

of Analysis

Reference Material - Primary Standard

Product Name: Methedrone-D₃ Hydrochloride 1.0 mg/ml in Methanol (as free base)

Catalogue Number: LGCAMP1343.04-01

Lot Number: 14565

CAS Number: 1231710-62-5

Molecular Formula: C11H12D3NO2 HCI

232.72 Molecular Weight: Solvent: Methanol

Not less than 1 ml Volume per Ampoule:

2 to 8 °C, dark Long-term Storage:

Expiry Date: June-2016

Intended Use: The primary aim of this material is for identification, calibration and quantification.

Component	Concentration ("free base")	Uncertainty
see product name	1.000 mg/ml ²	$U = 0.004 \text{ mg/ml}^{-3}$
Uncertainty of the concentration is expressed as an expanded uncertainty in accordance with ISO 17025 and Guide 34 at the		

about 95 % level of confidence using a coverage factor of k = 2 and has been calculated by statistical analysis of our production system and incorporates uncertainty of the purity, material density and balance and weighing technique. Concentration based on material weighings and material purity factor (assay of the neat material).

The solution's concentration and homogeneity are verified by independent method.

LGC certifies that this standard meets the specification stated in this certificate and warrants this product to meet the stated acceptance criteria through the retest date when stored unopened as recommended. Product should be used shortly after opening to avoid concentration changes due to evaporation. Warranty does not apply to ampoules stored after opening.

Release Date:

Luckenwalde, August 2012

Signed:

Dr. Sabine Schröder Unit for Reference Materials

Н

x HCI

LGC Quality - ISO Guide 34:2009 | ISO/IEC 17025:2005 | ISO 9001:2008



¹ Ampoules are overfilled to ensure a minimum 1 ml volume fill. We advise laboratories to use measured volumes of this

standard solution before diluting to the desired concentration.

The value is based on the results of analytical techniques, which calibration and verification was carried out with standards traceable to SI-units. The value is expressed on a "free base" basis.

The concentration with its uncertainty is valid in the range between 19 °C and 25 °C.

The identity is verified by data from international scientific literature.

Gravimetrically prepared using qualified balances calibrated annually by accredited calibration service. Calibration verification

performed daily prior to use utilizing weights traceable to SI via other mass standards.

The uncertainty "U" is the expanded uncertainty estimated in accordance with the Guide to the Expression of Uncertainty in Measurement (GUM). It is corresponding to a level of confidence of about 95 %. Standard uncertainties are indicated with "u".



Additional information	
Concentration ("as is")	1.186 mg/ml

Verification of Concentration and Homogeneity		
Lot Number	Verified Concentration (mg/ml) Result Acceptance Criteria	% RSD - Homogeneity Result Acceptance Criteria
14565	0.994 ± 3 %	0.902 ≤ 3 %
Concentration verified by HPLC		

Solution Standard Assay Parameters

External Calibration (100 % amount)

Analysis Method HPLC

Column: Hypersil Gold (C18), 5 µm, 150 x 4.6 mm Number of Measurements: 6

Injector: Auto; 2 µl; 1.0 mg/ml in Methanol

Flow: 1.0 ml/min, 40 °C Detector: 290 nm

Conditions: mob. Phase A: Water + 0.1 % H₃PO₄,

mob. Phase B: Acetonitrile + 0.1 % H_3PO_4 0-8 min A/B 90/10, 8-11 min A/B to 50/50,

11-13 min A/B to 90/10, 13-20 min A/B 90/10 (v/v)

Neat Material Data

Product Name: Methedrone-D₃ Hydrochloride

CAS Number: 1231710-62-5 Molecular Formula: $C_{11}H_{12}D_3NO_2$ HCI

Molecular Weight: 232.72 Compound Lot: 14524

Test	Method	Result
Melting Point (°C)*	SOP 06-010	206 °C
¹ H-NMR Spectrum*	SOP 06-053	conform / complies to structure
IR Spectrum*	SOP 06-036	conform / complies to structure
Mass Spectrum (ESI)*	SOP 06-022	conform / complies to structure
Isotopic Purity by HRMS		0.07 % D ₂ , 99.93 % D ₃
Assay by quantitative NMR (free base)*	Quant. NMR	83.79 %
T	0.05.0/ /	1050/1 1 (() 1

The expanded uncertainty according to the assay is $U=0.35\,\%$ (about 95 % level of confidence using a coverage factor of k=2).

Assay by quantitative NMR	("as is")*	99.35 %
---------------------------	------------	---------

^{*:} Validated method performed by ISO/IEC 17025 accredited testing lab. Purity factor does not include adjustment for chiral and/or isotopic purity.

The assay of the neat material is verified by the 100 % method using HPLC, corrected with water

(KFT) and residual solvents.

LGCAMP1343.04-01 Lot Number 14565



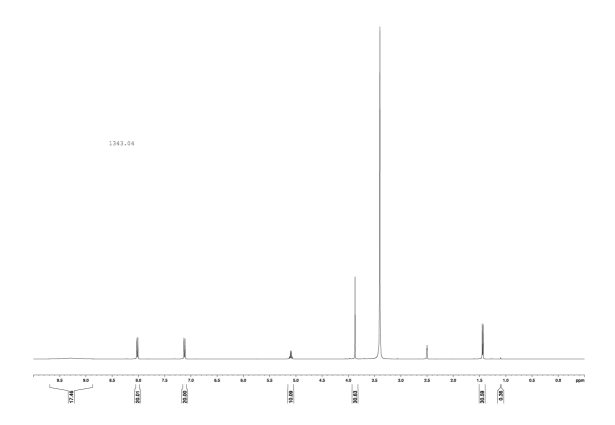
I. Identity

The identity of the reference substance (neat material) was established by the following analyses.

la. ¹H-NMR Spectrum

Conditions: 400 MHz, DMSO-d₆

The structure is confirmed with the signals of the spectrum and their interpretation.

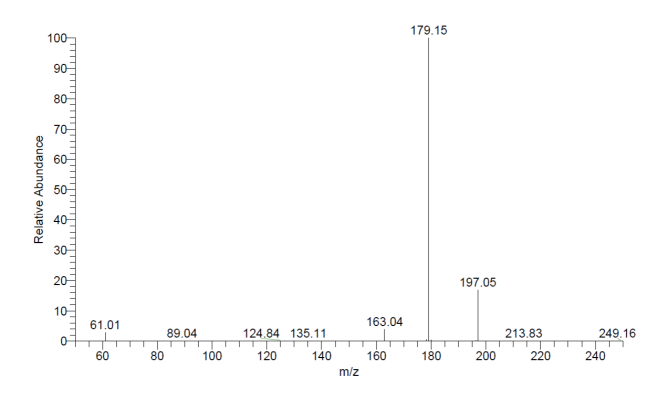






lb. Mass Spectrum

Method: 4.5 kV ESI; vaporization temperature: 200 °C, direct inlet



m/z	fragments (M = free base)
197.05	[MH]
179.15	[MH – CD ₃]
163.04	[M – NHCD ₃]
61.01	[C ₃ H ₅ D ₃ N]

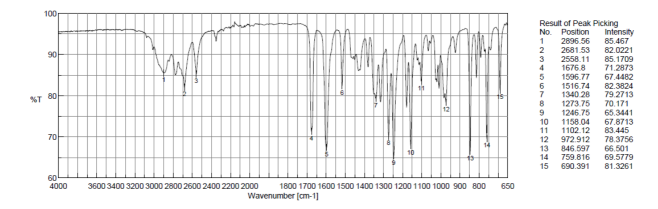
The signals of the mass spectrum and their interpretation are consistent with the structural formula.





Ic. IR Spectrum

Method: attenuated total reflection fourier transform infrared (ATR-FTIR) spectroscopy



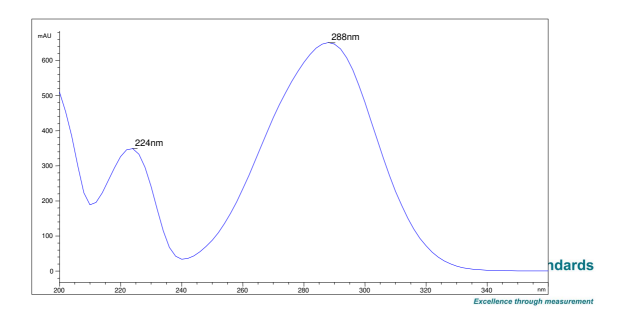
The signals of the IR spectrum and their interpretation are consistent with the structural formula.

ld. Melting Point

206 °C

le. UV Spectrum

Method: HPLC (DAD-detection)

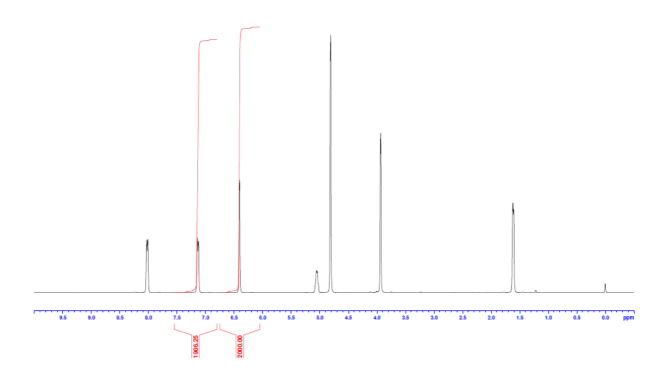


LGCAMP1343.04-01 Lot Number 14565



II. Assay by quantitative NMR spectroscopy

The assay of the reference substance was established by quantitative NMR spectroscopy using D_2O as the solvent and with Maleic acid (certified reference material, signal 6.05 - 6.75 ppm, 2 H) as internal standard.



Results:

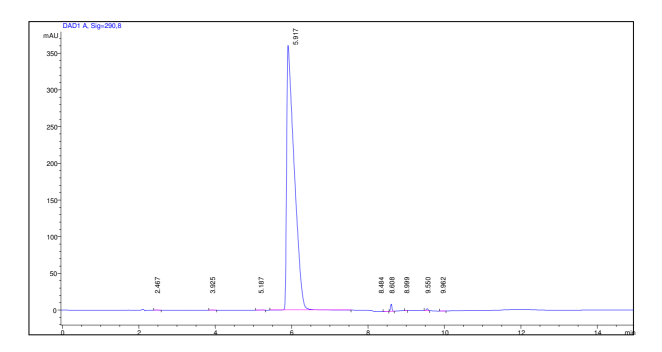




III. Purity

Illa. High Performance Liquid Chromatography (HPLC)

The purity of the reference substance (neat material) was analysed by high performance liquid chromatography (HPLC).



Area Percent Report - Sorted by Signal

Pk#	Retention Time	Area	Area %	
1	2.47	0.35	0.01	
2	3.93	0.38	0.01	
3	5.19	0.68	0.01	
4	5.92	4982.52	99.27	
5	8.48	0.69	0.01	
6	8.61	26.01	0.52	
7	9.00	0.67	0.01	
8	9.55	7.71	0.15	
9	9.96	0.39	0.01	
Totals		5019.41	100.00	

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

Excellence through measurement



Water

HPLC Conditions:

Column: **Conditions: Detector:** Injector: Hypersil Gold (C18) 1.0 ml/min, 40 °C DAD Auto

5 μm, 150 x 4.6 mm 0-5 min Water/Acetonitrile 90/10 290 nm 5 μl; 0.422 mg/ml in

> 5-7 min Water/Acetonitrile to 20/80 7-9 min Water/Acetonitrile to 90/10 9-15 min Water/Acetonitrile 90/10 (v/v);

0.1 % H₃PO₄

Results:

Arithmetic mean (n=3) 99.27 %

IIIb. **Water Content**

Method: coulometric Karl Fischer titration

Results:

Arithmetic mean (n=3) 0.18 % (mass fraction)

IIIc. **Residual Solvents**

Method: 1H-NMR

Result: 0.15 % tert-Butyl methyl ether

IV. Stability and Homogeneity

Accelerated stability studies indicate no significant instability. The given validity period is based on this data. This is backed up by historical data over the range of several years for the neat substance. Homogeneity assured by validated process of preparation (incl. ampoulation), verified by homogeneity testing (HPLC). **Standards**

Excellence through measurement



V. Further Information

General

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

This material conforms to the characteristics of a primary standard as described within ISO Guide 30 (Terms and definitions used in connection with reference materials).

The certified values quoted in this certificate are LGC's best estimate of the true values within the stated uncertainties and based on the techniques described in this certificate.

Handling of the RM

Before usage of the RM, it should be allowed to warm to room temperature. The concentration with its uncertainty is guaranteed in the range between 19 °C and 25 °C. The uncertainty accounts for the temperature-dependent density in this range.

Quality Control Assessment

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

Revision	Date	Reason for Revision
00	August 2012	Release of the Lot – initial version
01	August 2013	Copyright added

