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> ISO GUIDE 34 ISO/IEC 17025 ISO 13485

Certified Reference Material - Certificate of Analysis

Salicylic Acid-D₄, Primary Standard

Catalog Number:	S-042	он о	ISO 9001
Lot:	FN03211402		GMP/GLP
Retest:	May 2018	- Y Y YOH	
Description:	Salicylic Acid-D ₄ in Acetonitrile		
Packaging:	Solution in 2 mL amber USP Type I glass ampoule		
	containing not less than 1 mL of certified solution.	-	
Storage:	Store unopened in freezer (-10 °C to -25 °C).		
Shipping:	Ambient. See Stability Section.		
Intended Use:	This Certified Reference Material is suitable for thein vitro ide	entification, calibration, and quantificat	tion of the
	analyte(s) in analytical and R&D applications. Not suitable for	r human or animal consumption.	
Instructions for Use:	Users should quantitatively transfer desired volume using esta	blished good laboratory practices to	
	spike into matrix or to dilute to the desired concentration. Eac	h ampoule is intended for one-time use	2.
Safety:	Danger. See Safety Data Sheet		

· Retest Date - stability studies ongoing. Certificate of Analysis will be updated upon completion of retest.

- Ampoules are overfilled to ensure a minimum 1 mL volume can be transferred when using a 1 mL Class A volumetric pipette.
- For MS Applications, we advise laboratories not to mix lots during a single sequence.

Analyte	Certified Concentration Value
Salicylic Acid-D ₄	$100.0 \pm 0.5 \ \mu g/mL$
 coverage factor of k = 2 and has been density, balance, and weighing techni This standard is prepared gravimetric 	pressed as an expanded uncertainty in accordance with ISO 17025 and Guide 34 at the approximate 95% confidence interval using calculated by statistical analysis of our production system and incorporates uncertainty of the mass balance purity factor, material ique. ally and mass results are reported on the conventional basis for weighing in air. Nominal concentration is calculated based on: the Purity Factor of the analyte(s); measured mass of the solution; and the density of the pure diluent at 20 °C.
	atographic purity, residual water, residual solvents and residual inorganics. No adjustment required before use.

Metrological Traceability

This standard has been prepared and certified under the ISO Guide 34, ISO/IEC 17025, ISO 9001 and ISO 13485 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.

- 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 0.1% relative error. Balance calibration adjustments are performed weekly utilizing the balance's internal adjustment mechanism. Calibration verifications are performed pre-use. Weigh tapes from the calibration verification are included in the production batch record for this standard. Production data package available upon request.
- 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.
- Each raw material utilized has been identified and thoroughly characterized through the use of multiple analytical techniques. Spectral data is provided on subsequent
 pages of this COA. The density and material Mass Balance Purity Factor is traceable to the SI and higher order reference standards through mass measurement and
 instrument qualification and calibrations.

Cerilliant certifies that this standard meets the specifications stated in this certificate and warrants this product to meet the stated acceptance criteria through the expiration/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.



Darron Ellsworth, Quality Assurance Manager

May 7, 2014

Date

Solution Standard Verification

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

Solution standard verification demonstrates confirmation that the specified requirements for the Primary Standard have been fulfilled and validated under ISO 13485.

Standard Solution	n Assay Parameters		C	alibration Curve			
Analysis Method: UV/Vis			Ca	alibration Curve:	Linear Regression		
Wavelength:	230 nm		Nu	umber of Points:	4 1.000		
Slit Width:	1.0 nm		Li	nearity (r):			
Response:	0.5 s			• • • •			
Standard Solution	Lot Number	Actual Results	Acceptance Criteria	Actual Results	Acceptance Criteria		
New Lot	FN03211402	100.1	± 3%	0.2	≤ 3%		
Previous Lot	FN080411-03	99.0	± 3%				
 Within-sample and by verified by analysis. 	petween-sample homogeneity	of the New Lot is ensure a across the lot using a ra	d through rigorous production	process controls statistica	ntly prepared calibration solution. Ily analyzed to evaluate risk and omogeneity. % RSD results shown		

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:Salicylic Acid-D4Material Lot:PN070711-03	Chemical Formula CAS Number: Molecular Weight:	7864	5-17-0	
Material Cha	racterization Summary			
Analytical Test	Method	Method Results		
Primary Chromatographic Purity by HPLC/UV Analysis	SP10-0102	99.9%		
Secondary Purity Analysis by Thin Layer Chromatography	SP10-0106	Single Spot, $R_f = 0.38$		
Identity by LC/MS Analysis	SP10-0107	Consistent with Structure		
		0.00% D ₀ vs D ₄		
Isotopic Purity by LC/MS SIM Analysis	SP10-0107	0.00% D_0 to D_1	7.27% D ₃	
		0.22% D ₂	92.51% D ₄	
Identity by ¹ H-NMR Analysis	USP <761>, SP10-0116	Consistent with Structur		
Residual Solvent Analysis by GC/FID Headspace	AM1087 ¹	None Detected		
Residual Water Analysis by Karl Fischer Coulometry	USP <921>, SP10-0103	0.01%		
Inorganic Content by Microash Analysis	SP10-0135	< 0.2%		
Purity Factor		99.9	4%	

• 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 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.

Validated analytical method

Spectral and Physical Data

HPLC/UV



¹H NMR

Instrument: Solvent:





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Spectral and Physical Data (cont.)





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Spectral and Physical Data (cont.)

LC/MS

olumn: fobile Phase: fow Rate: can Range:	Zorbax Eclip 0.1% Formic Acetonitrile 0.4 mL/min 50-1200 amu	e acid in Wate (70::30)				1 1 5	Ionization: Data File N Instrument Sample Na Acquired:	lame: t: me:	W07141	XEVO G2 (11-03		
Salicylic acid- W07141133 10	-D4 2 (1.008) Cm (9 141.0492	5:117-(68:90+ 2	127:158))	Pf	RM-319_P	N07071	11-03				I-Jul-20111 Cone Volta collision Ene 1: TOF	ige 25 ergy 6
			eoretical [M and [M - H]									
%-												
				305.08	375							
97	140.0427 ¹⁴² .0591	.0523		304.08103	<u>06.0905</u>			469.125	4		633.1639	••• m/:

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Spectral and Physical Data (cont.)

Isotopic Purity by LC/MS SIM Analysis





Stability

Short term stability studies have been performed under accelerated conditions for a period of up to four weeks. Short term data is utilized to predict long term stability and 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.

Storage Condition	Mean Kinetic Temperature (MKT)	Time Period/Result		
Freezer	-15°C			
Refrigerator	4°C	No decrease in purity was noted after four weeks.		
Room Temperature	21°C	No decrease in purity was noted after rour weeks		
40°C	40°C			
Transport/Shipping : Stab	ility data supports transport of this produc	ct at ambient conditions.		
Short Term Storage: Stab	ility data supports short term storage up to	o 3 months at Refrigerate conditions.		
Long Term Stability: Long	g term stability has been assessed for Free	zer storage (-10 °C to -25 °C) conditions.		
	5 months has been established through rea			

COA Revision History

Revision No.	Date		Reason for Revision
00	May 7, 2014	Initial version	