

### Data Sheet

# CellGenix® Recombinant Human Thrombopoietin (rh TPO)

Preclinical Grade - Order No.: 1417-010 (10 μg), 1417-050 (50 μg)

#### **Product Characteristics**

Source E. coli

**Description** Truncated human TPO, accession # P40225, Ser22-Leu195

N-terminal Met and 6xHis-tag

Molecular mass 19.6 kDa

**Formulation** Lyophilized from a 0.2 μm-filtered aqueous solution.

**Intended use** For preclinical *ex vivo* use. Not intended for therapeutic use.

#### **Quality Parameters**

Activity  $\geq 10 \times 10^6 \text{ U/mg}$  calibrated against an in-house reference standard

Measured in a cell proliferation assay using a TPO-dependent cell line, MO7e

**Purity** ≥ 95 %, as determined by SDS-PAGE (under reducing and non-reducing conditions,

visualized by Coomassie staining)

Endotoxin < 1000 EU/mg, as determined by LAL gel clot test

Sterility Sterility test of the vialed product (direct inoculation)

**Mass per vial** 1417-010: 10 μg, 1417-050: 50 μg

Animal-derived ADCF Level 2: The final product contains neither animal- nor human-derived component-free materials. ADCF Level 2 cytokines are produced in our dedicated animal-free facility.

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process. All ADCF Level 2 cytokines are produced in E. coli.



### 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.

Store a 250 μg/ml cytokine solution:

• 4 weeks at 2°C to 8°C under sterile conditions after reconstitution. Store in the original container.

• 4 months at -20°C to -80°C under sterile conditions after reconstitution. Store in 60 μl aliquots in polypropylene cryogenic vials.

Avoid repeated freeze/thaw cycles.

## **Handling Instructions**

**Reconstitution** Recommended in sterile water to a final concentration of 100 μg/ml (for 10 μg vials)

or 250 μg/ml (for 50 μg 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.

