

Resource Summary Report

Generated by [RRID](#) on Aug 30, 2026

DDDDK tag antibody [M2] (SureLight® Allophycocyanin)

RRID:AB_1310127

Type: Antibody

Proper Citation

Abcam Cat# ab72569, RRID:AB_1310127

Antibody Information

URL: http://antibodyregistry.org/AB_1310127

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Target Antigen: DDDDK tag antibody [M2] (SureLight® Allophycocyanin)

Host Organism: mouse

Clonality: monoclonal

Comments: validation status unknown, seller recommendations provided in 2012: Flow Cyt, FRET; Flow Cytometry; Immunofluorescence; Other

Antibody Name: DDDDK tag antibody [M2] (SureLight® Allophycocyanin)

Description: This monoclonal targets DDDDK tag antibody [M2] (SureLight® Allophycocyanin)

Antibody ID: AB_1310127

Vendor: Abcam

Catalog Number: ab72569

Record Creation Time: 20250819T231458+0000

Record Last Update: 20250820T044519+0000

Ratings and Alerts

No rating or validation information has been found for DDDDK tag antibody [M2] (SureLight® Allophycocyanin).

No alerts have been found for DDDDK tag antibody [M2] (SureLight® Allophycocyanin).

Data and Source Information

Source: [Antibody Registry](#)

Usage and Citation Metrics

We found 6 mentions in open access literature.

Listed below are recent publications. The full list is available at [RRID](#) .

Edelstein HI, et al. (2025) Conversion of natural cytokine receptors into orthogonal synthetic biosensors. Nature chemical biology.

Howard MK, et al. (2025) Molecular basis of proton sensing by G protein-coupled receptors. Cell, 188(3), 671.

Cohen AA, et al. (2024) Mosaic sarbecovirus nanoparticles elicit cross-reactive responses in pre-vaccinated animals. Cell, 187(20), 5554.

Cohen AA, et al. (2024) Mosaic sarbecovirus vaccination elicits cross-reactive responses in pre-immunized animals. bioRxiv : the preprint server for biology.

Borges JP, et al. (2022) Glycine inhibits NINJ1 membrane clustering to suppress plasma membrane rupture in cell death. eLife, 11.

Edelstein HI, et al. (2020) Elucidation and refinement of synthetic receptor mechanisms. Synthetic biology (Oxford, England), 5(1), ysaa017.