## Recombinant Mouse HSPD1/HSP60 Protein (His Tag)

### Catalog No. PKSM040589

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

| Synonyms60kDa;Hsp60SpeciesMouseExpression HostE.coliSequenceLeu 2-Phe 573AccessionNP_034607.3Calculated Molecular Weight62.3 kDaObserved molecular weight58 kDaTagN-HisBioactivityNot validated for activityPropertiesPurity>95 % as determined by reducing SDS-PAGE.EndotoxinPlease contact us for more information.StorageGenerally, lyophilized proteins are stable for up to 12 months when stored at -20 to | Description                 |  |
|--|-----------------------------|--|
| Expression HostE.coliExquenceLeu 2-Phe 573AccessionNP_034607.3Calculated Molecular Weight62.3 kDaObserved molecular weight58 kDaTagN-HisBioactivityNot validated for activityPropertiesPurity> 95 % as determined by reducing SDS-PAGE.PurityPlease contact us for more information.   | Synonyms                    | 60kDa;Hsp60  |
| SequenceLeu 2-Phe 573AccessionNP_034607.3Calculated Molecular Weight62.3 kDaObserved molecular weight58 kDaTagN-HisBioactivityNot validated for activityPropertiesPurity>95 % as determined by reducing SDS-PAGE.PuntyPlase contact us for more information.   | Species                     | Mouse  |
| AccessionNP_034607.3Calculated Molecular Weight62.3 kDaObserved molecular weight58 kDaTagN-HisBioactivityNot validated for activityPropertiesPurity>95 % as determined by reducing SDS-PAGE.FundtoxinPlease contact us for more information.   | Expression Host             | E.coli   |
| Calculated Molecular Weight62.3 kDaObserved molecular weight58 kDaTagN-HisBioactivityNot validated for activityPropertiesPurity>95 % as determined by reducing SDS-PAGE.EndotoxinPlease contact us for more information.   | Sequence                    | Leu 2-Phe 573  |
| Observed molecular weight58 kDaTagN-HisBioactivityNot validated for activityPropertiesPurity>95 % as determined by reducing SDS-PAGE.EndotoxinPlease contact us for more information.  | Accession                   | NP_034607.3  |
| TagN-HisBioactivityNot validated for activityPropertiesPurity> 95 % as determined by reducing SDS-PAGE.EndotoxinPlease contact us for more information.  | Calculated Molecular Weight | 62.3 kDa   |
| BioactivityNot validated for activityPropertiesPurity> 95 % as determined by reducing SDS-PAGE.EndotoxinPlease contact us for more information.  | Observed molecular weight   | 58 kDa   |
| PropertiesPurity> 95 % as determined by reducing SDS-PAGE.EndotoxinPlease contact us for more information.   | Tag                         | N-His  |
| Purity> 95 % as determined by reducing SDS-PAGE.EndotoxinPlease contact us for more information.   | Bioactivity                 | Not validated for activity   |
| <b>Endotoxin</b> Please contact us for more information.   | Properties                  |  |
|  | Purity                      | > 95 % as determined by reducing SDS-PAGE.   |
| <b>Storage</b> Generally, lyophilized proteins are stable for up to 12 months when stored at -20 to  | Endotoxin                   | Please contact us for more information.  |
| $-80^{\circ}\text{C}$ . Reconstituted protein solution can be stored at 4-8°C for 2-7 days. Aliquots of reconstituted samples are stable at < -20°C for 3 months.  | Storage                     |  |
| <b>Shipping</b> This product is provided as lyophilized powder which is shipped with ice packs.  | Shipping                    | This product is provided as lyophilized powder which is shipped with ice packs.                          |
| FormulationLyophilized from sterile PBS, pH 7.4Normally 5 % - 8 % trehalose, mannitol and 0.01% Tween80 are added as<br>protectants before lyophilization.<br>Please refer to the specific buffer information in the printed manual.   | Formulation                 | Normally 5 % - 8 % trehalose, mannitol and 0.01% Tween80 are added as protectants before lyophilization. |
| <b>Reconstitution</b> Please refer to the printed manual for detailed information.   | Reconstitution              | Please refer to the printed manual for detailed information.   |

Data

| KDa  | MK | R |
|------|----|---|
| 116  | -  |   |
| 66.2 | -  |   |
| 45.0 | -  |   |
| 35.0 | -  |   |
| 25.0 | -  |   |
| 18.4 | -  |   |
| 14.4 | -  |   |
|      |    |   |

> 95 % as determined by reducing SDS-PAGE.

## Background

HSPD1, also known as HSP60, is a member of the chaperonin family. HSPD1 may function as a signaling molecule in the innate immune system. This protein is essential for the folding and assembly of newly imported proteins in the

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mitochondria. It may also prevent misfolding and promote the refolding and proper assembly of unfolded polypeptides generated under stress conditions in the mitochondrial matrix. HSPD1 gene is adjacent to a related family member and the region between the 2 genes functions as a bidirectional promoter. Several pseudogenes have been associated with this gene. Mutations associated with this gene cause autosomal recessive spastic paraplegia 13.Defects in HSPD1 are a cause of spastic paraplegia autosomal dominant type 13 (SPG13). Spastic paraplegia is a degenerative spinal cord disorder characterized by a slow, gradual, progressive weakness and spasticity of the lower limbs. Defects in HSPD1 are the cause of leukodystrophy hypomyelinating type 4 (HLD4); also called mitochondrial HSP60 chaperonopathy or MitCHAP-60 disease. HLD4 is a severe autosomal recessive hypomyelinating leukodystrophy. HSPD1 is cinically characterized by infantile-onset rotary nystagmus, progressive spastic paraplegia, neurologic regression, motor impairment, profound mental retardation. Death usually occurrs within the first two decades of life.