

Catalog Number: HZ-1320-GMP

## **Data Sheet**



## GMP HumanKine® IL-1 alpha (Recombinant Human)

**Animal Component-Free** 

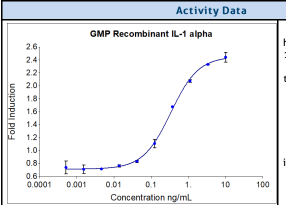
**Human cell expressed** 

Tag-Free

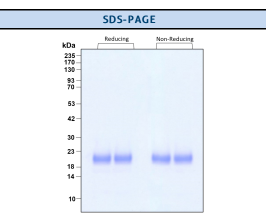
**Endotoxin Free** 

Product Description				
Alternative Names	Hematopoietin 1, Pro-interleukin-1-alpha, IL1 alpha, IL1, IL1A, IL-1A, IL1-ALPHA, IL1F1, IL-1F1, preinterleukin 1 alpha, IL1F1hematopoietin-1, interleukin 1, alpha, interleukin-1 alpha, LAF, LEM			
Accession Number	P01583			
Source	Human Embryonic Kidney cells (HEK293). HEK293-derived IL-1 alpha protein			
Species Reactivity	human, mouse			
Adventitious Virus	Master Cell Bank tested Negative for Adventitious Viruses			

Specifications					
Test	Method	Specification			
Activity	Does-dependent proliferation of the D10.G4.1 mouse helper t lymphocyte cell line.	0.125-1.25 ng/mL in D10.G4.1 cells			
Molecular Mass	SDS-PAGE	22 kDa reduced and non-reduced, monomer, glycosylated			
Purity	SDS-PAGE	>95%			
Endotoxin	LAL	LAL <0.1 EU/μg			
Mycoplasma	PCR	Not Detected			



GMP-grade recombinant human IL-1 alpha (Cat no: HZ-1320-GMP) stimulates doesdependent proliferation of the D10.G4.1 mouse helper tlymphocyte cell line. Viable cell number was quantitatively assessed by PrestoBlue® Cell Viability Reagent. D10.G4.1 cells were treated with increasing concentrations of recombinant IL-1 alpha for 72 hours. The EC50 was determined using a 4-



Purity of GMP-grade recombinant human IL-1 alpha was determined by SDS- polyacrylamide gel electrophoresis. The protein was resolved in an SDS- polyacrylamide gel in reducing and non-reducing conditions and stained using Coomassie blue.

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Document #: FR-QA118-101 Rev 0 Data Sheet Version #: Proteintech Group, Inc.

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	Preparation				
Shipping Temperature ambient temperature					
Formulation 1x PBS, See Certificate of Analysis for details					
Reconstitution  Briefly centrifuge the vial before opening. It is recommended to reconstitute the protein to 0.2 mg/mL in sterile 2 containing 0.1% endotoxin-free recombinant human serum albumin (HSA). Gently swirl or tap vial to make the protein to 0.2 mg/mL in sterile 2 containing 0.1% endotoxin-free recombinant human serum albumin (HSA).					

	Product Form	Temperature Conditions	Storage Time (From Date of Receipt)	
	Lyophilized	-20°C to -80°C	Until Expiry Date	
Stability and Storage	Lyophilized	Room Temperature	2 weeks	
	Reconstituted as per CofA	-20°C to -80°C	6 months	
	Reconstituted as per CofA	4°C	1 week	
	Avoid repeated freeze-thaw cycles.			

## Proteintech GMP Quality Policy HumanKine® GMP Proteins

Invitro recombinant protein production can be prone to variability due to various reasons ranging from quality of raw materials to inconsistency in the process. Therefore, HumanKine®, a proteintech brand's GMP proteins are produced and tested under an ISO 13485 certified quality management system in a clean room facility. Proteintech manufactures the GMP HumanKine® products according to the applicable sections in the following documents: USP Chapter 1043 (Ancillary Materials for Cell, Gene, and Tissue-Engineered Products, USP Chapter 92 (Growth Factors and Cytokines Used in Cell Therapy Manufacturing), WHO TRS, No. 822, 1992 Annex 1 (Good Manufacturing Practices for Biological Products), Ph. Eur. General Chapter 5.2.12, and EudraLex – Volume 4 – Part IV (Guidelines on GMP specific to ATMPs). Proteintech strives to achieve the utmost quality GMP raw material ensuring all applicable guidelines are taken into consideration.

The QMS is built to provide our customers with consistent and pure product delivered by documented processes, qualified personnel, validated processes, qualified equipment, qualified vendors, and a stringent final product release process. Although the final product release process is important, Proteintech performs in-process QC steps after each major manufacturing stage. Production records and facilities may be available for an inspection by approved personnel.

Our GMP policy covers all the aspects of production; from raw materials, facility, equipment, and Instruments to training and personal hygiene of staff. It also guarantees that the process is explicit, validated and well documented for transparency and traceability.

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