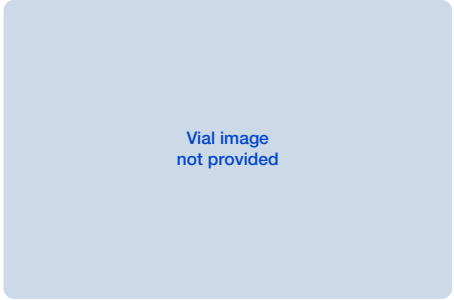


ORDERED BY  ZYCARE PEPTIDES.COM Your Peptide Brand	PRODUCT NAME	SAMPLE
	SKU CATEGORY	BATCH TESTED
	Tesamorelin + Ipamorelin 8mg Blend 8 mg SQC.906 Identify, Purity, Assay, Heavy metals and Microbial.	20254567L07CIU100 2026-04-18

DESCRIPTION	Vial with white powder sealed with plastic security cap. Flag tag attached at neck with manufacturer / product SKU number, PO, and expiration
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PERFORMING LAB	Analytical Formulations, Inc. LAB Zycare Peptides Link Zycarepeptide.com Lab certified result data: 2026-04-24
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Vial image not provided

Submitted vial — YPB.222



DIGITALLY SIGNED BY
Zycare Peptides

ISSUED	2026-04-29
ISSUER	Sectigo RSA Document Signing CA
ALGORITHM	RSA-SHA512
FIELD	Purity Signature
TYPE	adbe.pkcs.deatached
CERT SERIAL	10251389K04CC6010
CERT SHA-256	T3467E58CFKE3JK9.....AF9CBFFB76366623
TIMESTAMP	Sectigo RFC 3161 TSA

Click this panel in Adobe Reader to validate the embedded signature. Cryptographic validity is also shown at [verify.zycarepeptides.com](#).

ANALYTICAL RESULTS

COMPONENT	ANALYTE	SPECIFICATION	RESULT	P/F
Tesamorelin + Ipamorelin 8 mg Blend	Qualitative ID (Lambda Max)	TS λ max is a match compared to its characteristic reference standard.	Matches	✓
	Percent Purity	NLT 98% $r_{xy} = \frac{\sum (x_i - \bar{x})(y_i - \bar{y})}{\sqrt{\sum (x_i - \bar{x})^2} \cdot \sqrt{\sum (y_i - \bar{y})^2}}$	99.79%	✓
Tesamorelin + Ipamorelin 8 mg Blend	Quantitative Assay	NLT 95% label claim $\% = \frac{A_{TS}}{A_{STD}} \times \frac{C_{STD}}{C_{TS}} \times 100$	8.96 MG	✓
	Heavy Metals (Total Quantity)	NMT 150 ppb/vial (including Pb, Cd, Hg, Ni, Fe & Co)	<8 ppb	✓
	TAMC (Aerobic/Coliform)	NMT 1,000 CFU · Incubated 48 hrs @ 37°C	0 CFU	✓
	TYMC (Yeast & Molds)	NMT 100 CFU · Incubated 48 hrs @ 30°C	0 CFU	✓

METHODS

- Qualitative ID - Lambda Max
- Percent Purity - Correlation Coefficient
- Quantitative Assay - Beer-Lambert
- Heavy Metals - Total Quantity
- TAMC (Total Aerobic Microbial Counts)
- TYMC (Total Yeast and Mold Counts)

Microbial specifications follow the Substances for Pharmaceutical Use compendium. UV-Vis measurements performed on a Jenway 6715 UV/Vis Spectrophotometer.

PERFORMING LABORATORY & CERTIFICATION

PERFORMING LABORATORY

Analytical Formulations, Inc.

8940 Fourwinds Drive STE 107, Windcrest, TX 78239

<https://analyticalformulations.com>

A laboratory-authorized user certified on 2026-06-02 that the submitted result data and the Purity Analytics COA generated from that data accurately represent the laboratory's reported findings for the sample identified in this report.

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Laboratory role.

The performing laboratory identified above performed or reported the analytical measurements reflected in this Certificate of Analysis for the submitted sample only. The laboratory's role is limited to analytical testing and reporting of the sample as received. The laboratory does not manufacture, package, relabel, market, prescribe, recommend, dispense, distribute, or sell the product identified in this report.

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