

Express Biotech International

4650 Wedgewood Blvd., STE 103 Frederick, MD 21703 USA Tel: +001.301.228.2444

Fax: +001.301.560.6570

Email: info@xpressbio.com

COL G (Recombinant Collagenase Class I) COL H (Recombinant Collagenase Class II)

For Research Use Only

CATALOG NUMBERS

CG-001 – COL G, 75 U	CH-001 – COL H, 750 U
CG-002 – COL G, 300 U	CH-002 - COL H, 3000 U
CG-003 - COL G, 750 U	CH-003 - COL H, 7500 U

DESCRIPTION

COL G and COL H are recombinant collagenases (metalloproteinases) class I and class II respectively [1]. COL G and COL H are synthesized separately from *C. Histolyticum* genes by DNA recombination in *E. Coli BL21 AI* strain, bearing a Maltose Binding Protein (MBP) tag at the N-terminal end [2].

COL G and COL H are affinity chromatography purified proteins, highly pure, highly stable, lot-to-lot consistent, endotoxin-free (≤ 10 EU/mg, LAL assay) and animal-free.

CAS: 9001-12-1

EC: 3.4.24.3

Grade: Research Premium Grade

Form: Lyophilized white powder

Quality: Amylose Affinity Chromatography

Inhibitors: EDTA, EGTA, Cys, Hys, DTT, 2-

mercaptoethanol

Activators: Ca²⁺

Their molecular weights are ~135 kDa (COL G) and ~158.5 kDa (COL H). COL G and COL H are soluble in water or aqueous buffers and express their maximum activity at pH 8.

SUBSTRATES

COL G and COL H play different synergic roles in collagen digestion. COL G expresses a higher activity against native collagen, specifically hydrolyzing 3D-helix regions, while COL H expresses a lower activity against the 3D helix and a higher activity against linear collagen regions at the motif Pro-Y-Gly-Pro [3,4]. The mix of COL G and COL H expresses a synergic activity that results in efficient collagen digestion [5].

For tissue dissociation, protease addition is needed to hydrolyze non-collagenous proteins and other macromolecules present in the extracellular matrix [6].

ENZYMATIC ACTIVITY

COL G ≥ 3.0 Units/mg*

COL H ≥ 30.0 Units/mg*

*according to Grassmann, one Unit liberates 1 µmol of Gly-Pro-Ala from Carbobenzoxy-Gly-Pro-Gly-Gly-Pro-Ala-OH (Fluka 27673) in 1 min at pH 7.4, 37 °C [7].



Express Biotech International

4650 Wedgewood Blvd., STE 103 Frederick, MD 21703 USA Tel: +001.301.228.2444 Fax: +001.301.560.6570 Email: info@xpressbio.com

APPLICATIONS

For research use only.

Due to their high purity and specificity, COL G and COL H are especially indicated for the isolation of primary cells from liver, pancreas, heart, and cartilage, and stem cells from adipose tissue and others.

In these applications we recommend using a combination of COL G and COL H in a specific activity ratio, or according to the relevant isolation protocol in order to obtain an optimal collagen digestion in cell isolation. For other applications or suggestions, contact xpressbio.com or visit www.xpressbio.com.

PREPARATION METHOD

We recommend reconstituting the lyophilized COL G and COL H enzymes in the tissue-dissociation buffer by injecting the **buffer directly into the vial**. Do not exceed an enzyme concentration of 30 U/ml (COL G) or 300 U/ml (COL H) to avoid precipitates.

Keep the vial on ice and periodically shake until the enzyme is completely dissolved. Filter with 0.22 μm mesh for sterility.

Prepare a mix of COL G and COL H solutions in a specific activity ratio and dilute according to your protocol working solution concentration.

Add protease to the mix at 4 °C according to the specific application. Thermolysin, pronase or neutral protease/dispase can be normally used. **Protease must be added immediately before use** to avoid catalytic processes in the enzymatic blend. The amount of protease will define the aggressiveness of your enzyme mixture. For suggestions about your specific protocol and application please contact xpressbio@xpressbio.com or visit www.xpressbio.com.

STORAGE AND STABILITY

Lyophilized COL G and COL H are stable at -80 °C up to two years. We recommend splitting in aliquots the reconstituted solutions at need and storing them at -20 °C up to one month or -80 °C up to 6 months.

To use aliquots later on, they can be diluted in re-constitutive buffer or can be directly added into the enzyme working solution.

▲ Warning: We recommend avoiding multiple freeze-thaw cycles and exposure to frequent temperature changes. REFERENCES

- [1] Matsushita, O. et al. (1999) J. Bacteriol. 181(3): 923-933.
- [2] Salamone, M. et al. (2012) Chem. Eng. Trans. 27: 259-264.
- [3] Philominathan S.T. et al. (2009) J. Biol. Chem. 284(16): 10868-10876;
- [4] Matsushita, O. Et al. (1994) J. Bacteriol.176: 149-156
- [5] Breite, A.G. et al. (2011) Transplant Proc. 43(9): 3171-3175
- [6] Salamone, M. et al. (2014) Chem. Eng. Trans. 38: 247-252.
- [7] W. Grassmann, et al, (1960) Z. Physiol.Chemie 322:267