: HSA/LC 17-11-2020 Replaces: HSA/LC 18-04-2018

Batch Release Certificate

Product name: Specification No.: Batch No.: Manufacturing date: Retest date (5 years): Manufacturing sites: DKMA* No.: GMP certificate No.: FDA establishment No.: FDA facility classification: EDQM* certificate No.:		A0069 XXXXXX mmmm_yyyy mmmm_yyyy Pharmacosmos A/S, Roervangsvej 30, DK-4300 Holbaek, Denmark 254629 DK API-H 00125319, DK API-V 00125319 FEI 3002807874 Acceptable Certificates for Dextran 1, 40, 60 & 70 Ph.Eur. available							
					Method:	Parameter:		Results of analysis:	Limits:
					EP	Appearance of solution:		Complies	Clear and colorless
					DF005	Acidity or Alkalinity:		Complies	Complies
EP	Specific rotation, (+/-) °:		+x	+191 – +201					
LI030-1	Average molecular mass, Mw:		x,xxx	17,000 – 23,000					
EP	Nitrogen containing substances, ppm N:		x	≤110					
EP	Loss on drying (105°C, 5h), % w/w:		x.x	≤7.0					
EP	Sulphated ash, % w/w:		x.x	≤0.3					
EP	Bacterial endotoxins, IU/g:		Complies	<25					
EP	Microbial contamination, cfu/g:		Complies	≤100					
*) EDQM refers to We confirm that i	o 'European Directorato no class 1, class 2 and	class 3 solvent, cf. EP 5.4 and l	d Healthcare'. DKMA refers to 'Danish JSP <467> Residual solvents, is used	• ,					
packaging and qual	the above information ity control at the abo	ove mentioned site in full com	pliance with the GMP requirement	ngredient has been manufactured, including is for active starting materials and with the d found to be in compliance with GMP.					

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Date (dd.mm.yyyy):

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