

Resource Summary Report

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U-2932

RRID:CVCL_1896

Type: Cell Line

Proper Citation

Ubigen Cat# YC-C059, RRID:CVCL_1896

Cell Line Information

URL: https://web.expasy.org/cellosaurus/CVCL_1896

Proper Citation: Ubigen Cat# YC-C059, RRID:CVCL_1896

Description: Cell line U-2932 is a Cancer cell line with a species of origin Homo sapiens (Human)

Sex: Female

Disease: Diffuse large B-cell lymphoma activated B-cell type

Defining Citation: [PMID:12533045](#), [PMID:20054396](#), [PMID:23257783](#), [PMID:23292937](#), [PMID:23295736](#), [PMID:25485619](#), [PMID:25894527](#), [PMID:26589293](#), [PMID:27566572](#), [PMID:28196595](#), [PMID:29416618](#), [PMID:29533902](#), [PMID:29666304](#), [PMID:30165192](#), [PMID:31160637](#)

Comments: Donor information: Originally the patient was suffering from Hodgkin lymphoma., Characteristics: Genetically heterogeneous, consists of 2 subclones (PubMed=27566572)., Part of: MD Anderson Cell Lines Project., Part of: LL-100 blood cancer cell line panel.

Category: Cancer cell line

Organism: Homo sapiens (Human)

Name: U-2932

Synonyms: U2932

Cross References: EFO:EFO_0006499, ArrayExpress:E-MTAB-2706, ArrayExpress:E-MTAB-4527, ArrayExpress:E-MTAB-7721, ArrayExpress:E-MTAB-7722, BioSample:SAMN03473452, cancercellines:CVCL_1896, ChEMBL-Cells:ChEMBL4295421, ChEMBL-Targets:ChEMBL4296507, Cosmic:1487546, Cosmic:1517654, Cosmic:1945196, DSMZ:ACC-633, DSMZCellDive:ACC-633,

EGA:EGAS00001000610, GEO:GSM1035339, GEO:GSM1059802, GEO:GSM1374971, PharmacDB:U2932_1621_2019, PRIDE:PXD000589, Progenetix:CVCL_1896, PubChem_Cell_line:CVCL_1896, Ubigen:YC-C059, Wikidata:Q54973560

ID: CVCL_1896

Vendor: Ubigen

Catalog Number: YC-C059

Record Creation Time: 20250130T072907+0000

Record Last Update: 20260913T011657+0000

Ratings and Alerts

No rating or validation information has been found for U-2932.

No alerts have been found for U-2932.

Data and Source Information

Source: [Cellosaurus](#)

Usage and Citation Metrics

We found 32 mentions in open access literature.

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

Demel UM, et al. (2026) Aberrant SUMOylation Restricts the Targetable Cancer Immunopeptidome. Advanced science (Weinheim, Baden-Wurttemberg, Germany), 13(17), e11449.

Zhao J, et al. (2026) Construction and experimental verification of a prognostic model based on tumor-infiltrating lymphocytes related genes (TILRGs) in diffuse large B-cell lymphoma. SLAS technology, 39, 100425.

Enssle JC, et al. (2026) Pathogenesis of diffuse large B cell lymphoma proteogenotypes. Cancer cell, 44(7), 1437.

Chen Z, et al. (2025) YTHDF2 promotes ATP synthesis and immune evasion in B cell malignancies. Cell, 188(2), 331.

Duranti C, et al. (2025) Antineoplastic Activity of a Novel Trispecific Single-Chain Antibody Targeting the hERG1/??1 Integrin Complex and TRAIL Receptors. Molecular cancer therapeutics, 24(10), 1584.

Lüönd F, et al. (2025) DLBCL Cells Emerge after CD19 CAR T Cells with Cross-Antigen Resistance and a Gene Signature Predictive of Clinical CAR T-cell Response. Blood

cancer discovery, 6(6), 580.

Shi X, et al. (2025) NFS1, together with FXN, protects cells from ferroptosis and DNA damage in diffuse large B-cell lymphoma. *Redox biology*, 87, 103878.

McCrury M, et al. (2024) Bifunctional Inhibitor Reveals NEK2 as a Therapeutic Target and Regulator of Oncogenic Pathways in Lymphoma. *Molecular cancer therapeutics*, 23(3), 316.

Choi J, et al. (2024) Molecular targets of glucocorticoids that elucidate their therapeutic efficacy in aggressive lymphomas. *Cancer cell*, 42(5), 833.

Mehdi SH, et al. (2024) Development of New Diffuse Large B Cell Lymphoma Mouse Models. *Cancers*, 16(17).

Pieper NM, et al. (2024) Inhibition of bromodomain and extra-terminal proteins targets constitutively active NF κ B and STAT signaling in lymphoma and influences the expression of the antiapoptotic proteins BCL2A1 and c-MYC. *Cell communication and signaling : CCS*, 22(1), 415.

Eken JA, et al. (2024) Antigen-independent, autonomous B cell receptor signaling drives activated B cell DLBCL. *The Journal of experimental medicine*, 221(5).

Scuoppo C, et al. (2024) Repurposing NAMPT Inhibitors for Germinal Center B Cell-Like Diffuse Large B-Cell Lymphoma. *Blood cancer discovery*, 5(6), 417.

Zeng H, et al. (2023) Exosomal PD α L1 promotes the formation of an immunosuppressive microenvironment in gastric diffuse large B α cell lymphoma. *Oncology reports*, 49(5).

Scheich S, et al. (2023) Targeting N-linked Glycosylation for the Therapy of Aggressive Lymphomas. *Cancer discovery*, 13(8), 1862.

Venturutti L, et al. (2023) An Aged/Autoimmune B-cell Program Defines the Early Transformation of Extranodal Lymphomas. *Cancer discovery*, 13(1), 216.

Rodina A, et al. (2023) Systems-level analyses of protein-protein interaction network dysfunctions via epichaperomics identify cancer-specific mechanisms of stress adaptation. *Nature communications*, 14(1), 3742.

Johnson Z, et al. (2023) IOA-244 is a Non-ATP-competitive, Highly Selective, Tolerable PI3K Delta Inhibitor That Targets Solid Tumors and Breaks Immune Tolerance. *Cancer research communications*, 3(4), 576.

Pecori R, et al. (2022) ADAR RNA editing on antisense RNAs results in apparent U-to-C base changes on overlapping sense transcripts. *Frontiers in cell and developmental biology*, 10, 1080626.

Biggi AFB, et al. (2022) The Epstein-Barr Virus Hacks Immune Checkpoints: Evidence and Consequences for Lymphoproliferative Disorders and Cancers. *Biomolecules*, 12(3).