

Human papillomavirus type 16 isolate Amazonian, complete genome
GenBank: HM057182.1

FASTA Graphics

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LOCUS HM057182 7915 bp DNA circular VRL 25-JUL-2016

DEFINITION Human papillomavirus type 16 isolate Amazonian, complete genome.

ACCESSION HM057182

VERSION HM057182.1

KEYWORDS .

SOURCE Human papillomavirus type 16

ORGANISM Human papillomavirus type 16
Viruses; Monodnaviria; Shotokuvirae; Cossaviricota;
Papovaviricetes; Zurhausenvirales; Papillomaviridae;
Firstpapillomavirinae; Alphapapillomavirus.

REFERENCE 1 (bases 1 to 7915)

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TITLE Direct Submission

JOURNAL Submitted (03-APR-2010) Biotechnology, Molecular Diagnostic's Lab,
Amazonas' Federal University, Av. Gen. Rodrigo Otavio Jordao Ramos,
Manaus, Amazonas, Brazil

FEATURES Location/Qualifiers

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