



Desmin (ABT168) Mouse mAb

| Catalog No | BYab-15687 |
|--------------------|--|
| lsotype | lgG |
| Reactivity | Human;Mouse;Rat |
| Applications | IHC;WB; |
| Gene Name | DES |
| Protein Name | CMD1I;CSM1;CSM2;DES;DESM_HUMAN;Desmin;FLJ12025;FLJ39719;FLJ410 13;FLJ41793;Intermediate filament protein;OTTHUMP00000064865 |
| Immunogen | Synthesized peptide derived from human Desmin |
| Specificity | The antibody can specifically recognize human Desmin protein. |
| Formulation | PBS, pH7.2, 0.03% Porcolin 300, containing stabilizing protein |
| Source | Monoclonal Mouse IgG2b, Kappa |
| Purification | The antibody was affinity-purified from mouse ascites by affinity-chromatography using specific immunogen. |
| Dilution | IHC-p 1:200-400, WB 1:200-1000, |
| Concentration | 1 mg/ml |
| Purity | ≥90% |
| Storage Stability | -20°C/1 year |
| Synonyms | CMD1I;CSM1;CSM2;DES;DESM_HUMAN;Desmin;FLJ12025;FLJ39719;FLJ410 13;FLJ41793;Intermediate filament protein;OTTHUMP00000064865 |
| Observed Band | |
| Cell Pathway | Cytoplasmic |
| Tissue Specificity | Appendix/ Colon |
| Function | disease:Defects in DES are the cause of cardiomyopathy dilated type 1I (CMD1I) [MIM:604765]. Dilated cardiomyopathy is a disorder characterized by ventricular dilation and impaired systolic function, resulting in congestive heart failure and arrhythmia. Patients are at risk of premature death.,disease:Defects in DES are the cause of desmin-related cardio-skeletal myopathy (CSM) [MIM:601419]; also known as desmin-related myopathy (DRM). CSM is characterized by skeletal muscle weakness associated with cardiac conduction blocks, arrhythmias, restrictive heart failure, and by intracytoplasmic accumulation of desmin-reactive deposits in cardiac and skeletal muscle cells. A desmin-related myopathy can have a distal onset, it is then known as hereditary distal myopathy (HDM).,disease:Defects in DES are the cause of neurogenic scapuloperoneal |

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| | syndrome Kaeser type (Kaeser syndrome) [MIM:181400]. |
|---------------------------|---|
| Background | This gene encodes a muscle-specific class III intermediate filament. Homopolymers of this protein form a stable intracytoplasmic filamentous network connecting myofibrils to each other and to the plasma membrane. Mutations in this gene are associated with desmin-related myopathy, a familial cardiac and skeletal myopathy (CSM), and with distal myopathies. [provided by RefSeq, Jul 2008], |
| matters needing attention | Avoid repeated freezing and thawing! |
| Usage suggestions | This product can be used in immunological reaction related experiments. For more information, please consult technical personnel. |

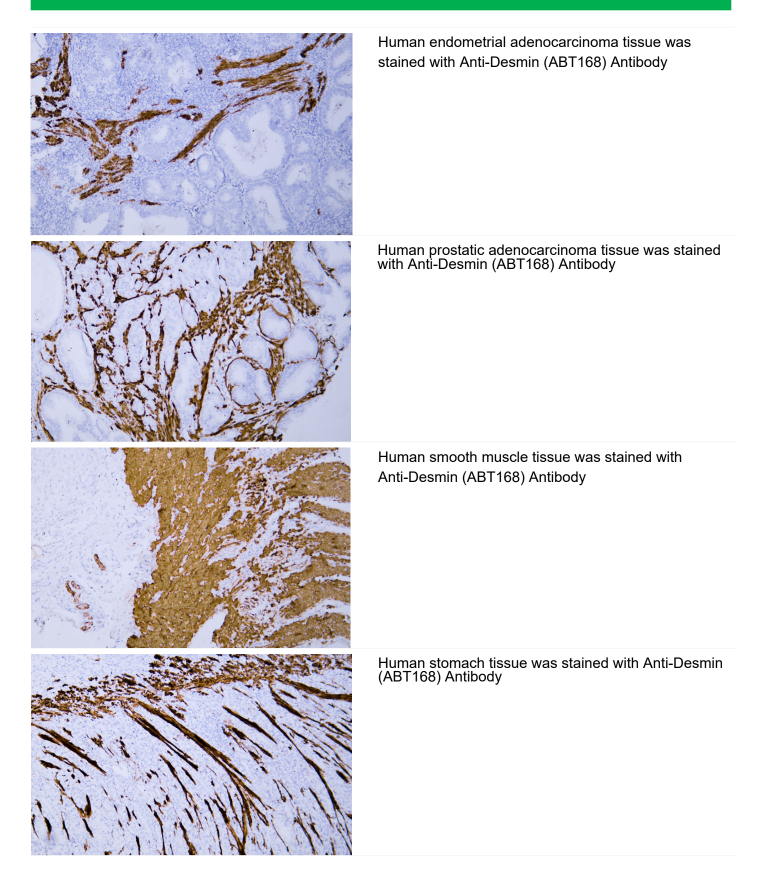
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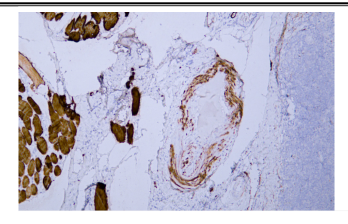
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Human tonsil tissue was stained with Anti-Desmin (ABT168) Antibody



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