Data Sheet



CellGenix[®] Recombinant Human Interleukin-1 beta (rh IL-1β) Preclinical Grade - Order No.: 1411-010 (10 μg), 1411-050 (50 μg)

Product Characteristics

Source	E. coli
Description	Human Interleukin-1 beta, accession # P01584, Ala117-Ser269 N-terminal Met Molecular mass 17.5 kDa
Formulation	Lyophilized from a 0.2 μ m-filtered solution containing 1.5 mM potassium phosphate, 8.1 mM sodium phosphate, 2.7 mM potassium chloride, and 137 mM sodium chloride, pH 7.5.
Intended use	For preclinical ex vivo use. Not intended for therapeutic use.

Quality Parameters

Activity	$\geq 100 \times 10^6$ IU/mg calibrated against NIBSC #86/680 Measured in a cell proliferation assay using an IL-1 β -dependent cell line, RPMI-1788
Purity	\geq 95 %, as determined by SDS-PAGE (under reducing and non-reducing conditions, visualized by silver staining)
Endotoxin	< 1000 EU/mg, as determined by LAL gel clot test
Sterility	Sterility test of the vialed product (direct inoculation)
Mass per vial	1411-010: 10 μg, 1411-050: 50 μg
Animal-derived component-free	ADCF Level 2: The final product contains neither animal- nor human-derived materials. ADCF Level 2 cytokines are produced in our dedicated animal-free facility. No animal-derived components are used throughout the complete production process. All ADCF Level 2 cytokines are produced in <i>E. coli</i> .



Shipment & Storage

Transport	Ambient temperature. Please refer to Technote (www.cellgenix.com)
Expiry	3 years from date of shipment
Storage & Stability	Store lyophilized cytokine at -20°C to -80 °C. Avoid repeated freeze/thaw cycles.

Handling Instructions

Reconstitution	Recommended in sterile water to a final concentration of 100 μ g/ml (for 10 μ g vials) or 250 μ g/ml (for 50 μ g vials).
Dilution	Recommended in CellGenix [®] serum-free media. For dilution with protein free medium, a carrier protein (0.1-1 % albumin or 1-10 % appropriate serum) has to be included. Failure to dilute product according to these instructions may result in loss of activity.

Quality Statement

Final manufacturing steps and QC are performed in a GMP facility.

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