

Feline leukemia virus, complete genome

NCBI Reference Sequence: NC_001940.1

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LOCUS NC_001940 8448 bp ss-RNA linear VRL 13-AUG-2018

DEFINITION Feline leukemia virus, complete genome.

ACCESSION NC_001940

VERSION NC_001940.1

DBLINK BioProject: [PRJNA485481](#)

KEYWORDS RefSeq.

SOURCE Feline leukemia virus

ORGANISM [Feline leukemia virus](#)
Viruses; Riboviria; Pararnavirae; Artverviricota; Revtraviricetes;
Ortervirales; Retroviridae; Orthoretrovirinae; Gammaretrovirus.

REFERENCE 1 (bases 1 to 8448)

AUTHORS Chen,H., Bechtel,M.K., Shi,Y., Phipps,A., Mathes,L.E., Hayes,K.A.
and Roy-Burman,P.

TITLE Pathogenicity induced by feline leukemia virus, Rickard strain,
subgroup A plasmid DNA (pFRA)

JOURNAL J. Virol. 72 (9), 7048-7056 (1998)

PUBMED [9696797](#)

REFERENCE 2 (bases 1 to 8448)

CONSRTM NCBI Genome Project

TITLE Direct Submission

JOURNAL Submitted (01-AUG-2000) National Center for Biotechnology
Information, NIH, Bethesda, MD 20894, USA

REFERENCE 3 (bases 1 to 8448)

AUTHORS Chen,H. and Roy-Burman,P.

TITLE Direct Submission

JOURNAL Submitted (08-MAR-1998) Department of Pathology, Univ. of
Southern
California, School of Medicine, 2011 Zonal Ave., HMR 212, Los
Angeles, CA 90033, USA

COMMENT PROVISIONAL [REFSEQ](#): This record has not yet been subject to final
NCBI review. The reference sequence was derived from [AF052723](#).
COMPLETENESS: full length.

FEATURES Location/Qualifiers

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