



CERTIFICATE OF ANALYSIS

REPORT ONE-CI10-091426
VERIFICATION CODE 3V8539G3
VERIFY AT UNITEDLABORATORIES.ORG

Client and sample information

CLIENT	One Aminos LLC	PRODUCT	CJC-1295 + Ipamorelin
ACTIVE IDENTITY	5 mg CJC-1295 no DAC + 5 mg Ipamorelin	LABELED CONTENT	5+5 mg
LABORATORY SAMPLE ID	ONE-CI10	REPORT NUMBER	ONE-CI10-091426
BATCH / LOT	ONE-CI10-091426	DATE CERTIFIED	09/14/2026
SAMPLE MATRIX	Lyophilized Powder	APPEARANCE	Conforms
VERIFICATION CODE	3V8539G3	OVERALL RESULT	PASS

7X certificate summary

Test	Method	Specification	Result	Status
Identity	LC-MS	C152H252N44O42 + C38H49N9O5	5 mg CJC-1295 no DAC + 5 mg Ipamorelin	PASS
Purity	RP-HPLC	>= 98.0%	99.54%	PASS
Net content	Quantitative HPLC	Report result	10.93 mg	REPORTED
Vial conformity	HPLC conformity	Consistent vials	3 vials conform	PASS
Heavy metals	ICP-MS	Analyte limits	Below limits	PASS
Sterility	Rapid sterility	No growth	No growth	PASS
Endotoxin	USP <85>	<= 0.5 EU/mL	< 0.05 EU/mL	PASS

OVERALL RESULT: PASS - REVIEWED AND APPROVED FOR ISSUE



IDENTITY, PURITY AND CONTENT

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METHOD	RP-HPLC	COLUMN	C18
DETECTOR	DAD	DETECTION	214 nm

Analytical results

Analyte	Specification	Result	Unit	Status
Peptide purity (HPLC)	>= 98.0%	99.54	%	PASS
Net content	Report result	10.93	mg	REPORTED
Identity (LC-MS)	C152H252N44O42 + C38H49N9O5	5 mg CJC-1295 no DAC + 5 mg Ipamorelin	-	PASS

HPLC conformity testing

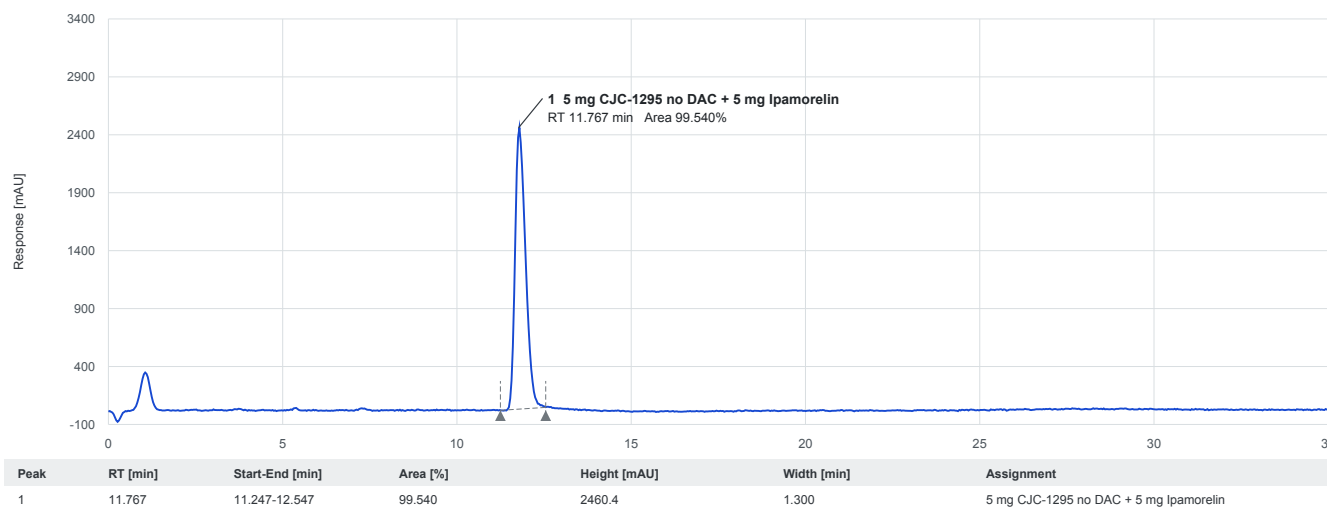
Sample	Purity	Net content	Identity	Result
Vial 1	99.570%	11.01 mg	Confirmed	PASS
Vial 2	99.557%	10.83 mg	Confirmed	PASS
Vial 3	99.493%	10.95 mg	Confirmed	PASS
Conformity mean	99.540%	10.93 mg	-	-

HPLC chromatogram

CHROMATOGRAPHY DATA REVIEW - DAD SIGNAL

Sample ID: ONE-CI10 Report: ONE-CI10-091426 Method: RP-HPLC / C18

Channel: DAD1 A, Sig=214.4 Ref=off Run window: 0.000-35.000 min



Reconstructed HPLC chromatogram - CJC-1295 + Ipamorelin, 5+5 mg



MICROBIOLOGY AND ENDOTOXIN

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Sterility screening

Test	Method	Specification	Result	Status
Bacteria (aerobic/anaerobic)	Rapid sterility	No growth	No growth	PASS
Fungi / yeast	Rapid sterility	Not detected	Not detected	PASS

INCUBATION PERIOD	2 days	TEMPERATURE	30-35 °C
SAMPLE MATRIX	Lyophilized Powder	FINAL DETERMINATION	PASS

STERILITY RESULT: NO GROWTH / NOT DETECTED - PASS

Bacterial endotoxin - USP <85>

TEST PERFORMED	Quantitative bacterial endotoxin	ASSAY	Kinetic LAL
DETECTION RANGE	0.01-1.0 EU/mL	STANDARD	E. coli O111:B4
DILUTION VOLUME	2.0 mL	MATRIX	LAL reagent water
SPECIFICATION BASIS	Supplied acceptance criterion	COMPENDIAL REFERENCE	USP <85>

Parameter	Result
Endotoxin level	< 0.05 EU/mL
Acceptance limit	<= 0.5 EU/mL
Sample CV	4.4%
Spike CV	4.2%
Spike recovery	106%
Final determination	PASS

Control	Expected	Observed	Status
Positive control	Detectable	As expected	PASS
Negative control	No signal	As expected	PASS

ENDOTOXIN RESULT: PASS



HEAVY METALS AND AUTHORISATION

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Elemental impurities - ICP-MS

Element	Result (ppm)	Acceptance limit (ppm)	Status
Lead (Pb)	< 0.5	<= 10	PASS
Arsenic (As)	< 0.15	<= 1.5	PASS
Cadmium (Cd)	< 0.05	<= 0.5	PASS
Mercury (Hg)	< 0.3	<= 3	PASS

Method and quality controls

SAMPLE PREPARATION	HNO3 / H2O2 matrix	CALIBRATION	Multi-element standard curve
INTERNAL STANDARDS	Sc, Ge, In, Bi	METHOD BLANK	PASS
CONTINUING CALIBRATION	PASS	DUPLICATE RPD	< 20%
SPECIFICATION BASIS	Supplied reporting limits	INSTRUMENT	ICP-MS

Element	Spike level	Recovery	Criterion
Lead	5 ppm	98%	70-150%
Arsenic	0.75 ppm	102%	70-150%
Cadmium	0.25 ppm	95%	70-150%
Mercury	1.5 ppm	91%	70-150%

ELEMENTAL IMPURITIES: ALL ANALYTES BELOW ACCEPTANCE LIMITS - PASS

Laboratory authorisation

Approved and authorised for issue

Bhavrat Singh
Owner, Analyst and Head Lab Director

Report number: ONE-CI10-091426
Batch / lot: ONE-CI10-091426
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Date certified: 09/14/2026



Authorised signature - Bhavrat Singh