

Resource Summary Report

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GM03767

RRID:CVCL_X111

Type: Cell Line

Proper Citation

Coriell Cat# GM03767, RRID:CVCL_X111

Cell Line Information

URL: https://web.expasy.org/cellosaurus/CVCL_X111

Proper Citation: Coriell Cat# GM03767, RRID:CVCL_X111

Description: Cell line GM03767 is a Finite cell line with a species of origin Homo sapiens (Human)

Sex: Male

Disease: Deletion 18p syndrome

Defining Citation: [PMID:6617268](#), [PMID:6661932](#)

Comments: Karyotypic information: 46,XY,del(18)(qter->p11) (Coriell=GM03767)., Population: Caucasian.

Category: Finite cell line

Organism: Homo sapiens (Human)

Name: GM03767

Synonyms: GM 3767

Cross References: CLO:CLO_0015491, BioSample:SAMN00808521, Coriell:GM03767, Wikidata:Q54838222

ID: CVCL_X111

Vendor: Coriell

Catalog Number: GM03767

Record Creation Time: 20220427T215931+0000

Record Last Update: 20260829T232820+0000

Ratings and Alerts

No rating or validation information has been found for GM03767.

No alerts have been found for GM03767.

Data and Source Information

Source: [CelloSaurus](#)

Usage and Citation Metrics

We have not found any literature mentions for this resource.