

Resource Summary Report

Generated by [RRID](#) on Sep 2, 2026

VU-SCC-1131

RRID:CVCL_XX18

Type: Cell Line

Proper Citation

RRID:CVCL_XX18

Cell Line Information

URL: https://web.expasy.org/cellosaurus/CVCL_XX18

Proper Citation: RRID:CVCL_XX18

Description: Cell line VU-SCC-1131 is a Cancer cell line with a species of origin Homo sapiens (Human)

Sex: Female

Disease: Fanconi anemia, complementation group C

Defining Citation: [PMID:15735012](#), [PMID:23613873](#), [PMID:25609062](#), [PMID:26122845](#), [PMID:31541927](#), [PMID:36912284](#)

Category: Cancer cell line

Organism: Homo sapiens (Human)

Name: VU-SCC-1131

Synonyms: VU1131, VU-1131, VU-1131-T2.8

Cross References: cancercellines:CVCL_XX18, IARC_TP53:21670, Wikidata:Q98134585

ID: CVCL_XX18

Record Creation Time: 20220427T221710+0000

Record Last Update: 20260830T000257+0000

Ratings and Alerts

No rating or validation information has been found for VU-SCC-1131.

No alerts have been found for VU-SCC-1131.

Data and Source Information

Source: [Cellosaurus](#)

Usage and Citation Metrics

We found 4 mentions in open access literature.

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

Kohabir KAV, et al. (2025) Protocol for high-precision CRISPR-Cas12a-based SNV detection on synthetic DNA, cell line cfDNA models, and liquid biopsies. STAR protocols, 6(2), 103696.

Kohabir KAV, et al. (2024) Synthetic mismatches enable specific CRISPR-Cas12a-based detection of genome-wide SNVs tracked by ARTEMIS. Cell reports methods, 4(12), 100912.

Nguyen HT, et al. (2023) Fanconi anemia-isogenic head and neck cancer cell line pairs: A basic and translational science resource. International journal of cancer, 153(1), 183.

Silva de Araujo BE, et al. (2022) Detection of cytogenetic changes and chromosomal aneuploidy with fluorescent in situ hybridization in cytological specimens of oral cancers in Fanconi anemia-Proof of concept. Clinical and experimental dental research, 8(1), 108.