

Certificate of Analysis – Certified Reference Material

N,N-DIETHYLETHANE-1,2-DIAMINE (Metoclopramide Impurity E)

Product no.: PHR2286-100MG

Lot no.: LRAC3776

Description of CRM: Clear Liquid

Expiry date: 31 August 2023

Storage: Refrigerator/Protect from Light

Certificate version: LRAC3776.2 (Note: Certificates may be updated due to

Pharmacopeial Lot Changes or the availability of new data. Check our website at: www.sigma-aldrich.com for the most

current version.)

Chemical formula: $C_6H_{16}N_2$ Molecular mass:116.2CAS No.:100-36-7

	CH ₃	
H ₂ N	✓_N_	_CH₃

Analyte	Certified Purity \pm associated uncertainty U , $U=k \cdot u$ ($k=$) (Mass Balance/basis)
Metoclopramide Impt E	99.8 % Ucrm = ± 0.4 %, k = 2.0 (Mass Balance / anhydrous basis)

Metrological traceability: Traceable to the SI and higher order standards from NIST through an unbroken

chain of comparisons. Additional traceability to Primary Standards is established through comparative assay determinations. See "Details on metrological

traceability" on page 2.

Measurement method: Where applicable, the certified value is based on a purity determination by mass

balance. See "Certification process details" on page 3.

Intended use: Intended for R&D and Analytical Use only. Not for drug, household or other uses.

Minimum sample size: 12 mg

Instructions for handling Determine water content titrimetrically at time of use and use on the anhydrous

and correct use: basis. The internal pressure of the container may be slightly different from the atmospheric pressure at the user's location. Open slowly and carefully to avoid

dispersion of the material. Attachment of a 20 mm aluminum crimp seal

recommended for unused portions.

Health and safetyAll chemical reference materials should be considered potentially hazardous and should be used only by qualified laboratory personnel. Please refer to the Safety

Data Sheet for detailed information about the nature of any hazard and appropriate

precautions to be taken.

Accreditation: Sigma-Aldrich RTC is accredited by the US accreditation authority ANAB as a

registered reference material producer AR-1470 in accordance with ISO 17034.

Certificate issue date: 05 April 2022



ISO 17034 AR-1470 ay Ome

[Andy Ommen; Quality Control]

Sham Statler

Shawn Stetler- QA Manager



Packaging:

100mg in amber ampule

Details on metrological traceability:

This standard has been gravimetrically prepared using balances that have been fully qualified and calibrated to ISO 17025 requirements. All calibrations utilize NIST traceable weights which are calibrated externally by a qualified ISO 17025 accredited calibration laboratory to NIST standards. Qualification of each balance includes the assignment of a minimum weighing by a qualified and ISO 17025 accredited calibration vendor taking into consideration the balance and installed environmental conditions to ensure compliance with USP tolerances of NMT 0.10% relative error. Fill volume to predetermined specifications is gravimetrically verified throughout the dispensing process using qualified and calibrated balances. Further traceability to a corresponding Primary Standard may be achieved through a direct comparison assay. Where a Primary Standard

Associated uncertainty:

Uncertainty values in this document are expressed as Expanded Uncertainty (U_{CRM}) corresponding to the 95% confidence interval. U_{CRM} is derived from the combined standard uncertainty multiplied by the coverage factor k, which is obtained from a t-distribution and degrees of freedom. The components of combined standard uncertainty include the uncertainties due to characterization, homogeneity, long term stability, and short term stability (transport). The components due to stability are generally considered to be negligible unless otherwise indicated by stability studies.

is available, the assay value will be included in the specified section of the COA.

TRACEABILITY COMPARISON

Comparative chromatographic identification analysis demonstrates direct traceability to Pharmacopeial standards through compendial method analysis

TRACEABILITY COMPARISON vs. USP REFERENCE STANDARD (anhydrous basis)

MilliporeSigma Lot LRAC3776 vs. EP Batch 2.0

Method: HPLC (ref.: Metoclopramide, Current Compendial Monographs)

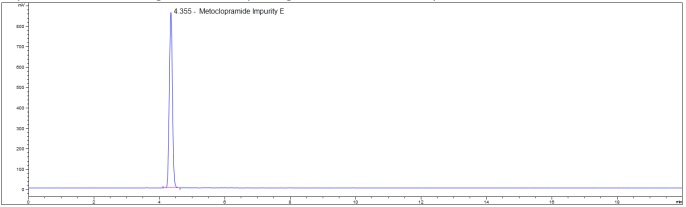
Column: Primesep 200, 4.6 x 150 mm, 5 um particle size

Mobile Phase: 0.1% Trifluoroacetic acid in water: 0.1% Trifluoroacetic acid in Acetonitrile (80:20)

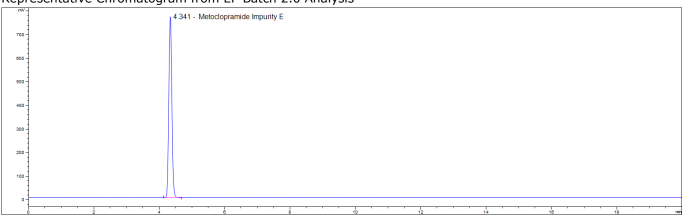
Flow Rate: 1.0 mL/min Column Temperature: 30 °C Injection Volume: 20 µL

Detector: ELSD

Representative Chromatogram from MilliporeSigma Lot LRAC3776 Analysis



Representative Chromatogram from EP Batch 2.0 Analysis



Certification process details:

The certified purity is determined by mass balance and calculated as

$$\% \ Purity = \left(\frac{(100-TCI)}{100} * \frac{(100-LOD)}{100} * \frac{(100-H2O)}{100} * \frac{(100-ROI)}{100} * \frac{(100-ROI)}{100} * \frac{(100-RS)}{100}\right) * 100\%$$

- TCI = Total Chromatographic Impurities
- LOD = Loss on Drying
- H₂O = Water content determined by Karl Fischer analysis
- ROI = Residue on Ignition
- RS = Residual Solvents

Methods for impurity determination may be added or deleted as required. The following techniques are applied:

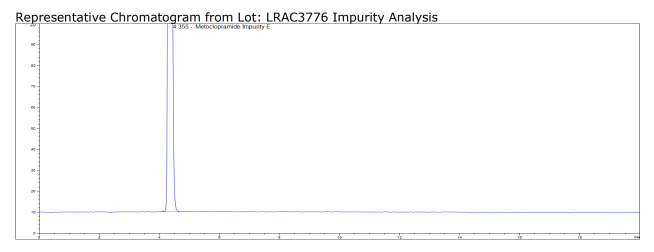
CHROMATOGRAPHIC IMPURITY ANALYSIS

METHOD: HPLC (ref.: Metoclopramide, Current Compendial Monographs)

See Traceability Comparison

Impurities Detected:

None



RESIDUAL SOLVENTS

Method: GC-MS Headspace (ref.: Adapted from Residual Solvents USP <467>)

Column: SPB-624 Carrier gas: He Flow: 1.2 mL/min Split Ratio: 1:5

Injection/Temperature: 1 mL/220 °C

Temperature Program: 40 °C for 5 min, 8 °C/min to 200 °C, hold 5 min

Solvents Detected:

Triethylamine: 0.176 %

WATER DETERMINATION

Method: Karl Fischer (ref.: Current Compendial Monographs)
Mean of three measurements, Water Content = **4.68** %

RESIDUE ANALYSIS

Method: Sulfated Ash (ref.: Current Compendial Monographs)

Sample Size: ~ 75 mg

Mean of three measurements, Residue = None

CERTIFIED PURITY BY MASS BALANCE

99.8 % $U_{crm} = \pm 0.4$ %, k = 2.0 (anhydrous basis)

Homogeneity assessment:

Homogeneity was assessed in accordance with ISO Guide 35. Completed units were sampled using a random stratified sampling protocol. The results of chemical analysis were then compared by Single Factor Analysis of Variance (ANOVA). The uncertainty due to homogeneity was derived from the ANOVA. Heterogeneity was not detected under the conditions of the ANOVA.

Analytical method: HPLC Sample size: 12 mg

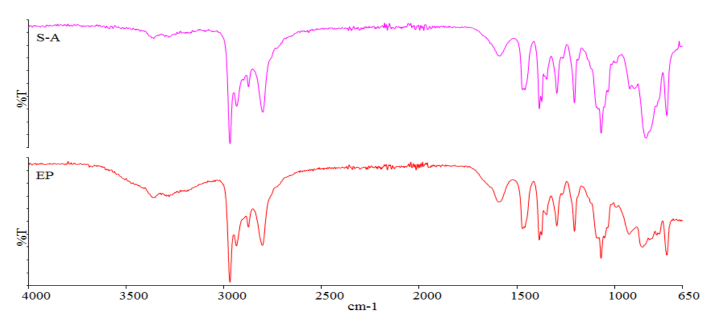
Stability assessment:

Significance of the stability assessment will be demonstrated if the analytical result of the study and the range of values represented by the Expanded Uncertainty do not overlap the result of the original assay and the range of its values represented by the Expanded Uncertainty. The method employed will usually be the same method used to characterize the assay value in the initial evaluation.

Long Term Stability Evaluation - An assessment, or re-test, versus a Compendial Reference Standard may be scheduled, within the 3 year anniversary date of a release of a Secondary Standard. The re-test interval will be determined on a case-by-case basis. Short Term Stability Study - It is useful to assess stability under reasonably anticipated, short term transport conditions by simulating exposure of the product to humidity and temperature stress. This type of study is conducted under controlled conditions of elevated temperature and humidity.

Identification Test:

INFRARED SPECTROPHOTOMETRY (Comparative identification analysis demonstrates direct traceability to Pharmacopeial standards)



S-A Lot LRAC3776 vs. EP Batch 2.0

Certificate of analysis revision history:

Certificate version	Date	Reason for version
LRAC3776.1	08 November 2019	Original Release
LRAC3776.2	05 April 2022	CoA Content Update

Disclaimer:

The purchaser is required to determine the suitability of this product for any particular application. Sigma-Aldrich RTC makes no warranty of any kind, express or implied, other than its products meet all quality control standards set by Sigma-Aldrich RTC. We do not guarantee that the product can be used for any particular application.

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