Product Name: Recombinant Human ITPase (C-6His) Catalog #: PEH0951



Summary

Name Inosine triphosphate pyrophosphatase/ITPA

Purity Greater than 95% as determined by reducing SDS-PAGE

Endotoxin level <1 EU/μg as determined by LAL test.

Construction Recombinant Human Inosine Triphosphate Pyrophosphatase is produced by

our E.coli expression system and the target gene encoding Ala2-Ala194 is

expressed with a 6His tag at the C-terminus.

Accession # Q9BY32

Host E.coli

Species Human

Predicted Molecular Mass 22.5 KDa

Formulation Supplied as a 0.2 µm filtered solution of 20mM Tris-HCl, 5% Trehalose, 300mM

NaCl, 30% Glycerol, 0.05% Tween 80, pH8.0.

Shipping The product is shipped on dry ice/polar packs. Upon receipt, store it immediately

at the temperature listed below.

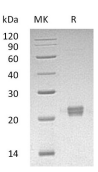
Stability&Storage Store at \leq -70°C, stable for 6 months after receipt. Store at \leq -70°C, stable for 3

months under sterile conditions after opening. Please minimize freeze-thaw

cycles.

Reconstitution

SDS-PAGE image



Background

Alternative Names Inosine Triphosphate Pyrophosphatase; ITPase; Inosine Triphosphatase; Non-

Canonical Purine NTP Pyrophosphatase; Non-Standard Purine NTP Pyrophosphatase; Nucleoside-Triphosphate Diphosphatase; Nucleoside-Triphosphate Pyrophosphatase; NTPase; Putative Oncogene Protein hlc14-06-p;

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Background

ITPA; C20orf37

Inosine Triphosphate Pyrophosphatase (ITPase) is a cytoplasmic enzyme that belongs to the HAM1 NTPase family. ITPase hydrolyzes the non-canonical purine nucleotides inosine triphosphate (ITP) and deoxyinosine triphosphate (dITP) to the monophosphate nucleotide (IMP) and diphosphate. The ITPase enzyme acts as a homodimer and does not distinguish between the deoxy- and ribose forms. ITPase probably excludes non-canonical purines from RNA and DNA precursor pools, thus preventing their incorporation into RNA and DNA and avoiding chromosomal lesions. Defects in ITPase is thought to be inherited and is characterized by an over-accumulation of ITP in erythocytes, leukocytes and fibroblasts.

Note

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