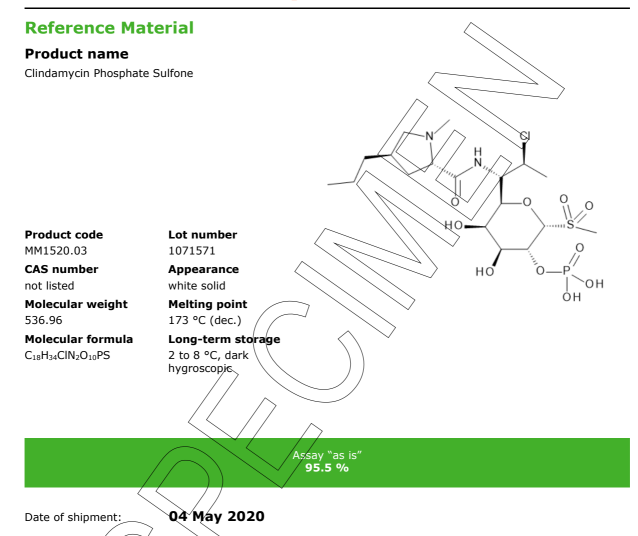


Certificate of Analysis

ISO 9001



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Producer confirms that this reference material (RM) meets the specification detailed on this Certificate of Analysis for **one year** from the date of shipment, provided the substance is stored under the recommended conditions unopened in the original

Release by:	Date of Release:	\mathcal{O}	Product Release
Dr. Sabine Schröder	Luckenwalde, 06 Apr 2020	Janol	Product Release

container.

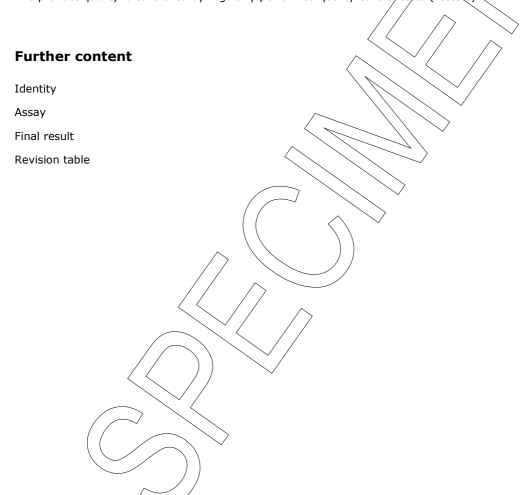


Product information

For laboratory use only. Not suitable for human or animal consumption.

Before usage of the RM, it should be allowed to warm to room temperature. No drying required, as the certified value is already corrected for the content of water and other volatile materials.

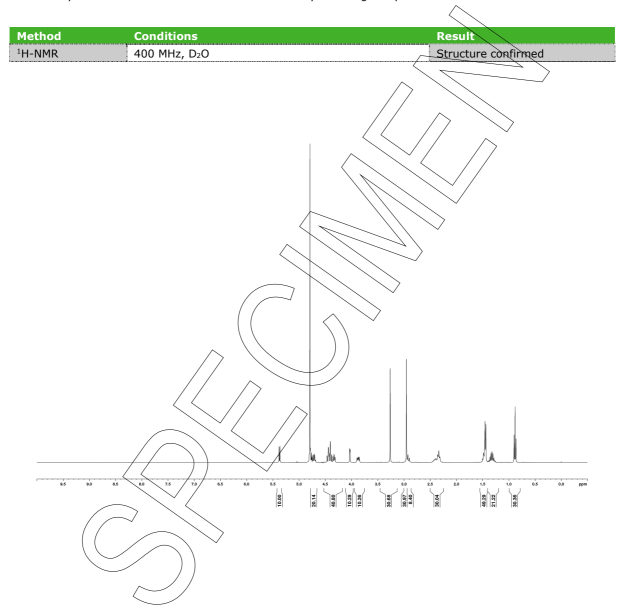
The product quality is controlled by regularly performed quality control tests (retests).





Identity

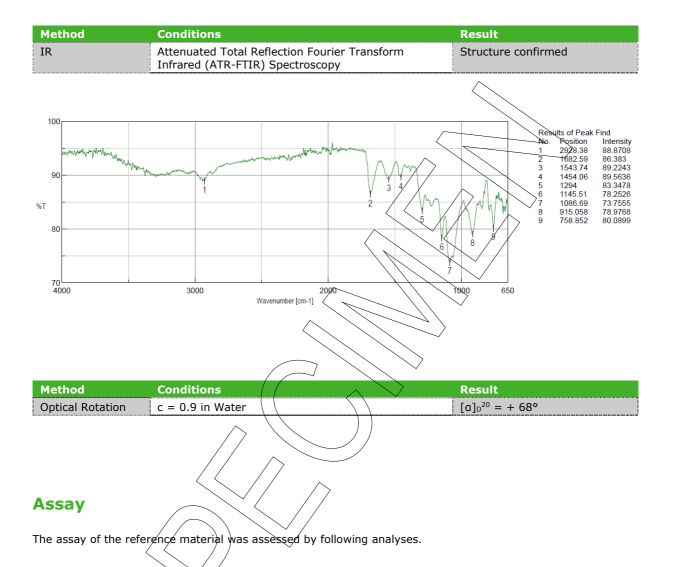
The identity of the reference material was established by following analyses.





Method	Conditions	Result
MS	3.5 kV ESI+; capillary temperature: 269 °C Theoretical value: 559.12525	Structure confirmed
10	29 217.06935 326.10420 365.13538 47.29236 537.14288 581.10693 7	703.20960 777.22797 828.19190 925.26566 700 800 900 1000



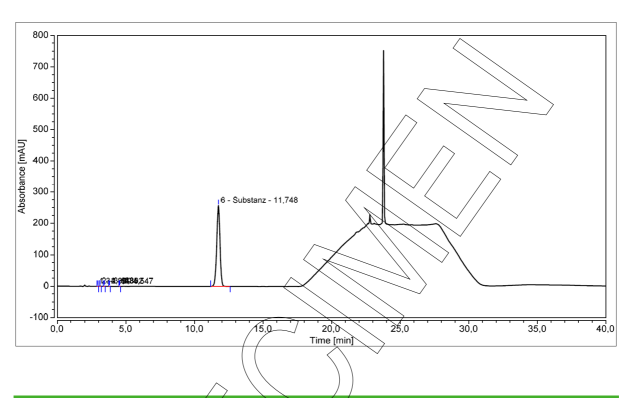


Purity by High Performance Liquid Chromatography (HPLC)

HPLC conditions:	
Column	Hypersil Gold C18; 5 μm, 150 x 4.6 mm
Column temperature	40 °C
Detector	DAD, 200 nm
Injector	Auto 7 μl; 0.193 mg/ml in Water
Flow rate	1.0 ml/min
Phase A	Water, 0.1 % H₃PO₄
Phase B	Acetonitrile, 0.1 % H₃PO₄
Gradient program	0-15 min A/B 85/15 15-20 min A/B to 20/80 20-25 min A/B 20/80 25-28 min A/B to 85/15 28-40 min A/B 85/15 (v/v)



HPLC chromatogram and peak table



Area percent report - sorted by signal				
Pk #	Retention time	Area	Area %	
1	2.940	/0.ø21	0.03	
2	3.133	0.069	0.10	
3	3,435	0.024	0.03	
4	3.802	0.018	0.03	
5 ((4.547	0.005	0.01	
6	11.748	71.401	99.81	
Totals		71.539	100.00	

The content of the analyte was determined as ratio of the peak area of the analyte and the cumulative areas of the purities, added up to 100 %. System peaks were ignored in calculation.

esult (n = 6)	99.84 %; SD = 0.04 %	
	22101 707 02 0101 70	



Volatile content

Water content		
Method	Karl Fischer titration	
Result (n = 3)	4.32 %; SD = 0.09 %	

Residual solvents		_	
Method	¹ H-NMR		
Result (n = 1)	No significant amounts of residual	sølν	

Final result

Assay "as is": 95.53 %

The assay "as is" is assessed by 100% method (mass balance) and is equivalent to the assay based on the not anhydrous and not dried substance respectively.

The calculation of the 100% method follows the formula:

Assay (%) = (100 % - volatile contents (%)) * Purity (%) 100 %

Volatile contents are considered as absolute contributions and purity is considered as relative contribution. Inorganic residues are excluded by additional tests.

Revision table

Revision	Date/	Reason for revision
00	06 Apr 2020	Release of the Certificate of Analysis – initial version

Product warranties for the RM are set out in the terms and conditions of purchase.