Product Name: Recombinant Human SGSH (C-6His)

Catalog #: PHH1578



Summary

Name Sulfamidase/SGSH

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

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

Construction Recombinant Human N-Sulphoglucosamine Sulphohydrolase is produced by

our Mammalian expression system and the target gene encoding Arg21-

Leu502 is expressed with a 6His tag at the C-terminus.

Accession # P51688

Host Human Cells

Species Human

Predicted Molecular Mass 55.72 KDa

Formulation Supplied as a 0.2 µm filtered solution of 20mM Tris-HCl, 150mM NaCl, 1mM

CaCl2, 10% Glycerol, pH 7.5.

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

at the temperature listed below.

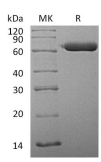
Stability&Storage Store at \leq -70°C, stable for 6 months after receipt. Store at \leq -70°C, stable for 3

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

cycles.

Reconstitution

SDS-PAGE image



Background

Alternative Names N-Sulphoglucosamine Sulphohydrolase; Sulfoglucosamine Sulfamidase;

Sulphamidase; SGSH; HSS

Background N-Sulphoglucosamine Sulphohydrolase (SGSH) is an important member of the

sulfatase family which is involved in the degradation of heparin sulfate. SGSH binds one calcium ion per subunit as a cofactor. SGSH catalyzes N-sulfo-D-glucosamine

Web: https://www.enkilife.com E-mail: order@enkilife.com techsupport@enkilife.com Tel: 0086-27-87002838

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and H2O to D-glucosamine and sulfate. SGSH deficiency is result in mucopolysaccharidosis type 3A (MPS3A), a recessive lysosomal storage disease characterized by neurological dysfunction but relatively mild somatic manifestations.

Note

For Research Use Only, Not for Diagnostic Use.

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