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The results of the tests, calibrations and/or measurements included in this document are traceable to Australia/national standards. NATA is a signatory to the APLAC Mutual Recognition Arrangement for the mutual recognition of the equivalence of reference materials certificates.

	eference Material Product Information Sheet tem conforms to ISO9001:2015 as certified by ECAAS Pty Ltd - Certification number 616061.				
Name	<i>N</i> -(2,6-dimethylphenyl)acetamide				
BP Name	Lidocaine Impurity C				
Synonym(s)	2',6'-Acetoxylidide, 2',6'-Dimethylacetanilide, N-Acetyl-2,6-xylidine				
Epichem Item #	EPL-AA239 Batch 1				
CAS #	2198-53-0				
Molecular Formula	C ₁₀ H ₁₃ NO				
Molecular Weight	163.22 g/mol				
Appearance	White powder				
Melting Point	180.6-182.1°C				
Combustion Analysis	Required (%): C:73.6, H:8.0, N:8.6. Found (%): C:73.2, H:8.0, N:8.9.				
Purity*	99.8%				
Date of Manufacture	27 June 2019				
Storage Requirements	Protect from heat, light and moisture.				
Special Precautions	This compound is for laboratory use only. Its toxicological properties may not have been fully established. It should be handled only by suitably qualified personnel.				
Intended Use	This compound is suitable for the identification of impurities and degradants in pharmaceutical materials. The purity assay is considered as relative contribution.				
Date of Shipment	ТВА				
	This certificate is valid for one year from the date of shipment provided the substance is unopened and stored under the recommended conditions.				
Retest Date	TBA (Proper Storage and Handling Required)				

* NATA accreditation does not cover the performance of this service

EPL-AA239 Batch 1

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I. Identity

The identity of this product was established using the following analyses:

Ia. ¹HNMR Spectrum

Conditions: 400 MHz, DMSO-d₆

¹HNMR spectrum consistent with chemical structure.

Acquisition Time (sec)	3,7547	Comment	LBC364-101@2F	H 1H DMSO {E:\dataextern	nal\epichem} cvgo	h 19			
Date	16 Jul 2019 17:2			Date Stamp	16 Jul 2019 17:				
ile Name		mpany\NMR files\LBC364	LBC364-101@2H*			Frequency (MHz)	400.13		
lucleus	1H	Number of Transients		Origin	spect	Original Points Count	24038		
wner	nmr	Points Count	32768	Pulse Sequence	zg	Receiver Gain	71.80		
W(cyclical) (Hz)	6402.05	Solvent	DMSO-d6	Spectrum Offset (Hz)	2795.8198	Spectrum Type	STANDARD		
weep Width (Hz)	6401.85	Temperature (degree C	24.996						
			20	+0°2	R		2.49		
			P			3.32	7	1	
							l_		

Consistent with structure, rotamers are observed (approx. 30:1). No residual solvent impurity signals visible.

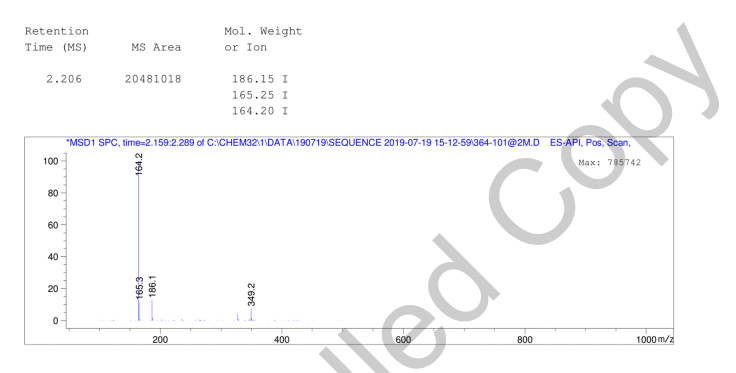
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Ib. Mass Spectrum

The mass spectrum of this material was analysed by Liquid Chromatography Mass Spectroscopy (LCMS) using inhouse EM005.WI08.

Method: ACN/water gradient (+ 0.1% formic acid).

ZORBAX SB-C8, 4.6 x 30 mm, 3.5 micron.



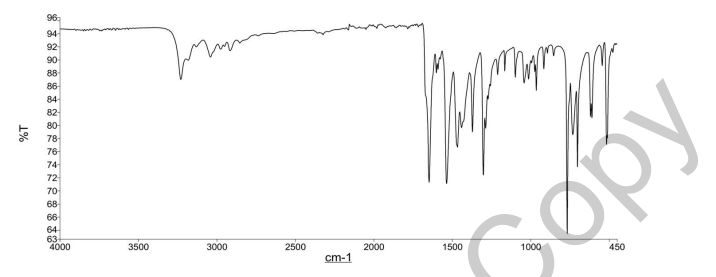
Theoretical value: 164.2 [M+H]⁺.

The signal of the Mass Spectrum is consistent with the theoretical value and its interpretation is consistent with the structural formula.

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Ic. IR Spectrum

The infra-red spectrum of this material was analysed by Fourier-Transform Infrared Spectroscopy (FTIR) using inhouse EM005.WI09.



The interpretation of the signals of the Fourier-Transform Infrared Spectrum is consistent with the structural formula.

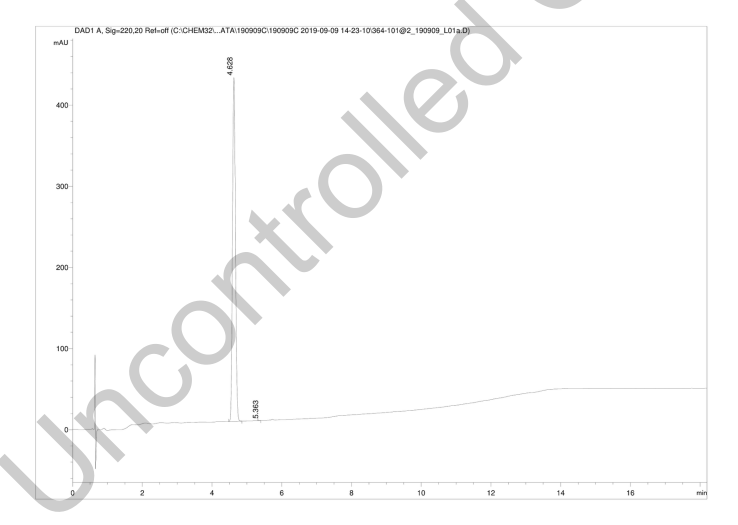
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II. Purity

The purity of this material was analysed by high performance liquid chromatography (HPLC) using in-house EM005.WI07.

HPLC Conditions:

Column	Conditions				Detector	Injector
Agilent Poroshell	45°C					Auto
120 EC C-18 4.6x50 mm	Time (min)	% Line A (Water + 0.1% (v/v) TFA)	% Line B (Acetonitrile + 0.1% (v/v) TFA)	Flow rate (mL/min)	220nm	1.0 μL 1.0mg/mL in 50% water / 50% acetonitrile (+0.1%
4.0x30 mm	0.00	95	5	1.0		(v/v) TFA)
2.7 micron	2.7 micron 6.00	65	35	1.0		
	12.00	5	95	1.0		
	17.00	5	95	1.0		
	18.00	95	5	1.0		
	21.00	95	5	1.0		



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Area Percent Report – Sorted by Signal

Peak Number	Retention Time (rounded)	Area	Area % (rounded)
1	4.63	2484.73	99.99
2	5.36	0.21	0.01
Totals			100 (rounded)

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

Results:

Average

100% (average of 10 duplicate analyses)

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III. Water Content

Method: Karl-Fischer titration using in-house EM005.WI04.

Results:

Average 0.1%

IV. Ash Content

Method: BP 2019 Ash Appendix XIJ Method II

Result:

Contains 0.1% ash.

V. Residual Solvents

Method: ¹HNMR

Result:

<0.1% by ¹H NMR analysis.

VI. Final Result

Chromatographic purity (HPLC)	100%
Water content	0.1%
Ash content	0.1%
Residual solvents	<0.1%
Purity*	99.8%

This purity is assessed to be 99.8%.

Product Reviewed By:

Product Released By:

John Moursounidis, PhD Head Reference Standards Boon Tan Quality Manager Release Date: 21 October 2019

**NATA accreditation does not cover the performance of this service.* The calculation of the purity follows the formula:

 $Purity(\%) = \frac{((Chromatographicpurity[HPLC])x(100 - (watercontent + ashcontent + volatilecontents)))}{100}$

100

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