

Certified Reference Material - Certificate of Analysis

Isotonitazene, Primary Measurement Standard

N,N-Diethyl-2-[[4-(1-methylethoxy)phenyl]methyl]-5-nitro-1H-benzimidazole-1-ethanamine HCl

Product No.:	I-055-1ML	Cerilliant Quality
Lot No.:	FE04142124	ISO 17034
Description of CRM:	Isotonitazene HCI in Methanol (Solution)	130 17034
	Nominal concentration is adjusted for HCI content.	ISO/IEC 17025
Retest Date:	June 2022 See Stability Section	ISO 14001
Storage:	Store unopened in freezer (-10 °C to -25 °C).	ISO 9001
Shipping:	Ambient. See Stability Section	100 7001
Chemical formula:	$C_{23}H_{30}N_4O_3 \cdot HCI$	
CAS No.:	119276-00-5	=⟨CH₃
Regulatory:	USDEA Exempt Canadian TK # 061-1834 H₃C√ ^N ⊂ _{CH}	3 •HCI CH3

Analyte	Certified Concentration \pm associated uncertainty U, $u = k * u$ (k=2)	
Isotonitazene	1.000 ± 0.006 mg/mL	
Metrological traceability:	Traceable to the SI and higher order standards from NIST through an unbroken chain of comparisons. See "Details on metrological traceability" on page 3.	
Measurement method:	The certified value is calculated from high precision weighing of thoroughly characterized starting material. See "Details about certification process" on page 3.	
Intended use:	This Certified Reference Material is suitable for the in vitro identification, calibration, and quantification of the analyte(s) in analytical and R&D applications. Not suitable for human or animal consumption.	
Minimum sample size:	1 μL for quantitative applications	
Instructions for handling and correct	Concentration is corrected for chromatographic purity, residual water, residual solvents, and residual inorganics. No adjustment required before use.	
use:	Users should quantitatively transfer desired volume using established good laboratory practices to spike into matrix or to dilute to the desired concentration. Each ampoule is intended for one-time use.	
	Nominal concentration is adjusted for HCI content. No adjustment required before use.	
Health and safety information:	Danger. Please refer to the Safety Data Sheet for detailed information about the nature of any hazard and appropriate precautions to be taken.	
Accreditation:	Cerilliant Corp. is accredited by the US accreditation authority ANAB as registered reference material producer AR-1353 in accordance with ISO 17034 and registered testing laboratory AT-1352 according to ISO/IEC 17025.	



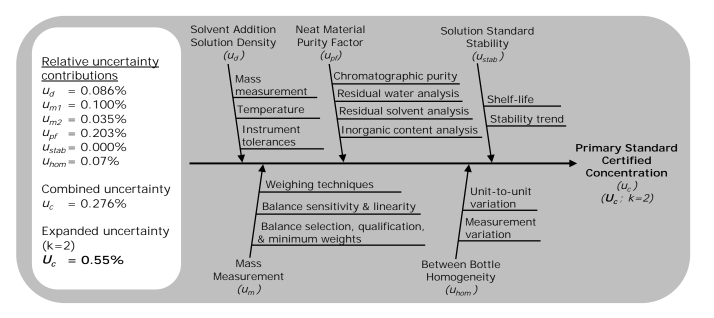
Darron Ellsworth, Quality Assurance Manager

June 16, 2021 Issue Date

Packaging:	2 mL amber USP Type 1 glass ampoule containing not less than 1 mL of certified solution. Ampoules are overfilled to ensure a minimum of 1 mL volume can be transferred when using a 1mL Class A volumetric pipette.
Details on starting materials:	Each raw material utilized has been identified and thoroughly characterized through the use of multiple analytical techniques and is assigned a Mass Balance Purity Factor. Spectral data is provided on subsequent pages of this CoA.
Certificate of Origin:	Cerilliant Corporation certifies no material of animal origin (BSE/TSE) was used in the preparation of this product. This material is a product of USA.

Associated uncertainty:

The uncertainty has been calculated by statistical analysis of all aspects of our production system and incorporated uncertainty of the mass balance purity factor, material density, balance, weighing technique, and homogeneity. Uncertainty components of the gravimetrically prepared Primary Standard concentration are shown in the figure below. Uncertainty is expressed as an expanded uncertainty in accordance with ISO 17034 at the approximate 95% confidence interval using a coverage factor of k=2. Uncertainty contribution from neat material homogeneity was established to be negligible through establishment of process controls and verification of the control process. Stability uncertainty was determined to be negligible by regression analysis.



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.
- The density and material Mass Balance Purity Factor of each raw material is traceable to the SI and higher order reference materials through mass measurement and instrument qualification and calibrations.

Details about certification process:

This standard has been prepared and certified under the ISO 17034, ISO/IEC 17025, and ISO 9001 standards. This standard meets the requirements of a Certified Reference Material and a Primary Standard as defined by ISO and is traceable to the SI and higher order standards through an unbroken chain of comparisons.

- Nominal concentration is calculated based on: the actual mass; Mass balance purity factor of the analyte(s); measured mass of the solution; and the density of the pure diluent at 20°C.
- Fill volume is gravimetrically verified throughout the dispensing process using qualified and calibrated balances.
- Concentration is verified against an independently prepared calibration solution gravimetrically prepared.
- Additional certification information available upon request.

Solution Standard Verification

Concentration accuracy and within- and between-bottle homogeneity are analytically verified against an independently prepared calibration solution.

Standard Solution Assay Parameters		Calibration Curve			
Analysis Method:	HPLC/UV		Calibration Curve: Linear Regres		Linear Regression
Column:	Ascentis Express C	18, 2.7 µm, 3.0 x 100 mm	Number of Points: 4		4
Mobile Phase:	Acetonitrile: 0.1% Phosphoric acid in Water		Linearity (r) : 1.000		1.000
	(35:65)				
Flow Rate:	1.2 mL/min				
Wavelength:	240 nm				
		Verified Concentration	(mg/mL)	%	RSD - Homogeneity
Standard Solution	Lot Number	Actual Result	s		Actual Results
New Lot	FE04142124	1.018			1.8

• Concentration is verified through multiple analyses and is calculated as the average of multiple analyses compared to an independently prepared calibration solution.

 Within-sample and between-sample homogeneity of the New Lot is ensured through rigorous production process controls statistically analyzed to evaluate risk and verified by analysis. Multiple samples pulled from across the lot using a random stratified sampling plan were analyzed to verify homogeneity. % RSD results shown above for the New Lot demonstrate ampoule-to-ampoule homogeneity.

Analyte Certification - Mass Balance Purity Factor

Each analyte is thoroughly identified and characterized using an orthogonal approach. A mass balance purity factor is assigned incorporating chromatographic purity and residual impurities. The mass balance purity factor and salt adjustment are utilized to calculate the weighing adjustment necessary to ensure accuracy of the solution standard concentration.

Material Name: Material Lot: Chemical Formula: CAS Number:	Isotonitazene HCI FC11112007 $C_{23}H_{30}N_4O_3 \bullet$ HCI 119276-00-5	Molecular Weig Molecular Weig Salt Adjustmer	ght (salt):	410.51 446.97 1.089
	Material Charact	erization Summary		
Analytical Test		Method	R	esults
Primary Chromatographic Purity by HPLC/UV Analysis		20384348	99.5%	
Secondary Chromatographic Purity by GC/FID Analysis		20384346	99.1%	
Identity by LC/MS Analysis		20384217	Consistent with Structure	
Identity by ¹ H-NMR Analysis		20384224	Consistent	with Structure
Residual Solvent Analysis by GC/FID Headspace		20397799 ¹	C	0.15%
Residual Water Analysis by Karl Fischer Coulometry		20398075 ¹	1	.11%
Inorganic Content by Microash Analysis		20384350	Below Qua	antitation Limit
Mass Balance Purity Factor			9	8.26%

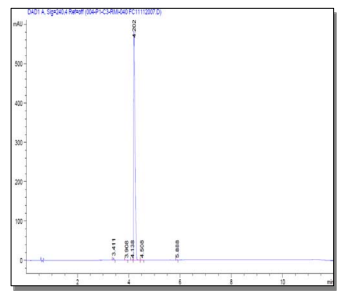
¹ Validated analytical method

• The primary chromatographic purity is calculated as the average of two independently performed analyses utilizing two different methods. Acceptance criteria requires the purity values to be within 0.5% of each other.

- The primary purity method was selected to optimize resolution of impurities while minimizing degradation of the analyte. Secondary purity methods with orthogonal detector capabilities from the primary purity method are used as controls to confirm an accurate purity value.
- The primary chromatographic purity value is used to calculate the Mass Balance Purity Factor.
- A secondary chromatographic purity method is utilized as a control.
- Mass Balance Purity Factor = [(100 wt% residual solvent wt% residual water wt% residual inorganics) x Chromatographic Purity/100].
- Mass Balance Purity Factor does not include adjustment for chiral and/or isotopic purity.

Spectral and Physical Data

HPLC/UV

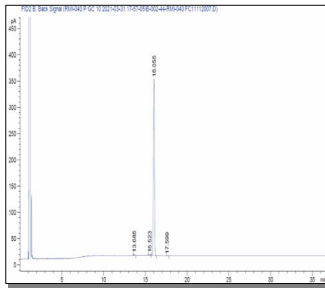


Column:	Ascentis E	xpress C1	8, 2.7 µ	ım,
	3.0 x 100	mm		
Mobile Phase:	: A: Acetoni	trile		
	B: 0.1% P	hosphoric	acid in	Water
Gradient:	Time (min)	% A	% B	
	0.0	15	85	
	8.0	80	20	
	10.0	80	20	
	10.1	15	85	
Flow Rate:	0.7 mL/mi	n		
Wavelength:	240 nm			
Sample Name	FC1111200	07		
Acquired:	March 27,	2021		
Peak # R	Ret Time	Area %		
1	3.41	0.39		
2	3.91	0.02		
3	4.14	0.03		
4	4.20	99.51		
5	4.51	0.04		

0.01

5.89

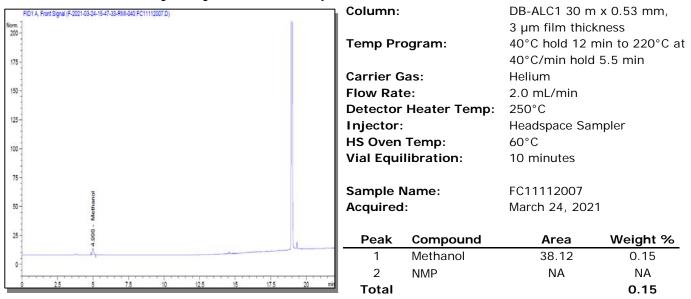
GC/FID



Column:		ms, 30 m x 0.53 mm ID, n film thickness		
Temp Prog	Jram: 40°C t	o 310°C at 40°C/min		
Injector Te		hold 30 min Cool-on-Column		
Detector T	emp: 325°C			
Sample Na				
Acquired:	Warch	31, 2021		
Peak #	Ret Time	Area %		
1	13.69	0.55		
2	15.52	0.34		
3	16.06	99.07		
4	17.60	0.03		

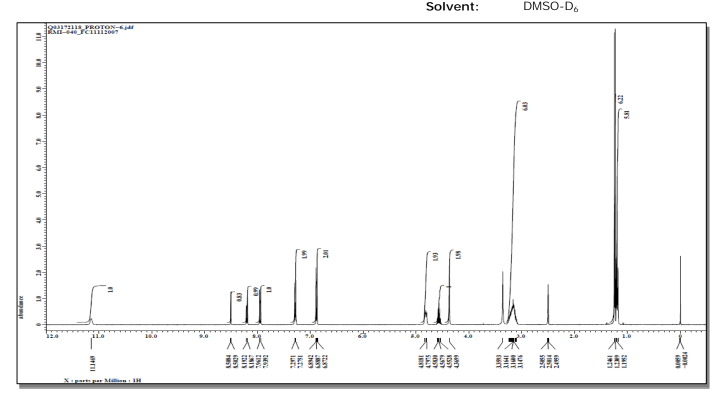
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Residual Solvent Analysis by GC/FID Headspace

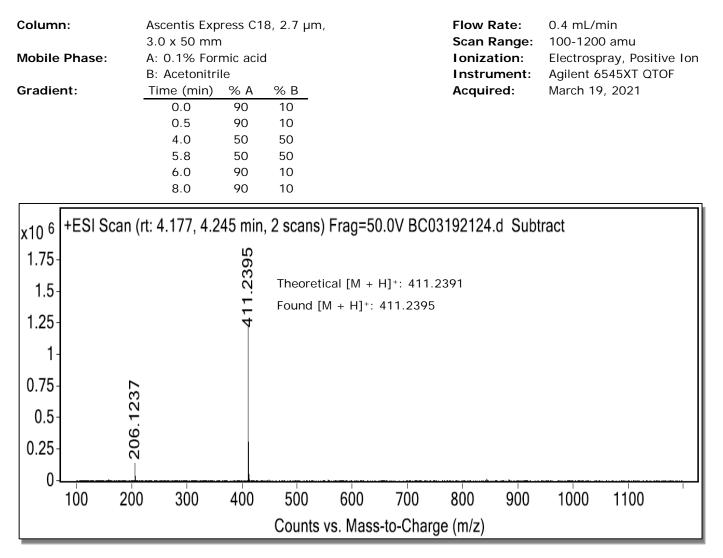


¹H NMR

Instrument:	JEOL ECS 400
Colvert	



LC/MS



Stability

Short term stability studies have been performed in multiple storage conditions for a period of up to four weeks. Short term data is utilized to support transport conditions and normal laboratory use. Real-time stability studies are performed at the recommended storage conditions over the life of the product.

Short Term Stability: A summary of stability findings for this product is listed below.

Storage Condition	Targeted Mean Kinetic Temperature (MKT)	Time Period/Result	
Freezer	-20°C		
Refrigerator	5°C	No decrease in purity was noted after four weeks.	
Room Temperature	20°C		
40°C	40°C		
Transport/Shipping: Stability studies support the transport of this product at ambient conditions.			

Commutability

This standard is a solution of a pure substance in an organic solvent and is a Primary Standard. This Primary Standard is suitable for use in the preparation of calibrators and/or controls in any biological matrix. This standard is not in a biological matrix and therefore commutability to methods or standards in biological matrices does not apply.

COA Revision History

Revision No.	Date	Reason for Revision
00	June 16, 2021	Initial version.



Certified Reference Material - Certificate of Analysis

Isotonitazene-¹³C₆, Primary Measurement Standard

N,N-Diethyl-2-[[4-(1-methylethoxy)phenyl-¹³C₆]methyl]-5-nitro-1H-benzimidazole-1-ethanamine HCl

Product No.:	I-057-1ML	
Lot No.:	FE04142125	Cerilliant Quality
Description of CRM:	Isotonitazene- $^{13}C_{6}$ HCl in Methanol (Solution)	ISO 17034
	Nominal concentration is adjusted for HCl content.	ISO/IEC 17025
Retest Date:	June 2022 See Stability Section	ISO 14001
Storage:	Store unopened in freezer (-10 °C to -25 °C). $O_2 N_{13c} = N_{13c} N_{13c}$	ISO 9001
Shipping:	Ambient. See Stability Section $13\dot{c}_{13}\dot{c}_{N}$	
Chemical formula:	$C_{17}^{13}C_{6}H_{30}N_{4}O_{3} \bullet HCI$	=< сн₃
CAS No.:		O-(CH ₃
Regulatory:	USDEA Exempt Canadian TK # 061-1836	Ŭ
Analyte	Certified Concentration ± associated uncertainty U, u =	
Isotonitazene- ¹³ C ₆	1.000 ± 0.006 mg/mL	
Metrological traceability:	Traceable to the SI and higher order standards from NIST throug chain of comparisons. See "Details on metrological traceability" o	
Measurement method:	The certified value is calculated from high precision weighing of t characterized starting material. See "Details about certification provide the set of t	horoughly
Intended use:	page 3. This Certified Reference Material is suitable for the in vitro identif calibration, and quantification of the analyte(s) in analytical and Not suitable for human or animal consumption.	•
Minimum sample size:	1 μ L for quantitative applications	
Instructions for	Concentration is corrected for chromatographic purity, residual w	ater, residual
handling and correct	solvents, and residual inorganics. No adjustment required before	use.
use:	Users should quantitatively transfer desired volume using establis laboratory practices to spike into matrix or to dilute to the desired Each ampoule is intended for one-time use.	d concentration.
	Nominal concentration is adjusted for HCl content. No adjustmer use.	it required before
	For MS Applications, we advise laboratories not to mix lots during sequence.) a single
Health and safety	Danger. Please refer to the Safety Data Sheet for detailed inform	ation about the
information:	nature of any hazard and appropriate precautions to be taken.	
Accreditation:	Cerilliant Corp. is accredited by the US accreditation authority AN reference material producer AR-1353 in accordance with ISO 170 testing laboratory AT-1352 according to ISO/IEC 17025.	



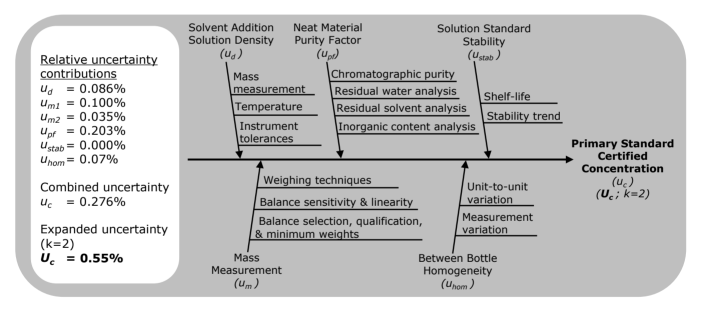
Darron Ellsworth, Quality Assurance Manager

June 16, 2021 Issue Date

Packaging:	2 mL amber USP Type 1 glass ampoule containing not less than 1 mL of certified solution. Ampoules are overfilled to ensure a minimum of 1 mL volume can be transferred when using a 1mL Class A volumetric pipette.
Details on starting materials:	Each raw material utilized has been identified and thoroughly characterized through the use of multiple analytical techniques and is assigned a Mass Balance Purity Factor. Spectral data is provided on subsequent pages of this CoA.
Certificate of Origin:	Cerilliant Corporation certifies no material of animal origin (BSE/TSE) was used in the preparation of this product. This material is a product of the USA.

Associated uncertainty:

The uncertainty has been calculated by statistical analysis of all aspects of our production system and incorporated uncertainty of the mass balance purity factor, material density, balance, weighing technique, and homogeneity. Uncertainty components of the gravimetrically prepared Primary Standard concentration are shown in the figure below. Uncertainty is expressed as an expanded uncertainty in accordance with ISO 17034 at the approximate 95% confidence interval using a coverage factor of k=2. Uncertainty contribution from neat material homogeneity was established to be negligible through establishment of process controls and verification of the control process. Stability uncertainty was determined to be negligible by regression analysis.



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.
- The density and material Mass Balance Purity Factor of each raw material is traceable to the SI and higher order reference materials through mass measurement and instrument qualification and calibrations.

Details about certification process:

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- Nominal concentration is calculated based on: the actual mass; Mass balance purity factor of the analyte(s); measured mass of the solution; and the density of the pure diluent at 20°C.
- Fill volume is gravimetrically verified throughout the dispensing process using qualified and calibrated balances.
- Concentration is verified against an independently prepared calibration solution gravimetrically prepared.
- Additional certification information available upon request.

Solution Standard Verification

Concentration accuracy and within- and between-bottle homogeneity are analytically verified against an independently prepared calibration solution.

Standard Solution	Calibration Curve				
Analysis Method:	HPLC/UV	Calibration Curve: Linear Regre		Linear Regression	
Column:	Ascentis Express C	18, 2.7 µm, 3.0 x 100 mm	Number of Points:		4
Mobile Phase:	Acetonitrile:0.1% F (35:65)	Phosphoric acid in Water	Linearity (r) : 1.000		1.000
Flow Rate:	1.2 mL/min				
Wavelength:	240 nm				
		Verified Concentration	(mg/mL)	%	RSD - Homogeneity
Standard Solution	Lot Number	Actual Result	s		Actual Results
New Lot	FE04142125	1.016			1.9

• Concentration is verified through multiple analyses and is calculated as the average of multiple analyses compared to an independently prepared calibration solution.

 Within-sample and between-sample homogeneity of the New Lot is ensured through rigorous production process controls statistically analyzed to evaluate risk and verified by analysis. Multiple samples pulled from across the lot using a random stratified sampling plan were analyzed to verify homogeneity. % RSD results shown above for the New Lot demonstrate ampoule-to-ampoule homogeneity.

Analyte Certification - Mass Balance Purity Factor

Each analyte is thoroughly identified and characterized using an orthogonal approach. A mass balance purity factor is assigned incorporating chromatographic purity and residual impurities. The mass balance purity factor and salt adjustment are utilized to calculate the weighing adjustment necessary to ensure accuracy of the solution standard concentration.

Material Name: Material Lot:	Isotonitazene- ¹³ C ₆ HCl FC10062003		[·] Weight (base): [·] Weight (salt):	416.47 452.93		
Chemical Formula: CAS Number:	$C_{17}^{13}C_6H_{30}N_4O_3 \bullet HCI$ NA	Salt Adjustment: 1.088				
	Material Characte	rization Summar	Y			
Analytical Test		Method	Res	ults		
Primary Chromatographic	Purity by HPLC/UV Analysis	20384348	99.8	3%		
Secondary Chromatograp	hic Purity by GC/FID Analysis	20384346	99.3	3%		
Identity by LC/MS Analys	is	20384217	Consistent w	Consistent with Structure		
			0.00% 130	C ₀ vs ¹³ C ₆		
Isotopic Purity and Distrib	oution by LC/MS SIM Analysis	20384217	0.00% ¹³ C ₀ vs ¹³ C	4.50% ¹³ C ₅		
			0.10% ¹³ C ₄	95.40% ¹³ C ₆		
Identity by ¹ H-NMR Analy	vsis	20384224	Consistent w	th Structure		
Residual Solvent Analysis	by GC/FID Headspace	20397799 ¹	0.49%			
Residual Water Analysis b	by Karl Fischer Coulometry	20398075 ¹	0.82	0.82%		
Inorganic Content by Mic	roash Analysis	20384350	Below Quant	itation Limit		
Mass Balance Purity Facto	or		98.4	6%		

¹ Validated analytical method

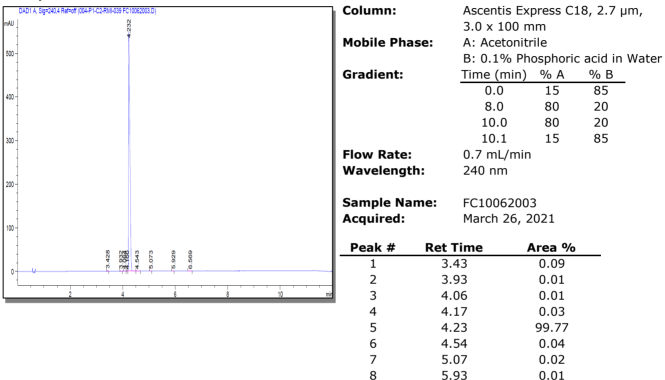
• The primary chromatographic purity is calculated as the average of two independently performed analyses utilizing two different methods. Acceptance criteria requires the purity values to be within 0.5% of each other.

• The primary purity method was selected to optimize resolution of impurities while minimizing degradation of the analyte. Secondary purity methods with orthogonal detector capabilities from the primary purity method are used as controls to confirm an accurate purity value.

- The primary chromatographic purity value is used to calculate the Mass Balance Purity Factor.
- A secondary chromatographic purity method is utilized as a control.
- Mass Balance Purity Factor = [(100 wt% residual solvent wt% residual water wt% residual inorganics) x Chromatographic Purity/100].
- Mass Balance Purity Factor does not include adjustment for chiral and/or isotopic purity.

Spectral and Physical Data

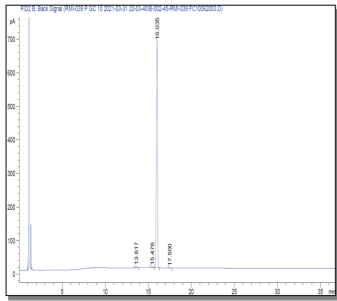
HPLC/UV



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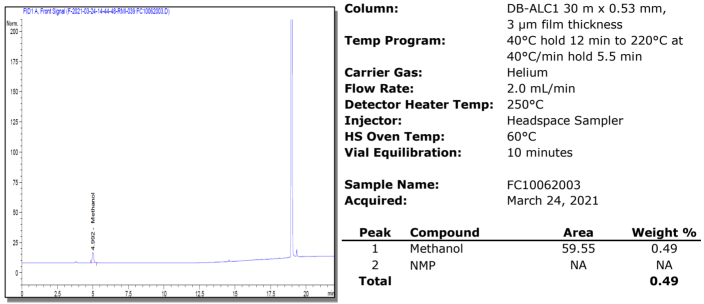
GC/FID



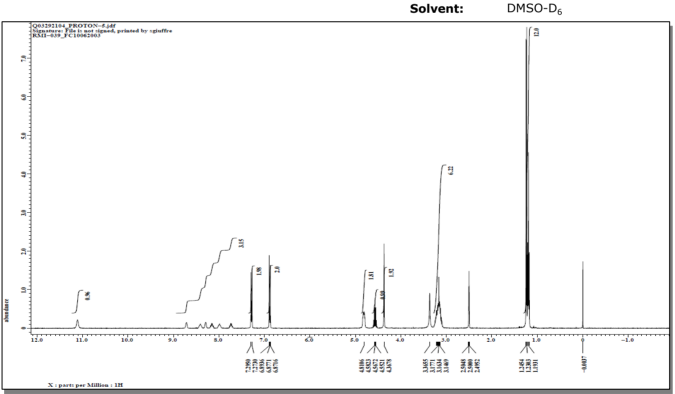
Column:		DB-35ms, 30 m x 0.53 mm ID,		
Temp Program:		1.0 μm film thickness 40°C to 310°C at 40°C/min hold 30 min		
Injector Te	emp: C	ool-on-Column		
Detector T	emp: 32	25°C		
Commite Name		210062003		
Sample Na Acquired:		March 31, 2021		
Acquireu:	1*1			
Peak #	Ret Tim	e Area %		
1	13.62	0.35		
2	15.48	0.26		
3	16.04	99.34		
4	17.50	0.05		

0.01

Residual Solvent Analysis by GC/FID Headspace



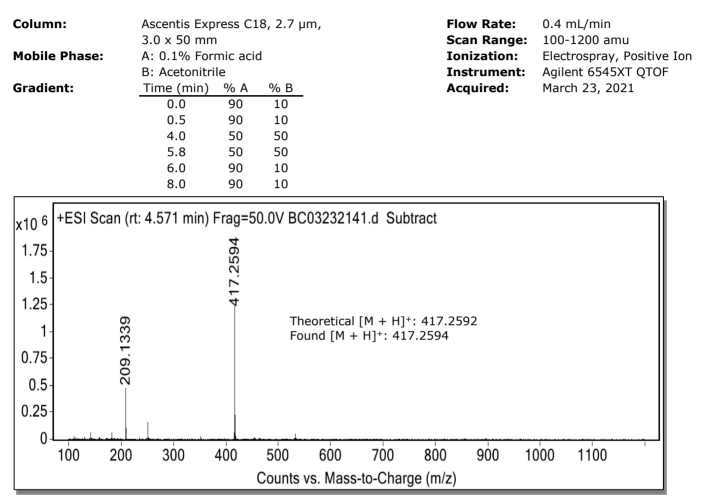
¹H NMR



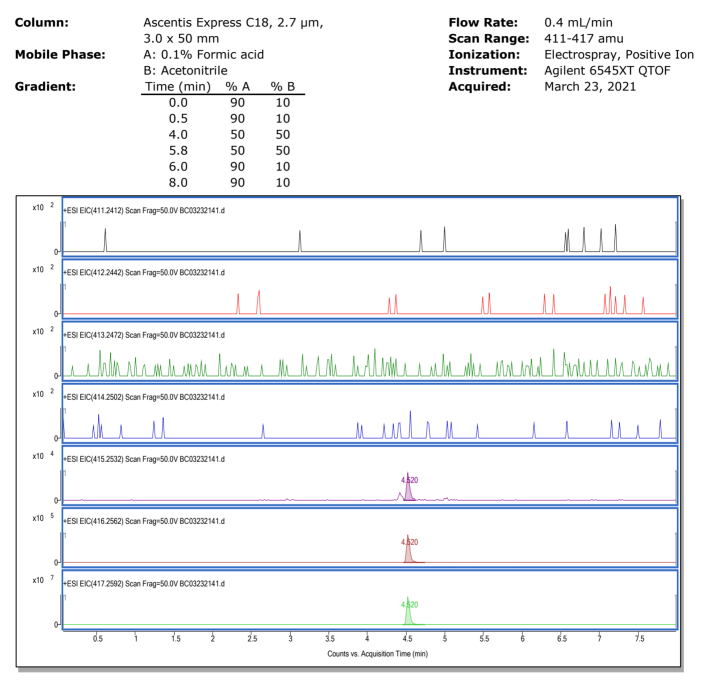
JEOL ECS 400

Instrument:

LC/MS



Isotopic Purity by LC/MS SIM



Stability

Short term stability studies have been performed in multiple storage conditions for a period of up to four weeks. Short term data is utilized to support transport conditions and normal laboratory use. Real-time stability studies are performed at the recommended storage conditions over the life of the product.

Short Term Stability: A summary of stability findings for a related product (I-055-1ML, Isotonitazene HCl) is listed below.

Storage Condition	Targeted Mean Kinetic Temperature (MKT)	Time Period/Result		
Freezer	-20°C			
Refrigerator	5°C	No decrease in purity was noted after		
Room Temperature	20°C	four weeks.		
40°C	40°C			
Transport/Shipping: Stability studies support the transport of this product at ambient conditions.				

Commutability

This standard is a solution of a pure substance in an organic solvent and is a Primary Standard. This Primary Standard is suitable for use in the preparation of calibrators and/or controls in any biological matrix. This standard is not in a biological matrix and therefore commutability to methods or standards in biological matrices does not apply.

COA Revision History

Revision No.	Date	Reason for Revision
00	June 16, 2021	Initial version.



Certified Reference Material - Certificate of Analysis

N-Phenethyl-4-piperidone (NPP), Primary Measurement Standard

Product No.: Lot No.: Description of CRM: Retest Date: Storage: Shipping: Chemical formula:	P-165-1ML FN06302103 N-Phenethyl-4-piperidone (NPP) in Acetonitrile (Solution) September 2022 See Stability Section Store unopened in freezer (-10 °C to -25 °C). Ambient. See Stability Section C ₁₃ H ₁₇ NO	Cerilliant Quality ISO 17034 ISO/IEC 17025 ISO 14001 ISO 9001	
CAS No.: Analyte	39742-60-4 Certified Concentration ± associated uncertainty <i>U</i> , <i>u</i> =	=k*u (k=2)	
N-Phenethyl-4-piperi	done (NPP) 1.000 ± 0.006 mg/mL		
Metrological traceability	: Traceable to the SI and higher order standards from NIST throug		
Measurement method:	chain of comparisons. See "Details on metrological traceability" on page 3. The certified value is calculated from high precision weighing of thoroughly characterized starting material. See "Details about certification process" on		
Intended use:	page 3. This Certified Reference Material is suitable for the in vitro identification, calibration, and quantification of the analyte(s) in analytical and R&D applications. Not suitable for human or animal consumption.		
Minimum sample size:	1 μ L for quantitative applications		
Instructions for handling and correct use:	Concentration is corrected for chromatographic purity, residual water, residual solvents, and residual inorganics. No adjustment required before use. Users should quantitatively transfer desired volume using established good laboratory practices to spike into matrix or to dilute to the desired concentration.		
Health and safety information: Accreditation:	Each ampoule is intended for one-time use. Danger. Please refer to the Safety Data Sheet for detailed informanature of any hazard and appropriate precautions to be taken. Cerilliant Corp. is accredited by the US accreditation authority AN reference material producer AR-1353 in accordance with ISO 170 testing laboratory AT-1352 according to ISO/IEC 17025.	IAB as registered	



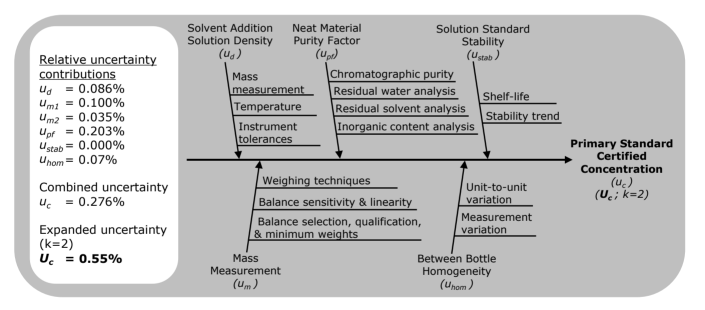
Darron Ellsworth, Quality Assurance Manager

August 06, 2021 Issue Date

Packaging:	2 mL amber USP Type 1 glass ampoule containing not less than 1 mL of certified solution. Ampoules are overfilled to ensure a minimum of 1 mL volume can be transferred when using a 1mL Class A volumetric pipette.
Details on starting materials:	Each raw material utilized has been identified and thoroughly characterized through the use of multiple analytical techniques and is assigned a Mass Balance Purity Factor. Spectral data is provided on subsequent pages of this CoA.
Certificate of Origin:	Cerilliant Corporation certifies no material of animal origin (BSE/TSE) was used in the preparation of this product. This material is a product of the USA.

Associated uncertainty:

The uncertainty has been calculated by statistical analysis of all aspects of our production system and incorporated uncertainty of the mass balance purity factor, material density, balance, weighing technique, and homogeneity. Uncertainty components of the gravimetrically prepared Primary Standard concentration are shown in the figure below. Uncertainty is expressed as an expanded uncertainty in accordance with ISO 17034 at the approximate 95% confidence interval using a coverage factor of k=2. Uncertainty contribution from neat material homogeneity was established to be negligible through establishment of process controls and verification of the control process. Stability uncertainty was determined to be negligible by regression analysis.



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.
- The density and material Mass Balance Purity Factor of each raw material is traceable to the SI and higher order reference materials through mass measurement and instrument qualification and calibrations.

Details about certification process:

This standard has been prepared and certified under the ISO 17034, ISO/IEC 17025, and ISO 9001 standards. This standard meets the requirements of a Certified Reference Material and a Primary Standard as defined by ISO and is traceable to the SI and higher order standards through an unbroken chain of comparisons.

- Nominal concentration is calculated based on: the actual mass; Mass balance purity factor of the analyte(s); measured mass of the solution; and the density of the pure diluent at 20°C.
- Fill volume is gravimetrically verified throughout the dispensing process using qualified and calibrated balances.
- Concentration is verified against an independently prepared calibration solution gravimetrically prepared.
- Additional certification information available upon request.

Solution Standard Verification

Concentration accuracy and within- and between-bottle homogeneity are analytically verified against an independently prepared calibration solution.

Standard Solution	Calibration Curve				
Analysis Method:	HPLC/UV	Calibration Curve: Linear Regress		Linear Regression	
Column:	Ascentis Express C18, 2.7 µm, 3.0 x 100 mm		Number of Points: 4		4
Mobile Phase:	Acetonitrile:0.1% P (12:88)	-		Linearity (r) : 1.000	
Flow Rate:	1.25 mL/min				
Wavelength:	210 nm				
		Verified Concentration	n (mg/mL)	%	RSD - Homogeneity
Standard Solution	Lot Number	Actual Result	S		Actual Results
New Lot	FN06302103	1.002			0.2

• Concentration is verified through multiple analyses and is calculated as the average of multiple analyses compared to an independently prepared calibration solution.

 Within-sample and between-sample homogeneity of the New Lot is ensured through rigorous production process controls statistically analyzed to evaluate risk and verified by analysis. Multiple samples pulled from across the lot using a random stratified sampling plan were analyzed to verify homogeneity. % RSD results shown above for the New Lot demonstrate ampoule-to-ampoule homogeneity.

Analyte Certification - Mass Balance Purity Factor

Each analyte is thoroughly identified and characterized using an orthogonal approach. A mass balance purity factor is assigned incorporating chromatographic purity and residual impurities. The mass balance purity factor is utilized to calculate the weighing adjustment necessary to ensure accuracy of the solution standard concentration.

Material Name: Material Lot:	N-Phenethyl-4-piperidone (N FN11112006	CAS Number: Molecular Weig	39742-60-4
Analytical Test	Material Characte		Deculto
Analytical Test		Method	Results
Primary Chromatograph	ic Purity by HPLC/UV Analysis	20384348	99.9%
Secondary Chromatogra	aphic Purity by GC/FID Analysis	20384346	> 99.9%
Identity by LC/MS Analy	/sis	20384217	Consistent with Structure
Identity by ¹ H-NMR Ana	lysis	20384224	Consistent with Structure
Residual Solvent Analysis by GC/FID Headspace		20397799 ¹	0.17%
Residual Water Analysis by Karl Fischer Coulometry		20398075 ¹	Below Quantitation Limit
Inorganic Content by Microash Analysis		20384350	Below Quantitation Limit
Mass Balance Purity Fac	tor		99.73%

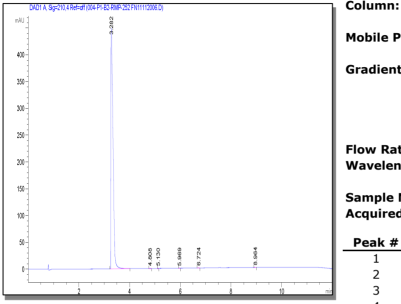
¹ Validated analytical method

• The primary chromatographic purity is calculated as the average of two independently performed analyses utilizing two different methods. Acceptance criteria requires the purity values to be within 0.5% of each other.

- The primary purity method was selected to optimize resolution of impurities while minimizing degradation of the analyte. Secondary purity methods with orthogonal detector capabilities from the primary purity method are used as controls to confirm an accurate purity value.
- The primary chromatographic purity value is used to calculate the Mass Balance Purity Factor.
- A secondary chromatographic purity method is utilized as a control.
- Mass Balance Purity Factor = [(100 wt% residual solvent wt% residual water wt% residual inorganics) x Chromatographic Purity/100].
- Mass Balance Purity Factor does not include adjustment for chiral and/or isotopic purity.

Spectral and Physical Data

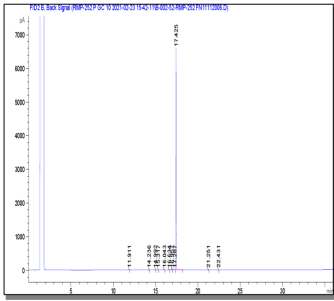
HPLC/UV



Column:	Ascentis Express C18, 2.7 µm,		
	3.0 x 100 m		
Mobile Phase:	A: Acetonitr	ile	
	B: 0.1% Ph	osphoric	c acid in Water
Gradient:	Time (min)	% A	% B
	0.0	2	98
	8.0	50	50
	10.0	50	50
	10.1	2	98
Flow Rate:	0.6 mL/min		
Wavelength:	210 nm		
Sample Name:	FN1111200	6	
Acquired:	February 23		
	, .	, -	
Peak # Ret	Time A	Area %	
1 3.	28	99.91	
2 4.	81	0.03	
3 5.	13	0.01	
4 5.	97	0.01	
5 6.	72	0.03	

0.01

GC/FID

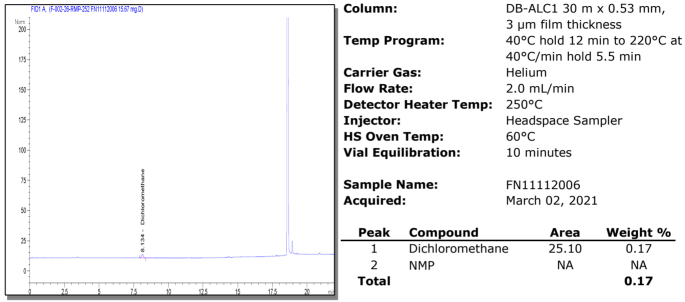


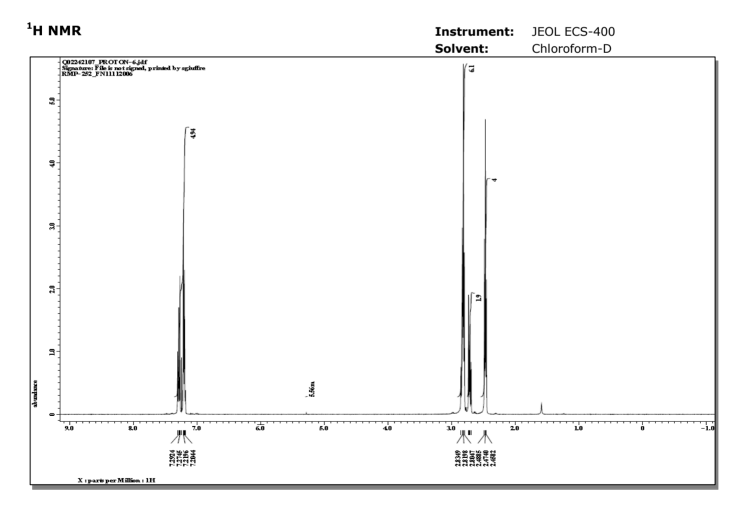
	Injector Temp:		DB-35ms, 30 m x 0.53 mm ID, 1.0 µm film thickness 40°C to 280°C at 10°C/min	
	Sample Na Acquired:	me:		112006 Jary 23, 2021
	Peak #	Ret T	ime	Area %
	1	11.	91	0.00
	2	14.	24	0.00
	3	14.	99	0.00
	4	15.	32	0.00
	5	16.	04	0.01
_	6	16.	63	0.00
min	7	16.	96	0.00
100	8	17.	29	0.01
	9	17.	43	99.96
	10	21.	25	0.01
	11	22.	43	0.00

8.96

6

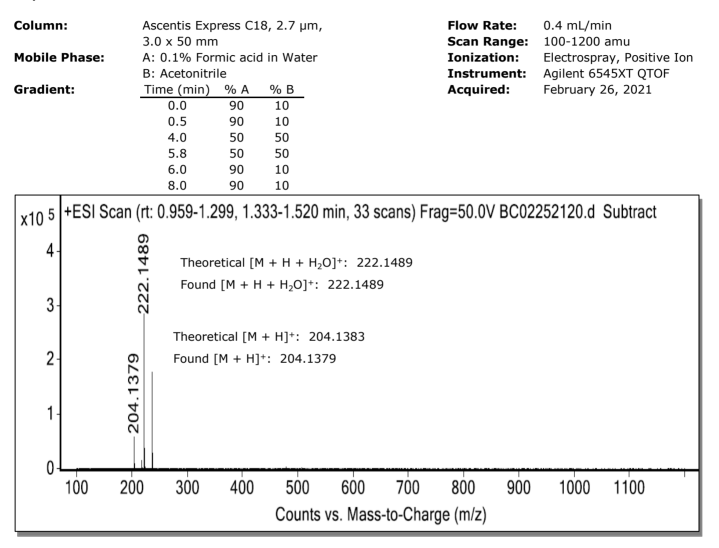
Residual Solvent Analysis by GC/FID Headspace





Cerilliant Corporation, 811 Paloma Drive, Suite A Round Rock, TX 78665, USA, Tel: 800-848-7837 / 512-238-9974

LC/MS



Stability

Short term stability studies have been performed in multiple storage conditions for a period of up to one week. Short term data is utilized to support transport conditions and normal laboratory use. Real-time stability studies are performed at the recommended storage conditions over the life of the product.

Short Term Stability: A summary of stability findings for this product is listed below.

Storage Condition	Targeted Mean Kinetic Temperature (MKT)	Time Period/Result		
Freezer	-20°C			
Refrigerator	5°C	No decrease in purity was noted after		
Room Temperature 20°C		one week.		
40°C	40°C			
Transport/Shipping: Stability studies support the transport of this product at ambient conditions.				

Commutability

This standard is a solution of a pure substance in an organic solvent and is a Primary Standard. This Primary Standard is suitable for use in the preparation of calibrators and/or controls in any biological matrix. This standard is not in a biological matrix and therefore commutability to methods or standards in biological matrices does not apply.

COA Revision History

Revision No.	Date	Reason for Revision
00	August 06, 2021	Initial version.



Certified Reference Material - Certificate of Analysis

N-Phenethyl-4-piperidone- ${}^{13}C_6$ (NPP- ${}^{13}C_6$), Primary Measurement Standard

	1-Phenethyl-4-piperidone- ¹³ C ₆				
Product No.:	P-167-1ML	Cerilliant Quality			
Lot No.:	FN06142104	ISO 17034			
Description of CRM:	N-Phenethyl-4-piperidone ⁻¹³ C ₆ (NPP ⁻¹³ C ₆) ISO/IEC 13				
	in Acetonitrile (Solution)	ISO 14001			
Retest Date:	September 2022 See Stability Section	10.0 0001			
Storage:	Store unopened in freezer (-10 °C to -25 °C). Ambient. See Stability Section $C_7^{13}C_{eH_{17}NO}$	ISO 9001			
Shipping:	Ambient. See Stability Section $N^{13}C_{13}C^{$				
Chemical formula:	$C_7^{13}C_6H_{17}NO$				
CAS No.:	NA				
AnalyteCertified Concentration ± associated uncertainty U, u = k * u (k =					
N-Phenethyl-4-piperio	done- $^{13}C_6$ (NPP- $^{13}C_6$) 1.000 ± 0.006 mg/mL				
Metrological traceability:	Metrological traceability: Traceable to the SI and higher order standards from NIST through an unbroken chain of comparisons. See "Details on metrological traceability" on page 3.				
Measurement method:	The certified value is calculated from high precision weighing of thoroughly characterized starting material. See "Details about certification process" on				
Intended use:	page 3. This Certified Reference Material is suitable for the in vitro identification, calibration, and quantification of the analyte(s) in analytical and R&D applications. Not suitable for human or animal consumption.				
Minimum sample size:	1 μL for quantitative applications				
Instructions for	Concentration is corrected for chromatographic purity, residual water, residual				
handling and correct	solvents, and residual inorganics. No adjustment required before use. Users should quantitatively transfer desired volume using established good laboratory practices to spike into matrix or to dilute to the desired concentration.				
use:					
	Each ampoule is intended for one-time use.				
	For MS Applications, we advise laboratories not to mix lots during a single				
	sequence.				
Health and safety	Danger. Please refer to the Safety Data Sheet for detailed inform	nation about the			
information:	nature of any hazard and appropriate precautions to be taken.				
Accreditation: Cerilliant Corp. is accredited by the US accreditation authority ANAB as reference material producer AR-1353 in accordance with ISO 17034 and testing laboratory AT-1352 according to ISO/IEC 17025.					



Darron Ellsworth, Quality Assurance Manager

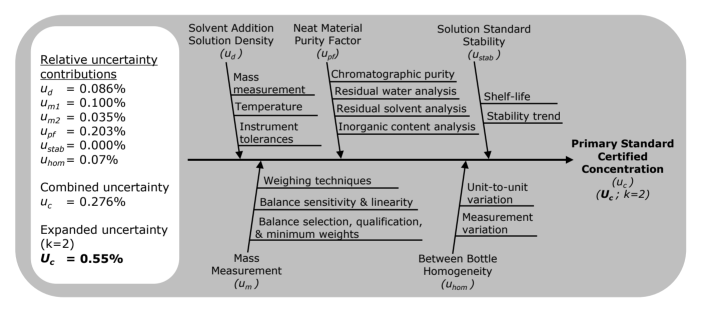
August 09, 2021

Issue Date

Packaging:	2 mL amber USP Type 1 glass ampoule containing not less than 1 mL of certified solution. Ampoules are overfilled to ensure a minimum of 1 mL volume can be transferred when using a 1mL Class A volumetric pipette.
Details on starting materials:	Each raw material utilized has been identified and thoroughly characterized through the use of multiple analytical techniques and is assigned a Mass Balance Purity Factor. Spectral data is provided on subsequent pages of this CoA.
Certificate of Origin:	Cerilliant Corporation certifies no material of animal origin (BSE/TSE) was used in the preparation of this product. This material is a product of the USA.

Associated uncertainty:

The uncertainty has been calculated by statistical analysis of all aspects of our production system and incorporated uncertainty of the mass balance purity factor, material density, balance, weighing technique, and homogeneity. Uncertainty components of the gravimetrically prepared Primary Standard concentration are shown in the figure below. Uncertainty is expressed as an expanded uncertainty in accordance with ISO 17034 at the approximate 95% confidence interval using a coverage factor of k=2. Uncertainty contribution from neat material homogeneity was established to be negligible through establishment of process controls and verification of the control process. Stability uncertainty was determined to be negligible by regression analysis.



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.
- The density and material Mass Balance Purity Factor of each raw material is traceable to the SI and higher order reference materials through mass measurement and instrument qualification and calibrations.

Details about certification process:

This standard has been prepared and certified under the ISO 17034, ISO/IEC 17025, and ISO 9001 standards. This standard meets the requirements of a Certified Reference Material and a Primary Standard as defined by ISO and is traceable to the SI and higher order standards through an unbroken chain of comparisons.

- Nominal concentration is calculated based on: the actual mass; Mass balance purity factor of the analyte(s); measured mass of the solution; and the density of the pure diluent at 20°C.
- Fill volume is gravimetrically verified throughout the dispensing process using qualified and calibrated balances.
- Concentration is verified against an independently prepared calibration solution gravimetrically prepared.
- Additional certification information available upon request.

Solution Standard Verification

Concentration accuracy and within- and between-bottle homogeneity are analytically verified against an independently prepared calibration solution.

Standard Solution Assay Parameters			Calibration Curve	
Analysis Method:	HPLC/UV		Calibration Curve	: Linear Regression
Column:	Ascentis Express C1	L8, 2.7 μm, 3.0 x 100 mm	Number of Points	: 4
Mobile Phase:	Acetonitrile:0.1% P (12:88)	hosphoric acid in Water	Linearity (r) :	1.000
Flow Rate:	1.25 mL/min			
Wavelength:	210 nm			
		Verified Concentration	n (mg/mL)	%RSD - Homogeneity
Standard Solution	Lot Number	Actual Result	s	Actual Results
Solution		0.992		0.6

 Concentration is verified through multiple analyses and is calculated as the average of multiple analyses compared to an independently prepared calibration solution.

 Within-sample and between-sample homogeneity of the New Lot is ensured through rigorous production process controls statistically analyzed to evaluate risk and verified by analysis. Multiple samples pulled from across the lot using a random stratified sampling plan were analyzed to verify homogeneity. % RSD results shown above for the New Lot demonstrate ampoule-to-ampoule homogeneity.

Analyte Certification - Mass Balance Purity Factor

Each analyte is thoroughly identified and characterized using an orthogonal approach. A mass balance purity factor is assigned incorporating chromatographic purity and residual impurities. The mass balance purity factor is utilized to calculate the weighing adjustment necessary to ensure accuracy of the solution standard concentration.

Material Name: Material Lot:	N-Phenethyl-4-piperidone- ^{1:} (NPP- ¹³ C ₆) FN06032001	³ C ₆ Chemical Forn CAS Number: Molecular Wei	N	^{,13} C ₆ H ₁₇ NO A)9.24
	Material Characte	erization Summary		
Analytical Test		Method	Re	esults
Primary Chromatograph	ic Purity by HPLC/UV Analysis	20384348	9	9.7%
Secondary Chromatogra	phic Purity by GC/FID Analysis	20384346	9	9.5%
Identity by GC/MS Analy	ysis	20384214	Consistent with Structure	
			0.48%	¹³ C ₀ vs ¹³ C ₆
			0.35% ¹³ C ₀	11.97% ¹³ C ₄
Isotopic Purity and Distr	ibution by GC/MS SIM Analysis	20384214	0.80% ¹³ C ₁	9.46% ¹³ C ₅
			0.90% ¹³ C ₂	74.34% ¹³ C ₆
			2.17% ¹³ C ₃	
Identity by ¹ H-NMR Ana	lysis	20384224	Consistent	with Structure
Residual Solvent Analys	is by GC/FID Headspace	20397799 ¹	0.10%	
Residual Water Analysis	by Karl Fischer Coulometry	20398075 ¹	1.25%	
Inorganic Content by Mi	croash Analysis	20384350	<	0.2%
Mass Balance Purity Fac	tor		98	8.39%

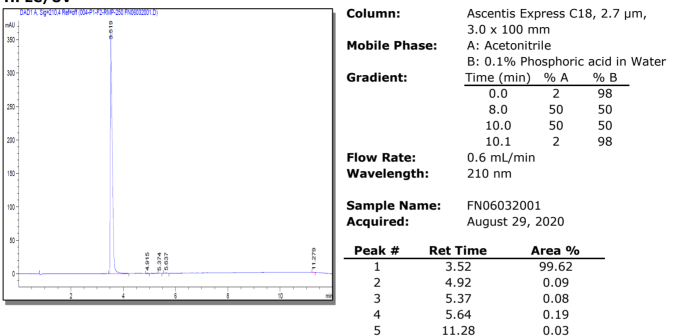
¹ Validated analytical method

• The primary chromatographic purity is calculated as the average of two independently performed analyses utilizing two different methods. Acceptance criteria requires the purity values to be within 0.5% of each other.

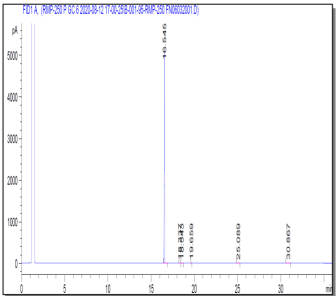
- The primary purity method was selected to optimize resolution of impurities while minimizing degradation of the analyte. Secondary purity methods with orthogonal detector capabilities from the primary purity method are used as controls to confirm an accurate purity value.
- The primary chromatographic purity value is used to calculate the Mass Balance Purity Factor.
- A secondary chromatographic purity method is utilized as a control.
- Mass Balance Purity Factor = [(100 wt% residual solvent wt% residual water wt% residual inorganics) x Chromatographic Purity/100].
- Mass Balance Purity Factor does not include adjustment for chiral and/or isotopic purity.

Spectral and Physical Data

HPLC/UV

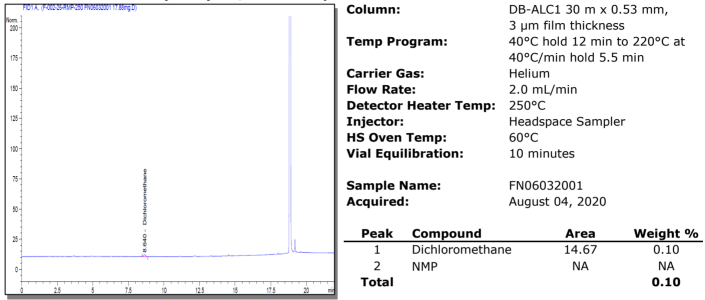


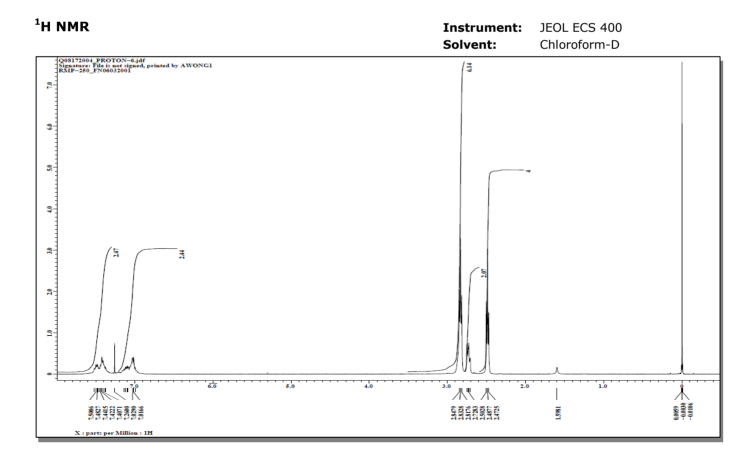
GC/FID



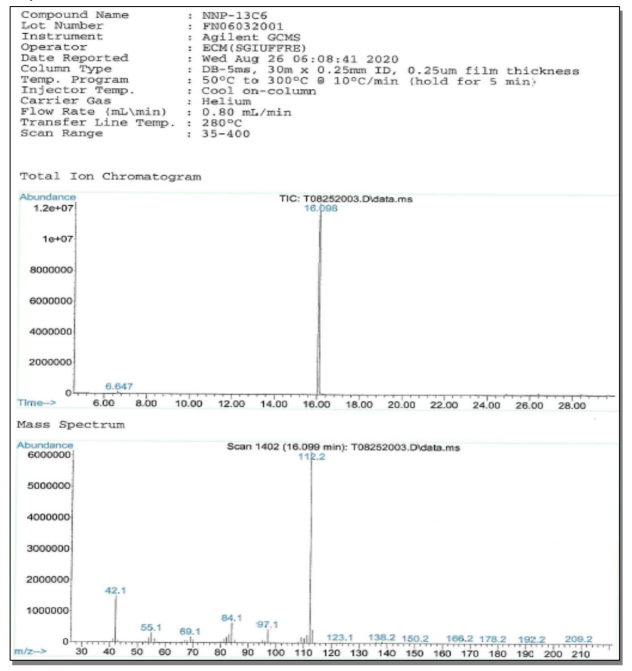
	Column: Temp Program: Injector Temp: Detector Temp:		DB-5ms, 30 m x 0.53 mm ID, 1.5 µm film thickness 40°C to 280°C at 10°C/min hold 12 min Cool-on-Column 325°C		
	Sample Na Acquired:	ime:		032001 st 12, 2020	
	Peak #	Ret T	ime	Area %	
	1	16.	55	99.53	
	2	18.	34	0.10	
	3	18.	54	0.11	
	4	19.	66	0.00	
	5	25.	09	0.14	
	6	30.	87	0.11	
mir					

Residual Solvent Analysis by GC/FID Headspace





GC/MS



Stability

Short term stability studies have been performed in multiple storage conditions for a period of up to one week. Short term data is utilized to support transport conditions and normal laboratory use. Real-time stability studies are performed at the recommended storage conditions over the life of the product.

Short Term Stability: A summary of stability findings for a related product (P-165-1ML, N-Phenethyl-4piperidone (NPP)) is listed below.

Storage Condition	Targeted Mean Kinetic Temperature (MKT)	Time Period/Result		
Freezer	-20°C			
Refrigerator	5°C	No decrease in purity was noted after		
Room Temperature	20°C	one week.		
40°C	40°C			
<i>Transport/Shipping:</i> Stability studies support the transport of this product at ambient conditions.				

Commutability

This standard is a solution of a pure substance in an organic solvent and is a Primary Standard. This Primary Standard is suitable for use in the preparation of calibrators and/or controls in any biological matrix. This standard is not in a biological matrix and therefore commutability to methods or standards in biological matrices does not apply.

COA Revision History

Revision No.	Date	Reason for Revision
00	August 09, 2021	Initial version.