



## Certificate of Analysis (CoA)

Date: 7/15/2026 Date Tested: 6/29/2026  
Customer: Peptides Optimized LLC Testing material: GHK-Cu  
Batch #: GK50-260629-B BT Sample ID: 005000396442597  
Labeled Peptide 50 mg Storage: R.T.  
Content/Potency:  
Visual Description: Small clear vial: white lyophilized powder, pink label, silver crimp, blue plastic cap.  
Labeled as: GHK-Cu Manufacturer: N/A  
Testing Purpose: FTIR and HPLC analysis for the identification, purity, potency and composition of a peptide product. It additionally reports elemental impurities (heavy metals), bacterial endotoxins and sterility as recorded in this certificate. It does not provide information on particulate matter.



| Test                                         | Method             | Specification                                                                  | Result                                                       | Status      |
|----------------------------------------------|--------------------|--------------------------------------------------------------------------------|--------------------------------------------------------------|-------------|
| General Appearance                           | Visual             | White to off-white powder                                                      | white powder                                                 | <b>PASS</b> |
| HPLC Purity of Peptide Assay                 | USP <621>          | NLT 98.0 % (area normalisation)                                                | 99.5 %                                                       | <b>PASS</b> |
| HPLC Potency Assay                           | USP <621>          | 50 mg (net peptide), 90.0 - 110.0 %                                            | 50.1 mg (100.2 %)                                            | <b>PASS</b> |
| Identity by Mass Spectrometry                | LC-MS (ESI)        | Observed mass within 1 Da of theoretical                                       | Conforms; observed mass matches GHK-Cu reference standard.   | <b>PASS</b> |
| FTIR Identification and Composition Analysis | USP <197A>         | Sample spectrum should confirm the content of peptide via characteristic bands | FTIR spectrum consistent with the GHK-Cu reference standard. | <b>PASS</b> |
| Heavy Metals (Elemental Impurities)          | USP <233> (ICP-MS) | Class 1 (Pb, Cd, As, Hg): meets USP <232>/ICH Q3D                              | Complies (< LOQ)                                             | <b>PASS</b> |

The results of the CoA relate only to the item(s) tested and applied to the sample as received.

COA # 1527140

ACCESS CODE **TAZ7FNC49SGV**

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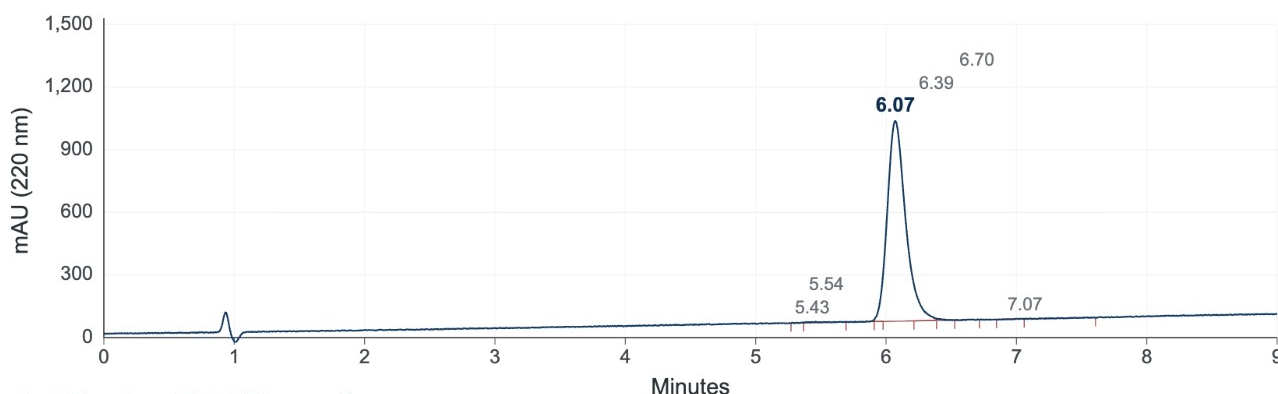
# Certificate of Analysis (CoA)

**Product:** GHK-Cu **Batch #:** GK50-260629-B **Content:** 50 mg **Tested:** 6/29/2026

## HPLC Purity Chromatogram @ 220 nm: GHK-Cu Lot GK50-260629-B

**Sample Name:** GHK-Cu 50mg  
**Sample ID / Lot:** GK50-260629-B  
**Method:** PEP\_TFA\_220\_GRAD.M  
**Column:** Kinetex EVO C18, 4.6 x 100 mm, 2.6 um  
**Signal:** VWD1 A, 220 nm  
**Mobile Phase:** A 0.1% TFA/H<sub>2</sub>O, B 0.1% TFA/MeCN (gradient)

**Injection Volume:** 10.0 uL  
**Acquired:** 6/29/2026 09:09  
**Vial:** 34 **Injection:** 1  
**Operator:** A. Ivanova  
**Flow:** 1.2 mL/min **Column Temp:** 30 C  
**System:** HPLC-02



### Peak Results - USP <621>, area %

| Pk           | RT (min) | Area (mAU*s) | Height (mAU) | Area %        | USP Res. | USP Tailing | USP Plates |
|--------------|----------|--------------|--------------|---------------|----------|-------------|------------|
| 1            | 5.43     | 18           | 1.9          | 0.19          | -        | -           | -          |
| 2            | 5.54     | 5            | 0.5          | 0.06          | -        | -           | -          |
| 3            | 6.07     | 9,435        | 959          | 99.50         | 1.98     | 1.33        | 9,494      |
| 4            | 6.39     | 5            | 0.5          | 0.05          | -        | -           | -          |
| 5            | 6.70     | 7            | 0.7          | 0.08          | -        | -           | -          |
| 6            | 7.07     | 13           | 0.9          | 0.13          | -        | -           | -          |
| <b>Total</b> |          | <b>9,482</b> |              | <b>100.00</b> |          |             |            |

OpenLab CDS | Processing method: USP\_PURITY\_AREAPCT | Result: main peak 99.5 % (area) | Page 1 of 1



## Certificate of Analysis (CoA)

**Product:** GHK-Cu    **Batch #:** GK50-260629-B    **Content:** 50 mg    **Tested:** 6/29/2026

### Heavy Metals Analysis (ICP-MS, USP <233>)

| Element       | Specification | Result       | Status |
|---------------|---------------|--------------|--------|
| Arsenic (As)  | NMT 1.5 ppm   | Not Detected | PASS   |
| Cadmium (Cd)  | NMT 0.5 ppm   | Not Detected | PASS   |
| Mercury (Hg)  | NMT 1.5 ppm   | Not Detected | PASS   |
| Lead (Pb)     | NMT 1 ppm     | Not Detected | PASS   |
| Chromium (Cr) | NMT 10 ppm    | Not Detected | PASS   |

Elemental impurities determined by ICP-MS. Limits are laboratory screening limits for research-use material, derived from ICH Q3D permitted daily exposures for the parenteral route.

| Test                      | Method   | Specification                | Result | Status |
|---------------------------|----------|------------------------------|--------|--------|
| Bacterial Endotoxins Test | USP <85> | NMT 40 EU/vial reconstituted | PASS   | PASS   |

*The results of the CoA relate only to the item(s) tested and applied to the sample as received.*

### Bacterial Endotoxins Test GHK-Cu Lot GK50-260629-B

| Sample ID     | Description | Volume (mL) | Endotoxin Limit (EU/vial) | Lysate Sensitivity ( $\lambda$ , EU/mL) | Result      |
|---------------|-------------|-------------|---------------------------|-----------------------------------------|-------------|
| GK50-260629-B | GHK-Cu      | 3           | 40 EU/vial                | 0.25                                    | .07 EU/vial |

The sample passed a valid gel clot endotoxin test based on the type, mode of administration and dosage.

### Sterility Test

| Test      | Method   | Specification       | Result | Status |
|-----------|----------|---------------------|--------|--------|
| Sterility | USP <71> | No microbial growth | PASS   | PASS   |

| Medium                              | Incubation Temp. | Duration | Units Tested | Observation |
|-------------------------------------|------------------|----------|--------------|-------------|
| Fluid Thioglycollate Medium (FTM)   | 30 - 35 °C       | 14 days  | 2            | No growth   |
| Soybean-Casein Digest Medium (SCDM) | 20 - 25 °C       | 14 days  | 2            | No growth   |

No microbial growth was observed. The sample meets the requirements of the sterility test (USP <71>).

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### Notes & Methodology

1. Peptide purity determined by RP-HPLC area normalisation at 220 nm. The value represents the percentage of the target peptide relative to all peptide-related peaks. Non-peptide process-related impurities, where detected, are excluded from the calculation.
2. Net peptide content (potency) calculated as assay content corrected for water (Karl Fischer) and counterion content, expressed as a percentage of the labelled claim.
3. Identity confirmed by FTIR against a reference standard (USP <197A>) and by chromatographic retention time comparison.
4. Elemental impurities screened by ICP-MS per USP <233> methodology. Acceptance criteria are laboratory screening limits for research-use material and do not constitute evaluation against a specific pharmacopoeial monograph.
5. Bacterial endotoxins determined by gel-clot method per USP <85>. Sterility per USP <71>.
6. The chromatogram reproduced in this certificate is representative of the batch tested.