

# **Certificate of Analysis**

# **Reference Material**

### Product name

5-Chloropyrimidine-2,4(1H,3H)-dione (5-Chlorouracil)

Ы **Product code** Lot number MM0593.05-0025 1030651 **CAS number** Appearance 1820-81-1 white solid Molecular weight Melting point (DSC) 146.53 319 °C Molecular formula Long-term storage 2 to 8 °C, dark C<sub>4</sub>H<sub>3</sub>CIN<sub>2</sub>O<sub>2</sub> Assay "as is" 100.00 % 06 Sep 2022 Date of shipment: Producer confirms that this reference material (RM) meets the specification detailed on this Certificate of Analysis for two years from the date of shipment, provided the substance is stored under the recommended conditions unopened in the original container.

 Release by:
 Date of Rélease:

 Dr. Sabine Schröder
 Luckenwalde, 22 Aug 2019

Organisation certified to ISO 9001 | DQS 102448 and GMP (EXCIPACT)

Producer: LGC GmbH Louis-Pasteur-Str. 30 D-14943 Luckenwalde Germany www.lgcstandards.com Page 1/7



### **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)

### **Further content**

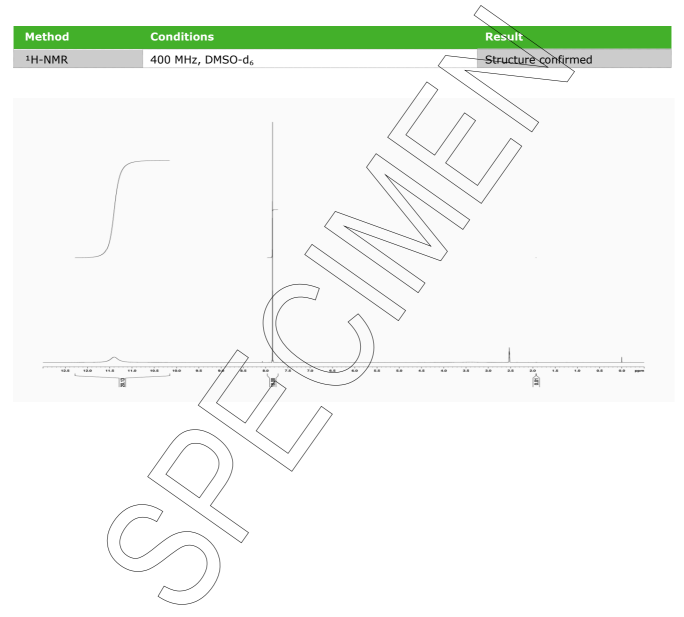
Identity Assay Final result

Revision table



# Identity

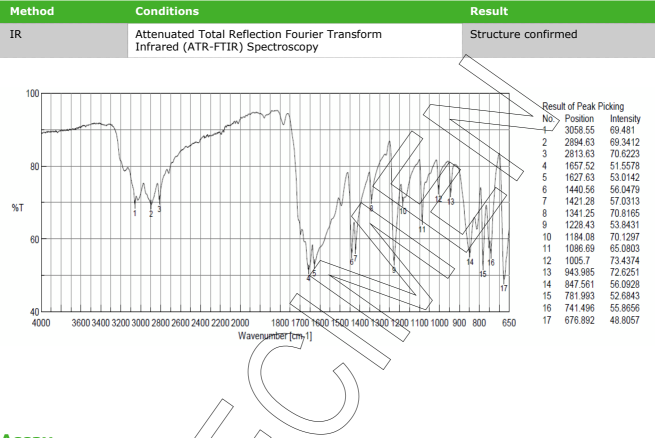
The identity of the reference material was established by following analyses.





Method	Conditions	Result
MS	3.5 kV ESI+; capillary temperature: 269 °C Theoretical value: 146.99558	Structure confirmed
Apex MM0593.05-FA	A163370 Scan: #41 RT: 0.42 min NL: 1.15E+007 Apex	FTMS +p ESI Full ms [80.00-1000.00]
90 - 80 - 70 -		
60- 50- 40-	214.00342	
30- 100.07576 20- 10- 102.03381 134	.03922 200.02428 282.96779 3/8.9651	7 377 92155
0-06089 114 09142	196/15393 199.48/94 248.00132 284.96705 328.9	7337 414.26964 m/z
80 100		<u>350 503</u>





# Assay

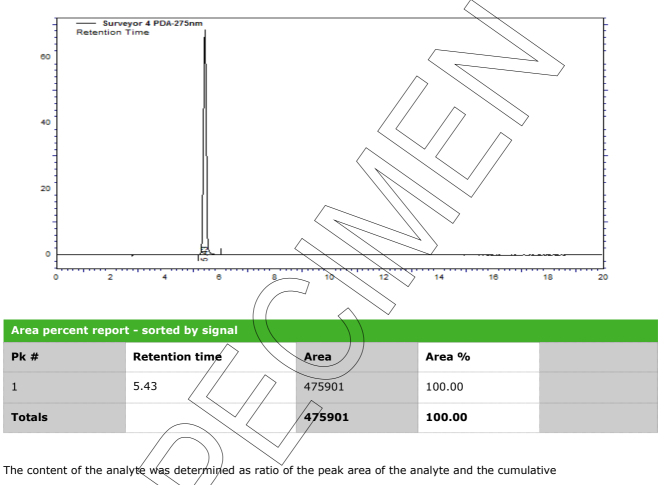
The assay of the reference material was assessed by following analyses.

## Purity by High Performance Liquid Chromatography (HPLC)

HPLC Conditions:	
Column	Discovery HS F5; 3 µm, 150 x 4.0 mm
Column temperature	40 °C
Detector	DAD, 275 nm
Injector	Auto 1.00 $\mu l;$ 0.037 mg/ml in Acetonitrile/Water 50/50 (v/v)
Flow rate	1.0 ml/min
Phase A	Water, 0.1 % H <sub>3</sub> PO <sub>4</sub>
Phase B	Acetonitrile, 0.1 % $H_3PO_4$
Gradient program	A/B 97/3 (v/v)



### HPLC chromatogram and peak table



areas of the purities, added up to  $100^{\circ}$ %. System peaks were ignored in calculation.

Result (n = 6)

100.00 %; SD < 0.01 %

### Volatile content

Water content		
Method	Karl Fischer titration	
Result	No significant amounts of water were detected (< $0.05$ %).	



Residual solvents	
Method	<sup>1</sup> H-NMR
Result (n = 1)	No significant amounts of residual solvents were detected (< 0.05 %).
Final result Assay "as is": 100.0	
	100% method (mass balance) and is equivalent to the assay based on the not
anhydrous and not dried substan	ce respectively.
The calculation of the 100% met	hod follows the formula:
Assay (%) = (100 % - volatile co	ontents (%)) *
Volatile contents are considered a	as absolute contributions and purity is considered as relative contribution.
Inorganic residues are excluded l	
Revision table	
Revision Date	Reason for revision
00 22 Aug 2019 Product warranties for the RM are	Release of the Certificate of Analysis - initial version e set out in the terms and conditions of purchase.