

Resource Summary Report

Generated by [NIF](#) on Sep 22, 2026

HEL 92.1.7

RRID:CVCL_2481

Type: Cell Line

Proper Citation

RRID:CVCL_2481

Cell Line Information

URL: https://web.expasy.org/cellosaurus/CVCL_2481

Proper Citation: RRID:CVCL_2481

Description: Cell line HEL 92.1.7 is a Cancer cell line with a species of origin Homo sapiens (Human)

Sex: Male

Disease: Erythroleukemia

Defining Citation: [PMID:20215515](#), [PMID:22460905](#), [PMID:30285677](#), [PMID:30894373](#), [PMID:31068700](#), [PMID:31978347](#)

Comments: Donor information: Originally the patient was suffering from Hodgkin lymphoma., Population: Caucasian., Part of: Tumor Immunology Bank (TIB) collection from Salk (transferred to ATCC in 1981)., Part of: Cancer Dependency Map project (DepMap) (includes Cancer Cell Line Encyclopedia - CCLE).

Category: Cancer cell line

Organism: Homo sapiens (Human)

Name: HEL 92.1.7

Synonyms: HEL92.1.7, HEL-92.1.7, HEL-92-1-7, HEL-92_1_7, HEL-92, HEL9217

Cross References: BTO:BTO_0003320, CLO:CLO_0003682, CLO:CLO_0003683, EFO:EFO_0002193, CLDB:cl5067, CLDB:cl7132, ArrayExpress:E-MTAB-38, ArrayExpress:E-MTAB-2770, ATCC:TIB-180, BCRC:60370, BCRJ:0308, BioSample:SAMN10987842, cancercellines:CVCL_2481, Cell_Model_Passport:SIDM00593, ChEMBL-Cells:ChEMBL4295457, ChEMBL-Targets:ChEMBL4296425, CLS:300462, Cosmic:683543, Cosmic:976723, DepMap:ACH-000005, ECACC:92111706, GEO:GSM887076, GEO:GSM888146, GEO:GSM1374526,

GEO:GSM1374527, IARC_TP53:21369, ICLC:HTL03002, IGRhCellID:HEL9217, LiGeA:CCLE_672, LINCS_LDP:LCL-1076, Lonza:1042, PharmacDB:HEL92_1_7_535_2019, Progenetix:CVCL_2481, PubChem_Cell_line:CVCL_2481, Wikidata:Q54882504

ID: CVCL_2481

Hierarchy: cvcl_0001

Record Creation Time: 20220427T220330+0000

Record Last Update: 20260920T003541+0000

Ratings and Alerts

No rating or validation information has been found for HEL 92.1.7.

No alerts have been found for HEL 92.1.7.

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](#) .

Peng M, et al. (2026) MMRN1-EGFR drives sialylglycan-Siglec immune evasion in AML leukemia stem cells. *Cell stem cell*, 33(5), 800.

Roy P, et al. (2026) A stress-induced PI3P complex controls autophagy in erythroid precursors. *iScience*, 29(4), 115182.

Tripathi C, et al. (2026) Nanobody-Engineered CLL-1 CAR T Cells: Optimizing Tumor-Specific Cytotoxicity and Minimizing Off-Tumor Toxicity. *Cancer research communications*, 6(3), 528.

Gao J, et al. (2026) Dual inhibition of tubulin and DNMT by the novel letermovir derivative H62 blocks leukemia. *Biochemical pharmacology*, 252, 118196.

Uible EE, et al. (2026) Scaffolding-dependent CASP1 constrains excessive cell-intrinsic inflammatory signaling in leukemia. *Cell chemical biology*, 33(1), 59.

Le BV, et al. (2026) Pol ϵ activity modulates sensitivity to standard therapies in DNMT3A-deficient leukemia. *Cell reports. Medicine*, 7(3), 102687.

Chancellor A, et al. (2025) The carbonyl nucleobase adduct M3Ade is a potent antigen for adaptive polyclonal MR1-restricted T cells. *Immunity*, 58(2), 431.

- Kumar S, et al. (2025) Development of PROTACs for targeted degradation of oncogenic TRK fusions. *bioRxiv : the preprint server for biology*.
- Fiskus W, et al. (2025) Preclinical efficacy of tasquinimod-based combinations in advanced myeloproliferative neoplasms in blastic phase. *Blood advances*, 9(21), 5598.
- Fiskus W, et al. (2025) Preclinical efficacy of CDK7 inhibitor-based combinations against myeloproliferative neoplasms transformed to AML. *Blood*, 145(6), 612.
- Liu T, et al. (2025) A generalized strategy to kill leukemic cells by targeting the regulatory systems governing mitochondrial membrane potential. *Cell reports*, 44(11), 116496.
- Graham K, et al. (2024) Discovery of YAP1/TAZ pathway inhibitors through phenotypic screening with potent anti-tumor activity via blockade of Rho-GTPase signaling. *Cell chemical biology*, 31(7), 1247.
- Saleiro D, et al. (2023) Targeting CHAF1B Enhances IFN Activity against Myeloproliferative Neoplasm Cells. *Cancer research communications*, 3(5), 943.
- Blöchl C, et al. (2023) Transcriptionally imprinted glycomic signatures of acute myeloid leukemia. *Cell & bioscience*, 13(1), 31.
- Nicosia L, et al. (2023) Therapeutic targeting of EP300/CBP by bromodomain inhibition in hematologic malignancies. *Cancer cell*, 41(12), 2136.
- Wei TT, et al. (2022) Cannabinoid receptor 1 antagonist genistein attenuates marijuana-induced vascular inflammation. *Cell*, 185(10), 1676.
- Roversi FM, et al. (2022) Novel inhibitor of hematopoietic cell kinase as a potential therapeutic agent for acute myeloid leukemia. *Cancer immunology, immunotherapy : CII*, 71(8), 1909.
- Yang BH, et al. (2022) Epigenetics-Associated Risk Reduction of Hematologic Neoplasms in a Nationwide Cohort Study: The Chemopreventive and Therapeutic Efficacy of Hydralazine. *Frontiers in oncology*, 12, 809014.
- Polyanskaya SA, et al. (2022) SCP4-STK35/PDIK1L complex is a dual phospho-catalytic signaling dependency in acute myeloid leukemia. *Cell reports*, 38(2), 110233.
- Cao Z, et al. (2021) ZMYND8-regulated IRF8 transcription axis is an acute myeloid leukemia dependency. *Molecular cell*, 81(17), 3604.