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# Recombinant Human Arginase-1/ARG1 Protein (E.coli, His Tag)

Catalog Number: PKSH032091

Note: Centrifuge before opening to ensure complete recovery of vial contents.

### Description

Species Human

Source E.coli-derived Human Arginase-1; ARG1 protein Met 1-lys322, with an C-terminal His

Calculated MW35.8 kDaObserved MW40 kDaAccessionP05089

**Bio-activity** Not validated for activity

#### **Properties**

**Purity** > 95 % as determined by reducing SDS-PAGE.

**Concentration** Subject to label value.

**Endotoxin**  $< 1.0 \text{ EU per } \mu\text{g of the protein as determined by the LAL method.}$ 

Storage Storage Store at  $< -20^{\circ}$ C, stable for 6 months. Please minimize freeze-thaw cycles.

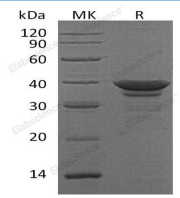
**Shipping** This product is provided as liquid. It is shipped at frozen temperature with blue ice/gel

packs. Upon receipt, store it immediately at < - 20°C.

Formulation Supplied as a 0.2 μm filtered solution of 20mM Tris-HCl, 150mM NaCl, 20% Glycerol,

1mM DTT, pH 7.4.

#### Data



> 95 % as determined by reducing SDS-PAGE.

## Background

ARGI is a member of the ureohydrolase family of enzymes. ARGI can catalyze the hydrolysis of arginine to ornithine and urea. In the urea cycle, ARGI catalyzes the fifth and final step, a series of biochemical reactions in mammals during which the body disposes of harmful ammonia. ARGI is a cytosolic enzyme and expressed widely in the liver as part of the urea cycle. Inherited deficiency of this ARGI causes argininemia, which is an autosomal recessive disorder characterized by hyperammonemia.

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