CLUSTAL 2.1 multiple sequence Alignment

MF288709.1	ATGGCGCGCTTTGAGGATCCAACACGGCGACCCTACAAGCTACCAGATCT
MF288709.1 MF288708.1	
	ATGGCGCGCTTTGAGGATCCAACACGGCGACCCTACAAGCTACCAGATCT
MF288713.1	ATGGCGCGCTTTGAGGATCCAACACGGCGACCCTACAAGCTACCTGATCT
MF288710.1	ATGGCGCGCTTTGAGGATCCAACACGGCGACCCTACAAGCTACCTGATCT
MF288712.1	ATGGCGCGCTTTGAGGATCCAACACGGCGACCCTACAAGCTACCTGATTT
MF288711.1	ATGGCGCGCTTTGAGGATCCAACACGGCGACCCTACAAGCTACCTGATTT
MF288714.1	ATGGCGCGCTTTGAGGATCCAACACGGCGACCCTACAAGCTACCTGATCT
MF288716.1	ATGGCGCGCTTTGAGGATCCAACACGGCGACCCTACAAGCTACCTGATCT
MF288715.1	ATGGCGCGCTTTGAGGATCCAACACGGCGACCCTACAAGCTACCTGATCT
MF288717.1	ATGGCGCGCTTTGAGGATCCAACACGGCGACCCTACAAGCTACCTGATCT
MF288727.1	ATGGCGCGCTTTGAGGATCCAACACGGCGACCCTACAAGCTACCTGATCT
MF288726.1	ATGGCGCGCTTTGAGGATCCAACACGGCGACCCTACAAGCTACCTGATCT
MF288724.1	ATGGCGCGCTTTGAGGATCCAACACGGCGACCCTACAAGCTACCTGATCT
MF288721.1	ATGGCGCGCTTTGAGGATCCAACACGGCGACCCTACAAGCTACCTGATCT
MF288720.1	ATGGCGCGCTTTGAGGATCCAACACGGCGACCCTACAAGCTACCTGATCT
MF288719.1	ATGGCGCGCTTTGAGGATCCAACACGGCGACCCTACAAGCTACCTGATCT
MF288718.1	ATGGCGCGCTTTGAGGATCCAACACGGCGACCCTACAAGCTACCTGATCT
MF288725.1	ATGGCGCGCTTTGAGGATCCAACACGGCGACCCTACAAGCTACCTGATCT
MF288723.1	ATGGCGCGCTTTGAGGATCCAACACGGCGACCCTACAAGCTACCTGATCT
MF288722.1	ATGGCGCGCTTTGAGGATCCAACACGGCGACCCTACAAGCTACCTGATCT

MF288709.1	GTGCACGGAACTGAACACTTCACTGCAAGACATAGAAATAACCTGTGTAT
MF288708.1	GTGCACGGAACTGAACACTTCACTGCAAGACATAGAAATAACCTGTGTAT
MF288713.1	GTGCACGGAACTGAACACTTCACTGCAAGACATAGAAATAACCTGTGTAT
MF288710.1	GTGCACGGAACTGAACACTTCACTGCAAGACATAGAAATAACCTGTGTAT
MF288712.1	GTGCACGGAACTGAACACTTCACTGCAAGACATAGAAATAACCTGTGTAT
MF288711.1	GTGCACGGAACTGAACACTTCACTGCAAGACATAGAAATAACCTGTGTAT
MF288714.1	GTGCACGGAACTGAACACTTCACTGCAAGACATAGAAATAACCTGTGTAT
MF288716.1	GTGCACGGAACTGAACACTTCACTGCAAGACATAGAAATAACCTGTGTAT
MF288715.1	GTGCACGGAACTGAACACTTCACTGCAAGACATAGAAATAACCTGTGTAT
MF288717.1	GTGCACGGAACTGAACACTTCACTGCAAGACATAGAAATAACCTGTGTAT
MF288727.1	GTGCACGGAACTGAACACTTCACTGCAAGACATAGAAATAACCTGTGTAT
MF288726.1	GTGCACGGAACTGAACACTTCACTGCAAGACATAGAAATAACCTGTGTAT
MF288724.1	GTGCACGGAACTGAACACTTCACTGCAAGACATAGAAATAACCTGTGTAT
MF288721.1	GTGCACGGAACTGAACACTTCACTGCAAGACATAGAAATAACCTGTGTAT
MF288721.1 MF288720.1	GTGCACGGAACTGAACACTTCACTGCAAGACATAGAAATAACCTGTGTAT GTGCACGGAACTGAACACTTCACTGCAAGACATAGAAATAACCTGTGTAT
MF288720.1	GTGCACGGAACTGAACACTTCACTGCAAGACATAGAAATAACCTGTGTAT
MF288720.1 MF288719.1	GTGCACGGAACTGAACACTTCACTGCAAGACATAGAAATAACCTGTGTAT GTGCACGGAACTGAACACTTCACTGCAAGACATAGAAATAACCTGTGTAT
MF288720.1 MF288719.1 MF288718.1	GTGCACGGAACTGAACACTTCACTGCAAGACATAGAAATAACCTGTGTAT GTGCACGGAACTGAACACTTCACTGCAAGACATAGAAATAACCTGTGTAT GTGCACGGAACTGAACACTTCACTGCAAGACATAGAAATAACCTGTGTAT
MF288720.1 MF288719.1 MF288718.1 MF288725.1	GTGCACGGAACTGAACACTTCACTGCAAGACATAGAAATAACCTGTGTAT GTGCACGGAACTGAACACTTCACTGCAAGACATAGAAATAACCTGTGTAT GTGCACGGAACTGAACACTTCACTGCAAGACATAGAAATAACCTGTGTAT GTGCACGGAACTGAACACTTCACTGCAAGACATAGAAATAACCTGTGTAT
MF288720.1 MF288719.1 MF288718.1 MF288725.1 MF288723.1	GTGCACGGAACTGAACACTTCACTGCAAGACATAGAAATAACCTGTGTAT GTGCACGGAACTGAACACTTCACTGCAAGACATAGAAATAACCTGTGTAT GTGCACGGAACTGAACACTTCACTGCAAGACATAGAAATAACCTGTGTAT GTGCACGGAACTGAACACTTCACTGCAAGACATAGAAATAACCTGTGTAT GTGCACGGAACTGAACACTTCACTGCAAGACATAGAAATAACCTGTGTAT

CLUSTAL	2.1	multiple	sequence	Alignment
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CLUSTAL 2.1 multiple sequence Al	ignment
MF288708.1	ATTGCAAGACAGTATTGGAACTTACAGAGGTATTTGAATTTGCATTTAAA
MF288713.1	ATTGCAAGACAGTATTGGAACTTACAGAGGTATTTGAATTTGCATTTAAA
MF288710.1	ATTGCAAGACAGTATTGGAACTTACAGAGGTATTTGAATTTGCATTTAAA
MF288712.1	ATTGCAAGACAGTATTGGAACTTACAGAGGTATTTGAATTTGCATTTAAA
MF288711.1	ATTGCAAGACAGTATTGGAACTTACAGAGGTATTTGAATTTGCATTTAAA
MF288714.1	ATTGCAAGACAGTATTGGAACTTACAGAGGTATTTGAATTTGCATTTAAA
MF288716.1	ATTGCAAGACAGTATTGGAACTTACAGAGGTATTTGAATTTGCATTTAAA
MF288715.1	ATTGCAAGACAGTATTGGAACTTACAGAGGTATTTGAATTTGCATTTAAA
MF288717.1	ATTGCAAGACAGTATTGGAACTTACAGAGGTATTTGAATTTGCATTTAAA
MF288727.1	ATTGCAAGACAGTATTGGAACTTACAGAGGTATTTGAATTTGCATTTAAA
MF288726.1	ATTGCAAGACAGTATTGGAACTTACAGAGGTATTTGAATTTGCATTTAAA
MF288724.1	ATTGCAAGACAGTATTGGAACTTACAGAGGTATTTGAATTTGCATTTAAA
MF288721.1	ATTGCAAGACAGTTTTGGAACTTACAGAGGTATTTGAATTTGCATTTAAA
MF288720.1	ATTGCAAGACAGTTTTGGAACTTACAGAGGTATTTGAATTTGCATTTAAA
MF288719.1	ATTGCAAGACAGTTTTGGAACTTACAGAGGTATTTGAATTTGCATTTAAA
MF288718.1	ATTGCAAGACAGTTTTGGAACTTACAGAGGTATTTGAATTTGCATTTAAA
MF288725.1	ATTGCAAGACAGTATTGGAACTTACAGAGGTATTTGAATTTGCATTTAAA
MF288723.1	ATTGCAAGACAGTATTGGAACTTACAGAGGTATTTGAATTTGCATTTAAA
MF288722.1	ATTGCAAGACAGTATTGGAACTTACAGAGGTATTTGAATTTGCATTTAAA
	******** *******************
MF288709.1	GATTTATTTGTGGTGTATAGAGACAGTATACCGCATGCTGCATGCCATAA
MF288708.1	GATTTATTTGTGGTGTATAGAGACAGTATACCGCATGCTGCATGCCATAA
MF288713.1	GATTTATTTGTGGTGTATAGAGACAGTATACCGCATGCTGCATGCCATAA
MF288710.1	GATTTATTTGTGGTGTATAGAGACAGTATACCGCATGCTGCATGCCATAA
MF288712.1	GATTTATTTGTGGTGTATAGAGACAGTATACCGCATGCTGCATGCCATAA
MF288711.1	GATTTATTTGTGGTGTATAGAGACAGTATACCGCATGCTGCATGCCATAA
MF288714.1	GATTTATTTGTGGTGTATAGAGACAGTATACCGCATGCTGCATGCCATAA
MF288716.1	GATTTATTTGTGGTGTATAGAGACAGTATACCGCATGCTGCATGCCATAA
MF288715.1	GATTTATTTGTGGTGTATAGAGACAGTATACCGCATGCTGCATGCCATAA
MF288717.1	GATTTATTTGTGGTGTATAGAGACAGTATACCGCATGCTGCATGCCATAA
MF288727.1	GATTTATTTGTGGTGTATAGAGACAGTATACCGCATGCTGCATGCCATAA
MF288726.1	GATTTATTGTGGTGTATAGAGACAGTATACCGCATGCTGCATGCCATAA
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MF288724.1	GATTTATTTGTGGTGTATAGAGACAGTATACCGCATGCTGCATGCCATAA
MF288721.1	GATTTATTTGTGGTGTATAGAGACAGTATACCGCATGCTGCATGCCATAA
MF288720.1	GATTTATTTGTGGTGTATAGAGACAGTATACCGCATGCTGCATGCCATAA
MF288719.1	GATTTATTTGTGGTGTATAGAGACAGTATACCGCATGCTGCATGCCATAA
MF288718.1	GATTTATTTGTGGTGTATAGAGACAGTATACCGCATGCTGCATGCCATAA
MF288725.1	GATTTATTTGTGGTGTATAGAGACAGTATACCGCATGCTGCATGCCATAA
MF288723.1	
	GATTTATTTGTGGTGTATAGAGACAGTATACCGCATGCTGCATGCCATAA
MF288722.1	GATTTATTTGTGGTGTATAGAGACAGTATACCGCATGCTGCATGCCATAA

MF288709.1	ATGTATAGATTTTTATTCTAGAATTAGAGAATTAAGACATTATTCAGACT
MF288708.1	ATGTATAGATTTTTATTCTAGAATTAGAGAATTAAGACATTATTCAGACT
MF288713.1	ATGTATAGATTTTTATTCTAGAATTAGAGAATTAAGACATTATTCAGACT
MF288710.1	ATGTATAGATTTTATTCTAGAATTAGAGAATTAAGACATTATTCAGACT

CLUSTAL	2.1	multiple	sequence	Alignment
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CLUSTAL 2.1 multiple seq	quence Alignment
MF288712.1	ATGTATAGATTTTTATTCTAGAATTAGAGAATTAAGACATTATTCAGACT
MF288711.1	ATGTATAGATTTTTATTCTAGAATTAGAGAATTAAGACATTATTCAGACT
MF288714.1	ATGTATAGATTTTTATTCTAGAATTAGAGAATTAAGACATTATTCAGACT
MF288716.1	ATGTATAGATTTTTATTCTAGAATTAGAGAATTAAGACATTATTCGGACT
MF288715.1	ATGTATAGATTTTTATTCTAGAATTAGAGAATTAAGACATTATTCGGACT
MF288717.1	ATGTATAGATTTTTATTCTAGAATTAGAGAATTAAGACATTATTCAGACT
MF288727.1	ATGTATAGATTTTTATTCTAGAATTAGAGAATTAAGACATTATTCAGACT
MF288726.1	ATGTATAGATTTTTATTCTAGAATTAGAGAATTAAGACATTATTCAGACT
MF288724.1	ATGTATAGATTTTTATTCTAGAATTAGAGAATTAAGACATTATTCAGACT
MF288721.1	ATGTATAGATTTTTATTCTAGAATTAGAGAATTAAGACATTATTCAGACT
MF288720.1	ATGTATAGATTTTTATTCTAGAATTAGAGAATTAAGACATTATTCAGACT
MF288719.1	ATGTATAGATTTTTATTCTAGAATTAGAGAATTAAGACATTATTCAGACT
MF288718.1	ATGTATAGATTTTTATTCTAGAATTAGAGAATTAAGACATTATTCAGACT
MF288725.1	ATGTATAGATTTTTATTCTAGAATTAGAGAATTAAGACATTATTCAGACT
MF288723.1	ATGTATAGATTTTTATTCTAGAATTAGAGAATTAAGACATTATTCAGACT
MF288722.1	ATGTATAGATTTTTATTCTAGAATTAGAGAATTAAGACATTATTCAGACT

MF288709.1	CTGTGTATGGAGACACATTGGAAAAACTAACTAACACTGGGTTATACAAT
MF288708.1	CTGTGTATGGAGACACTTGGAAAAACTAACTAACACTGGGTTATACAAT
MF288713.1	CTGTGTATGGAGACACTTGGAAAAACTAACTAACACTGGGTTATACAAT
MF288710.1	CTGTGTATGGAGACACATTGGAAAAACTAACTAACACTGGGTTATACAAT
MF288712.1	CTGTGTATGGAGACACATTGGAAAAACTAACTAACACTGGTTATACAAT
MF288711.1	CTGTGTATGGAGACACATTGGAAAAACTAACTAACACTGGGTTATACAAT
MF288714.1	CTGTGTATGGAGACACATTGGAAAAACTAACTAACACTGGGTTATACAAT
MF288716.1	
MF288715.1	CTGTGTATGGAGACACATTGGAAAAACTAACTAACACTGGGTTATACAAT
MF288717.1	CTGTGTATGGAGACACATTGGAAAAACTAACTAACACTGGGTTATACAAT
111 200 / 1 / 1 1	CTGTGTATGGAGACACATTGGAAAAACTAACTAACACTGGGTTATACAAT
MF288727.1	CTGTGTATGGAGACACATTGGAGAAACTAACTAACACTGGGTTATACAAT
MF288726.1	CTGTGTATGGAGACACATTGGAAAAACTAACTAACACTGGGTTATACAAT
MF288724.1	CTGTGTATGGAGACACATTGGAAAAACTAACTAACACTGGGTTATACAAT
MF288721.1	CTGTGTATGGAGACACATTGGAAAAACTAACTAACACTGGGTTATACAAT
MF288720.1	CTGTGTATGGAGACACTTGGAAAAACTAACTAACACTGGGTTATACAAT
MF288719.1	CTGTGTATGGAGACACTTGGAAAAACTAACTAACACTGGGTTATACAAT
MF288718.1	CTGTGTATGGAGACACTTGGAAAAACTAACTAACACTGGGTTATACAAT
MF288725.1	CTGTGTATGGAGACACATTGGAAAAACTAACTAACACTGGGTTATACAAT
MF288723.1	CTGTGTATGGAGACACATTGGAAAAACTAACTAACACTGGGTTATACAAT
MF288722.1	CTGTGTATGGAGACACATTGGAAAAACTAACTAACACTGGGTTATACAAT

MF288709.1	TTATTAATAAGGTGCCTGCGGTGCCAGAAACCGTTGAATCCAGCAGAAAA
MF288708.1	TTATTAATAAGGTGCCTGCGGTGCCAGAAACCGTTGAATCCAGCAGAAAA
MF288713.1	TTATTAATAAGGTGCCTGCGGTGCCAGAAACCGTTGAATCCAGCAGAAAA
MF288710.1	TTATTAATAAGGTGCCTGCGGTGCCAGAAACCGTTGAATCCAGCAGAAAA
MF288712.1	TTATTAATAAGGTGCCTGCGGTGCCAGAAACCGTTGAATCCAGCAGAAAA
MF288711.1	TTATTAATAAGGTGCCTGCGGTGCCAGAAACCGTTGAATCCAGCAGAAAA
MF288714.1	TTATTAATAAGGTGCCTGCGGTGCCAGAAACCGTTGAATCCAGCAGAAAA
111 200 / 11.1	IIII IIIIIII OO OO OO OO OO OO OO OO OO

CLUSTAL.	2 1	multiple	seguence	Alignment
CHOSTAL	Z.I	MATCINIC	SEUUELICE	ATTAIMEN

CLUSIAL Z.1 MUICIPIE SEQUENCE	Allgiment
MF288716.1	TTATTAATAAGGTGCCTGCGGTGCCAGAAACCGTTGAATCCAGCAGAAAA
MF288715.1	TTATTAATAAGGTGCCTGCGGTGCCAGAAACCGTTGAATCCAGCAGAAAA
MF288717.1	TTATTAATAAGGTGCCTGCGGTGCCAGAAACCGTTGAATCCAGCAGAAAA
MF288727.1	TTATTAATAAGGTGCCTGCGGTGCCAGAAACCGTTGAATCCAGCAGAAAA
MF288726.1	TTATTAATAAGGTGCCTGCGGTGCCAGAAACCGTTGAATCCAGCAGAAAA
MF288724.1	TTATTAATAAGGTGCCTGCGGTGCCAGAAACCGTTGAATCCAGCAGAAAA
MF288721.1	TTATTAATAAGGTGCCTGCGGTGCCAGAAACCGTTGAATCCAGCAGAAAA
MF288720.1	TTATTAATAAGGTGCCTGCGGTGCCAGAAACCGTTGAATCCAGCAGAAAA
MF288719.1	TTATTAATAAGGTGCCTGCGGTGCCAGAAACCGTTGAATCCAGCAGAAAA
MF288718.1	TTATTAATAAGGTGCCTGCGGTGCCAGAAACCGTTGAATCCAGCAGAAAA
MF288725.1	TTATTAATAAGGTGCCTGCGGTGCCAGAAACCGTTGAATCCAGCAGAAAA
MF288723.1	
	TTATTAATAAGGTGCCTGCGGTGCCAGAAACCGTTGAATCCAGCAGAAAA
MF288722.1	TTATTAATAAGGTGCCTGCGGTGCCAGAAACCGTTGAATCCAGCAGAAAA

MF288709.1	ACTTAGACACCTTAATGAAAAACGACGATTTCACAACATAGCTGGGCACT
MF288708.1	ACTTAGACACCTTAATGAAAAACGACGATTTCACAACATAGCTGGGCACT
MF288713.1	ACTTAGACACCTTAATGAAAAACGACGATTTCACAACATAGCTGGGCACT
MF288710.1	ACTTAGACACCTTAATGAAAAACGACGATTTCACAACATAGCTGGGCACT
112 200 / 2012	
MF288712.1	ACTTAGACACCTTAATGAAAAACGACGATTTCACAACATAGCTGGGCACT
MF288711.1	ACTTAGACACCTTAATGAAAAACGACGATTTCACAACATAGCTGGGCACT
MF288714.1	ACTTAGACACCTTAATGAAAAACGACGATTTCACAACATAGCTGGGCACT
MF288716.1	ACTTAGACACCTTAATGAAAAACGACGATTTCACAACATAGCTGGGCACT
MF288715.1	ACTTAGACACCTTAATGAAAAACGACGATTTCACAACATAGCTGGGCACT
MF288717.1	ACTTAGACACCTTAATGAAAAACGACGATTTCACAACATAGCTGGGCACT
MF288727.1	ACTTAGACACCTTAATGAAAAACGACGATTCCACAACATAGCTGGGCACT
MF288726.1	ACTTAGACACCTTAATGAAAAACGACGATTCCACAACATAGCTGGGCACT
MF288724.1	ACTTAGACACCTTAATGAAAAACGACGATTCCACAACATAGCTGGGCACT
MF288721.1	
	ACTTAGACACCTTAATGAAAAACGACGATTCCACAACATAGCTGGGCACT
MF288720.1	ACTTAGACACCTTAATGAAAAACGACGATTCCACAACATAGCTGGGCACT
MF288719.1	ACTTAGACACCTTAATGAAAAACGACGATTCCACAACATAGCTGGGCACT
MF288718.1	ACTTAGACACCTTAATGAAAAACGACGATTCCACAACATAGCTGGGCACT
MF288725.1	ACTTAGACACCTTAATGAAAAACGACGATTCCACAACATAGCTGGGCACT
MF288723.1	ACTTAGACACCTTAATGAAAAACGACGATTCCACAACATAGCTGGGCACT
MF288722.1	ACTTAGACACCTTAATGAAAAACGACGATTCCACAACATAGCTGGGCACT
112 200 / 22 12	************
MED 0 0 0 0 0 1	
MF288709.1	ATAGAGGCCAGTGCCATTCGTGCTGCAACCGAGCACGACAGGAACGACTC
MF288708.1	ATAGAGGCCAGTGCCATTCGTGCTGCAACCGAGCACGACAGGAACGACTC
MF288713.1	ATAGAGGCCAGTGCCATTCGTGCTGCAACCGAGCACGACAGGAACGACTC
MF288710.1	ATAGAGGCCAGTGCCATTCGTGCTGCAACCGAGCACGACAGGAACGACTC
MF288712.1	ATAGAGGCCAGTGCCATTCGTGCTGCAACCGAGCACGACAGGAACGACTC
MF288711.1	ATAGAGGCCAGTGCCATTCGTGCTGCAACCGAGCACGACAGGAACGACTC
MF288714.1	ATAGAGGCCAGTGCCATTCGTGCTGCAACCGAGCACGACAGGAACGACTC
MF288716.1	ATAGAGGCCAGTGCCATTCGTGCTGCAACCGAGCACGACAGGAACGACTC
MF288715.1	ATAGAGGCCAGTGCCATTCGTGCTGCAACCGAGCACGACAGGAACGACTC
MF288717.1	ATAGAGGCCAGTGCCATTCGTGCTGCAACCGAGCACGACAGGAACGACTC

CLUSTAL	2.1	multiple	sequence	Alignment
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MF288719.1

MF288718.1

MF288725.1

MF288723.1

MF288722.1

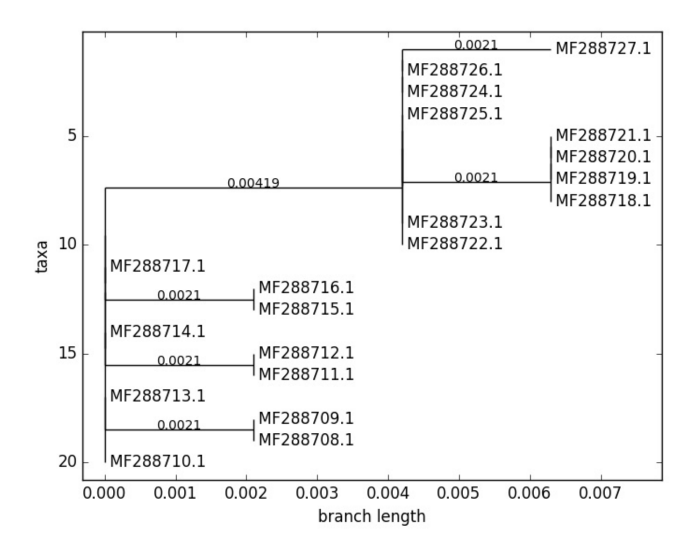
Chopiam 2.1 marcipic bequence ming	······································
MF288727.1	ATAGAGGCCAGTGCCATTCGTGCTGCAACCGAGCACGACAGGAAAGACTC
MF288726.1	ATAGAGGCCAGTGCCATTCGTGCTGCAACCGAGCACGACAGGAAAGACTC
MF288724.1	ATAGAGGCCAGTGCCATTCGTGCTGCAACCGAGCACGACAGGAAAGACTC
MF288721.1	ATAGAGGCCAGTGCCATTCGTGCTGCAACCGAGCACGACAGGAAAGACTC
MF288720.1	ATAGAGGCCAGTGCCATTCGTGCTGCAACCGAGCACGACAGGAAAGACTC
MF288719.1	ATAGAGGCCAGTGCCATTCGTGCTGCAACCGAGCACGACAGGAAAGACTC
MF288718.1	ATAGAGGCCAGTGCCATTCGTGCTGCAACCGAGCACGACAGGAAAGACTC
MF288725.1	ATAGAGGCCAGTGCCATTCGTGCTGCAACCGAGCACGACAGGAAAGACTC
MF288723.1	ATAGAGGCCAGTGCCATTCGTGCTGCAACCGAGCACGACAGGAAAGACTC
MF288722.1	ATAGAGGCCAGTGCCATTCGTGCTGCAACCGAGCACGACAGGAAAGACTC
	*********** ****
MF288709.1	CAACGACGCAGAAACACAAGTATAA
MF288708.1	CAACGACGCAGAAACACAAGTATAA
MF288713.1	CAACGACGCAGAAACACAAGTATAA
MF288710.1	CAACGACGCAGAAACACAAGTATAA
MF288712.1	CAACGACGCAGAAACACAAGTATAA
MF288711.1	CAACGACGCAGAAACACAAGTATAA
MF288714.1	CAACGACGCAGAAACACAAGTATAA
MF288716.1	CAACGACGCAGAAACACAAGTATAA
MF288715.1	CAACGACGCAGAAACACAAGTATAA
MF288717.1	CAACGACGCAGAAACACAAGTATAA
MF288727.1	CAACGACGCAGAAACACAAGTATAA
MF288726.1	CAACGACGCAGAAACACAAGTATAA
MF288724.1	CAACGACGCAGAAACACAAGTATAA
MF288721.1	CAACGACGCAGAAACACAAGTATAA
MF288720.1	CAACGACGCAGAAACACAAGTATAA

CAACGACGCAGAGAAACACAAGTATAA

CAACGACGCAGAGAAACACAAGTATAA

CAACGACGCAGAGAAACACAAGTATAA

CAACGACGCAGAGAAACACAAGTATAA



Consensus sequence

>/hpv_18_prueba/hpv_18_prueba_E6 Consensus threshold=0.7
ATGGCGCGCTTTGAGGATCCAACACGGCGACCCTACAAGCTACCTGATCTGTGCACGGAA
CTGAACACTTCACTGCAAGACATAGAAATAACCTGTGTATATTGCAAGACAGTATTGGAA
CTTACAGAGGTATTTGAATTTGCATTTAAAGATTTATTTGTGGTGTATAGAGACAGTATA
CCGCATGCTGCATGCCATAAATGTATAGATTTTTATTCTAGAATTAGAGAATTAAGACAT
TATTCAGACTCTGTGTATGGAGACACATTGGAAAAACTAACACTGGGTTATACAAT
TTATTAATAAGGTGCCTGCGGTGCCAGAAACCGTTGAATCCAGCAGAAAAACTTAGACAC
CTTAATGAAAAACGACGATTXCACAACATAGCTGGGCACTATAGAGGCCAGTGCCATTCG
TGCTGCAACCGAGCACGACAGGAAXGACTCCAACGACGCAGAGAAACACAAGTATAA

Primers data

ACCEPTABLE LEFT PRIMERS

0	-based #	self se	lf h	aiı	r- qua	al-			
#	sequence	start	ln	N	GC%	Tm	any_th	end_th pi	n lity
0	TGCATGGACCTAAGGCAACA	1	20	0	50.00	59.596	0.34	0.00 41.41	2.404
1	TGCATGGACCTAAGGCAACAT	1	21	0	47.62	59.994	0.34	0.00 41.41	3.006
2	ATGCATGGACCTAAGGCAACA	0	21	0	47.62	59.994	1.29	0.00 41.41	3.006
3	GCATGGACCTAAGGCAACA	2	19	0	52.63	57.735	0.00	0.00 41.41	3.265
4	TGCATGGACCTAAGGCAAC	1	19	0	52.63	57.735	0.34	0.00 0.00	3.265
5	GCATGGACCTAAGGCAACAT	2	20	0	50.00	58.232	0.00	0.00 41.41	3.768
6	ATGCATGGACCTAAGGCAAC	0	20	0	50.00	58.232	1.29	0.00 0.00	3.768
7	GCATGGACCTAAGGCAACATT	2	21	0	47.62	58.897	0.00	0.00 41.41	4.103
8	CATGGACCTAAGGCAACATTG	3	21	0	47.62	57.212	0.00	0.00 41.41	5.788

ACCEPTABLE RIGHT PRIMERS

0-based #	self self hair- qual-	
# sequence	start ln N GC% ${ m Tm}$ any_th	end_th pin lity
0 GGATGCACACCACGGACA	308 18 0 61.11 59.968 6.86	0.00 0.00 0.032
1 CTGCTGGGATGCACACCA	314 18 0 61.11 59.965 6.10	4.92 38.12 0.035
2 ACTGCTGGGATGCACACC	315 18 0 61.11 59.965 0.00	0.00 38.12 0.035
3 TGCTCGAAGGTCGTCTGC	254 18 0 61.11 59.741 6.26	6.26 43.45 0.259
4 GCTCGTCGGGCTGGTAAA	160 18 0 61.11 59.736 10.59	0.00 0.00 0.264
5 TGCTGGGATGCACCAC	313 18 0 61.11 60.282 3.46	0.78 38.12 0.282
6 TTGTGGTTCGGCTCGTCG	170 18 0 61.11 60.355 0.00	0.00 0.00 0.355
7 TGACGTTGTGGTTCGGCT	175 18 0 55.56 59.498 0.00	0.00 35.95 0.502
8 ATGCACACCACGGACACA	306 18 0 55.56 59.492 0.00	0.00 0.00 0.508
9 TGCACACCACGGACACAC	305 18 0 61.11 60.513 0.00	0.00 0.00 0.513
10 GCACACCACGGACACACA	304 18 0 61.11 60.513 0.00	0.00 0.00 0.513
11 GCTCGAAGGTCGTCTGCT	253 18 0 61.11 59.431 6.06	0.00 43.45 0.569
12 GTGACGTTGTGGTTCGGC	176 18 0 61.11 59.370 0.00	0.00 35.47 0.630
13 ACGTTGTGGTTCGGCTCG	173 18 0 61.11 60.661 0.00	0.00 0.00 0.661
14 CGTTGTGGTTCGGCTCGT	172 18 0 61.11 60.661 0.00	0.00 0.00 0.661
15 GGGATGCACACCACGGAC	309 18 0 66.67 60.746 12.79	0.00 0.00 0.746
16 CTGGGATGCACACCACGG	311 18 0 66.67 60.747 2.08	0.00 0.00 0.747
17 AGCTGCTGGAATGCTCGA	265 18 0 55.56 59.015 0.00	0.00 30.41 0.985
18 CAGCTGCTGGAATGCTCG	266 18 0 61.11 58.901 21.71	0.00 34.51 1.099
19 GCTCGTCGGGCTGGTAAAT	160 19 0 57.89 60.153 10.59	0.00 0.00 1.153
20 ATGCTCGAAGGTCGTCTGC	255 19 0 57.89 60.154 6.26	6.26 43.45 1.154
21 GACGTTGTGGTTCGGCTC	174 18 0 61.11 58.759 0.00	0.00 31.86 1.241
22 GTTGTGGTTCGGCTCGTC	171 18 0 61.11 58.759 0.00	0.00 0.00 1.241
23 TGGGATGCACACCACGGA	310 18 0 61.11 61.245 6.10	0.00 0.00 1.245
24 GATGCACACCACGGACAC	307 18 0 61.11 58.746 0.00	0.00 0.00 1.254
25 AGCTGCTGGAATGCTCGAA	265 19 0 52.63 59.702 0.00	0.00 30.41 1.298
26 TACTGCTGGGATGCACACC	316 19 0 57.89 59.700 0.00	0.00 38.12 1.300
27 TGTGGTTCGGCTCGTCG	169 17 0 64.71 59.693 0.00	0.00 0.00 1.307
28 TGGGATGCACACCACGG	310 17 0 64.71 59.592 6.10	0.00 0.00 1.408
29 GCTGGGATGCACACCACG	312 18 0 66.67 61.434 3.27	0.42 0.00 1.434

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Pr	imers data										
30	CAGCTGCTGGAATGCTCGA	266	19	0	57.	89	60.449	21.71	0.00 3	34.51	1.449
31	GTGGTTCGGCTCGTCGG	168	17	0	70.	59	60.495	0.00	0.00	0.00	1.495
32	GAAGGTCGTCTGCTGAGCT	249	19	0	57.	89	59.411	5.83	0.00	0.00	1.589
33	TGACGTTGTGGTTCGGCTC	175	19	0	57.	89	60.593	0.00	0.00 3	35.95	1.593
34	GATGCACACCACGGACACA	307	19	0	57.	89	60.597	0.00	0.00	0.00	1.597
35	GTGTGACGTTGTGGTTCGG	178	19	0	57.	89	59.360	0.00	0.00	0.00	1.640
36	TGCTGGAATGCTCGAAGGT	262	19	0	52.	63	59.321	0.00	0.00	0.00	1.679
37	GCTGCTGGAATGCTCGAAG	264	19	0	57.	89	59.280	0.00	0.00	0.00	1.720
38	GGGATGCACACCACGGA	309	17	0	64.	71	59.258	12.79	0.00	0.00	1.742
39	ACAGCTGCTGGAATGCTCG	267	19	0	57.	89	60.744	31.00	5.48 3	35.51	1.744
40	GGCTCGTCGGGCTGGTAA	161	18	0	66.	67	61.767	8.47	0.00	0.00	1.767
41	AAGGTCGTCTGCTGAGCT	248	18	0	55.	56	58.215	5.83	0.00	0.00	1.785
42	AGGTCGTCTGCTGAGCTT	247	18	0	55.	56	58.215	5.83	0.00	0.00	1.785
43	GCTCGAAGGTCGTCTGCTG	253	19	0	63.	16	60.804	6.06	0.00 4	13.45	1.804
44	CGAAGGTCGTCTGCTGAGC	250	19	0	63.	16	60.804	0.00	0.00 3	37.02	1.804
45	CTCGAAGGTCGTCTGCTGA	252	19	0	57.	89	59.123	0.00	0.00 4	13.45	1.877
46	TCGAAGGTCGTCTGCTGAG	251	19	0	57.	89	59.123	0.00	0.00 4	13.45	1.877
47	GTGACGTTGTGGTTCGGCT	176	19	0	57.	89	60.886	0.00	0.00 3	35.95	1.886
48	ATGCACACCACGGACACAC	306	19	0	57.	89	60.892	0.00	0.00	0.00	1.892
49	GCTGCTGGAATGCTCGAA	264	18	0	55.	56	58.105	0.00	0.00	0.00	1.895
50	ACCACGGACACAAAGGA	300	19	0	52.	63	59.091	0.00	0.00	0.00	1.909
51	TGGAATGCTCGAAGGTCGT	259	19	0	52.	63	59.028	0.00	0.00 4	13.45	1.972
52	GCTCGTCGGGCTGGTAA	160	17	0	64.	71	59.026	10.59	0.00	0.00	1.974
53	TCGTCGGGCTGGTAAATGT	158	19	0	52.	63	59.023	0.00	0.00	0.00	1.977
54	TGCACACCACGGACACA	305	17	0	58.	82	59.010	0.00	0.00	0.00	1.990
55	CACCACGGACACAAAGG	301	19				58.978	0.00	0.00	0.00	2.022
56	GAAGGTCGTCTGCTGAGCTT	249					60.038	7.47	7.47	38.04	2.038
57	AGGTCGTCTGCTGAGCTTTC	247					60.038	0.00	0.00	0.00	2.038
58	GGTCGTCTGCTGAGCTTTCT	246					60.038	0.00	0.00	0.00	2.038
59	AAGGTCGTCTGCTGAGCTT	248					58.946		4.25		2.054
	AGGTCGTCTGAGCTTT	247					58.946	5.83	0.00	0.00	2.054
	TCGAAGGTCGTCTGCTGA	251					57.925	0.00	0.00 4		2.075
	CTCGTCGGGCTGGTAAATGT	159					60.108	0.00	0.00	0.00	2.108
	CTGGAATGCTCGAAGGTCGT	260					60.109	0.00	0.00 4		2.109
	TGGAATGCTCGAAGGTCGTC	259					60.109	0.00	0.00 4		2.109
	GCACACCACGGACACAA	304					61.110	0.00	0.00	0.00	2.110
66		177					57.888	0.00	0.00	0.00	2.112
	GGCTCGTCGGGCTGGTA	161					61.167	8.47	0.00	0.00	2.167
	GGAATGCTCGAAGGTCGTCT	258					59.826	0.00	0.00 4		2.174
	TGTGACGTTGTGGTTCGGC	177					61.175	0.00	0.00 3		2.175
	GAAGGTCGTCTGCTGAGC	249					57.823	0.00	0.00	0.00	2.177
	ACGGACACAAAGGACAGG	297					60.179	0.00	0.00	0.00	2.179
	CTCGAAGGTCGTCTGCTGAG	252					60.179	0.00	0.00 4		2.179
	GGACACAAAGGACAGGGT	295					59.817	0.00	0.00	0.00	2.183
	GGTTCGGCTCGTCGG	166					59.806	0.00	0.00	0.00	2.194
	CACACCACGGACACACAA	303					57.790	0.00	0.00	0.00	2.210
76	CGTTGTGGTTCGGCTCG	172	Τ./	U	64.	/ 1	58.750	0.00	0.00	0.00	2.250

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Primers data			
77 TGCTCGAAGGTCGTCTGCT	254 19	0 57.89 61.269 1	1.58 0.00 43.45 2.269
78 TGCTGGGATGCACACCA	313 17	0 58.82 58.729	6.10 4.92 38.12 2.271
79 ACAGCTGCTGGAATGCTC	267 18	0 55.56 57.684 3	1.00 0.00 35.51 2.316
80 CCACGGACACAAAGGAC	299 19	0 57.89 58.683	0.00 0.00 0.00 2.317
81 TCGTCGGGCTGGTAAATGTT	158 20	0 50.00 59.678	0.00 0.00 0.00 2.322
82 TTACTGCTGGGATGCACACC	317 20	0 55.00 60.322	0.00 0.00 38.12 2.322
83 GGATGCACACCACGGACAC	308 19	0 63.16 61.325	6.86 0.00 0.00 2.325
84 CTGCTGGGATGCACACCAC	314 19	0 63.16 61.338	3.46 0.78 38.12 2.338
85 TGTGGTTCGGCTCGTCGG	169 18	0 66.67 62.352	0.00 0.00 0.00 2.352
86 TGTGTGACGTTGTGGTTCG	179 19	0 52.63 58.625	0.00 0.00 0.00 2.375
87 CACACCACGGACACAAAG	303 20	0 55.00 59.623	0.00 0.00 0.00 2.377
88 GCTGGAATGCTCGAAGGTC	261 19	0 57.89 58.612	0.00 0.00 0.00 2.388
89 CTGCTGGAATGCTCGAAGG	263 19	0 57.89 58.609	0.00 0.00 0.00 2.391
90 TGCTGGAATGCTCGAAGGTC	262 20	0 55.00 60.392	0.00 0.00 0.00 2.392
91 CTGCTGGAATGCTCGAAGGT	263 20		0.00 0.00 0.00 2.392
92 AAGGTCGTCTGCTGAGCTTT	248 20	0 50.00 59.605	4.25 0.00 34.76 2.395
93 GCACACCACGGACACAC	304 17	0 64.71 58.565	0.00 0.00 0.00 2.435
94 CGTCGGGCTGGTAAATGTTG	157 20	0 55.00 59.554	0.00 0.00 0.00 2.446
95 GGTCGTCTGCTGAGCTTTC	246 19	0 57.89 58.549	0.00 0.00 0.00 2.451
96 CACCACGGACACAAAGGA	301 20	0 55.00 60.463	0.00 0.00 0.00 2.463
97 CCACGGACACACAAAGGACA	299 20	0 55.00 60.463	0.00 0.00 0.00 2.463
98 ACCACGGACACACAAAGGAC	300 20	0 55.00 60.463	0.00 0.00 0.00 2.463
99 CTCGAAGGTCGTCTGCTG	252 18		0.00 0.00 0.00 2.467
100 CGAAGGTCGTCTGCTGAG	250 18		0.00 0.00 43.43 2.467
101 CACACCACGGACACAAA	303 19		0.00 0.00 37.02 2.407
102 ACCACGGACACAAAGG	300 18		0.00 0.00 0.00 2.472
103 AACAGCTGCTGGAATGCTC	268 19		
104 GTTCGGCTCGTCGGGC	165 16		
105 CGTCGGGCTGGTAAATGT	157 18	0 55.56 57.390 0 57.89 58.383	0.00 0.00 0.00 2.610 0.00 0.00 0.00 2.617
106 CGGACACAAAGGACAGG			
107 ACACACAAAGGACAGGGTGT	293 20		0.00 0.00 44.69 2.624
108 TGCTGGAATGCTCGAAGG	262 18		0.00 0.00 0.00 2.624
109 GCTGGAATGCTCGAAGGT	261 18		0.00 0.00 0.00 2.626
110 GACGTTGTGGTTCGGCTCG	174 19		0.00 0.00 31.86 2.655
111 CGTTGTGGTTCGGCTCGTC	172 19		0.00 0.00 0.00 2.655
112 GTTGTGGTTCGGCTCGTCG	171 19		0.00 0.00 0.00 2.655
113 GGAATGCTCGAAGGTCGTC	258 19		0.00 0.00 43.45 2.667
114 CTGGAATGCTCGAAGGTCG	260 19		0.00 0.00 44.56 2.671
115 CTCGTCGGGCTGGTAAATG	159 19		0.00 0.00 0.00 2.678
116 GGTCGTCTGCTGAGCTTT	246 18		5.83 0.00 0.00 2.683
117 GCTGGGATGCACACCAC	312 17		0.36 0.00 0.00 2.716
118 TTGTGTGACGTTGTGGTTCG	180 20		0.00 0.00 0.00 2.722
119 AATGCTCGAAGGTCGTCTGC	256 20		6.26 6.26 43.45 2.738
120 AGCTGCTGGAATGCTCGAAG	265 20		0.00 0.00 30.41 2.744
121 ACACCACGGACACAAAGG	302 20		0.00 0.00 0.00 2.747
122 TTCGGCTCGTCGGGCT	164 16		
123 ACACCACGGACACAAAG	302 19	0 52.63 58.225	0.00 0.00 0.00 2.775

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124	TGGTTCGGCTCGTCGGG	167	17	0	70.59	61.808	0.00	0.00	0.00	2.808
125	ACTGCTGGGATGCACACCA	315	19	0	57.89	61.833	6.10	4.92	38.12	2.833
126	AACAGCTGCTGGAATGCT	268	18	0	50.00	57.163	31.00	8.45	35.51	2.837
127	CGTCGGGCTGGTAAATGTT	157	19	0	52.63	58.151	0.00	0.00	0.00	2.849
128	CCACGGACACAAAGGA	299	18	0	55.56	57.132	0.00	0.00	0.00	2.868
129	CGGACACAAAGGACAGGG	296	20	0	60.00	60.885	0.00	0.00	0.00	2.885
130	AGAAACAGCTGCTGGAATGC	271	20	0	50.00	59.113	11.85	0.18	35.51	2.887
131	GAAACAGCTGCTGGAATGCT	270	20	0	50.00	59.113	12.08	0.00	35.51	2.887
132	AAACAGCTGCTGGAATGCTC	269	20	0	50.00	59.113	31.00	0.00	35.51	2.887
133	TGGAATGCTCGAAGGTCG	259	18	0	55.56	57.089	0.00	0.00	44.56	2.911
134	GGAATGCTCGAAGGTCGT	258	18	0	55.56	57.087	0.00	0.00	43.45	2.913
135	TCGTCGGGCTGGTAAATG	158	18	0	55.56	57.076	0.00	0.00	0.00	2.924
136	CACGGACACAAAGGACAG	298	20	0	55.00	59.063	0.00	0.00	0.00	2.937
137	GTGTTCAGAAACAGCTGCTGG	277	21	0	52.38	60.002	31.00	14.58	45.78	3.002
138	ACACACAAAGGACAGGGTGTT	293	21	0	47.62	59.995	0.00	0.00	42.16	3.005
139	TCGGCTCGTCGGGCT	163				60.018		0.00	0.00	3.018
140	GGACACAAAGGACAGGG	295				57.979	0.00	0.00	0.00	3.021
141		266				61.023			34.51	3.023
142	GGATGCACACCACGGAC	308				57.975	6.86	0.00	0.00	3.025
143	CTGGGATGCACACCACG	311				57.970	0.00	0.00	0.00	3.030
	AAACAGCTGCTGGAATGCT	269				57.953			35.51	3.047
145	CTGCTGGGATGCACACC	314				57.947	0.00		38.12	3.053
146	TGGTTCGGCTCGTCGG	167				58.943	0.00	0.00	0.00	3.057
147	CACGGACACAAAGGACA	298				57.927	0.00	0.00	0.00	3.073
148	TGTGTGACGTTGTGGTTCGG	179				61.085	0.00	0.00	0.00	3.085
149	TGTTCAGAAACAGCTGCTGGA	276	21			60.134				3.134
150	GAAACAGCTGCTGGAATGCTC	270	21			60.135			35.51	3.135
151	CGGCTCGTCGGGCTG	162				59.845		8.50	0.00	3.155
152	TGACGTTGTGGTTCGGC	175		0	58.82	57.822	0.00	0.00	35.47	3.178
153	TTCGGCTCGTCGGGCTG	164	17			62.201	6.26			3.201
	GTTCGGCTCGTCGGGCT	165				62.203			0.00	3.203
155	GGTCGTCTGCTGAGCTTTCTA	246	21	0	52.38	59.797	0.00	0.00	0.00	3.203
156	GCTCGAAGGTCGTCTGC	253	17	0	64.71	57.778	0.00	0.00	43.45	3.222
157	TTACTGCTGGGATGCACAC	317	19	0	52.63	57.752	0.00	0.00	38.12	3.248
158	CTGGGATGCACACCACGGA	311	19	0	63.16	62.253	2.08	0.00	0.00	3.253
159	GAATGCTCGAAGGTCGTCTG	257	20	0	55.00	58.727	0.00	0.00	43.45	3.273
160	ACACAAAGGACAGGGTGTTCA	291	21	0	47.62	59.716	0.00	0.00	41.36	3.284
161	AACAGCTGCTGGAATGCTCG	268	20	0	55.00	61.304	31.00	5.48	35.51	3.304
162	GGCTCGTCGGGCTGGTAAA	161	19	0	63.16	62.305	8.47	0.00	0.00	3.305
163	GACACAAAGGACAGGGTG	294	20	0	55.00	58.691	0.00	0.00	37.29	3.309
164	TGTTCAGAAACAGCTGCTGG	276	20	0	50.00	58.688	31.00	14.58	40.45	3.312
165	TGCACACCACGGACACA	305	19	0	57.89	62.317	0.00	0.00	0.00	3.317
166	ATTGTGTGACGTTGTGGTTCG	181		0	47.62	59.671	0.00	0.00	0.00	3.329
167	ACGGACACAAAGGACAG	297				57.620	0.00	0.00	0.00	3.380
168	CAGAAACAGCTGCTGGAATGC	272		0	52.38	60.402	12.65	0.18	35.51	3.402
169	GACACACAAAGGACAGGGTGT	294				60.408	0.00		44.69	3.408
170	GAAACAGCTGCTGGAATGC	270	19			57.586			35.51	3.414

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171	GCTGGAATGCTCGAAGGTCG	261	20	0	60.00	61.424	0.00	0.00	44.56	3.424
172	AATGCTCGAAGGTCGTCTG	256	19	0	52.63	57.574	0.00	0.00	43.45	3.426
173	GCTCGTCGGGCTGGTAAATG	160	20	0	60.00	61.429	10.59	0.00	0.00	3.429
174	ACGTTGTGGTTCGGCTCGT	173	19	0	57.89	62.435	0.00	0.00	0.00	3.435
175	TACTGCTGGGATGCACACCA	316	20	0	55.00	61.488	6.10	4.92	38.12	3.488
176	ACACACAAAGGACAGGGTG	293	19	0	52.63	57.505	0.00	0.00	37.29	3.495
177	CACACAAAGGACAGGGTGT	292	19	0	52.63	57.505	0.00	0.00	44.69	3.495
178	GTCGTCTGCTGAGCTTTCT	245	19	0	52.63	57.494	0.00	0.00	0.00	3.506
179	GACGTTGTGGTTCGGCT	174	17	0	58.82	57.486	0.00	0.00	31.86	3.514
180	ACGTTGTGGTTCGGCTC	173	17	0	58.82	57.486	0.00	0.00	0.00	3.514
181	GTTGTGGTTCGGCTCGT	171	17	0	58.82	57.486	0.00	0.00	0.00	3.514
182	ATGCACACCACGGACAC	306	17	0	58.82	57.460	0.00	0.00	0.00	3.540
183	GGCTCGTCGGGCTGGT	161	16	0	75.00	61.543	8.47	0.00	0.00	3.543
184	TGGGATGCACACCACGGAC	310	19	0	63.16	62.545	6.10	0.00	0.00	3.545
185	GGGATGCACACCACGGACA	309	19	0	63.16	62.545	12.79	0.00	0.00	3.545
186	AGAAACAGCTGCTGGAATGCT	271	21	0	47.62	60.547	14.60	5.01	35.51	3.547
187	GTTCAGAAACAGCTGCTGGA	275	20	0	50.00	58.402	31.00	16.80	35.51	3.598
188	ATGCTCGAAGGTCGTCTGCT	255	20	0	55.00	61.600	11.58	0.55	43.45	3.600
189	AGGTCGTCTGAGCT	247	17	0	58.82	57.400	5.83	0.00	0.00	3.600
190	TCGGCTCGTCGGGCTG	163	16	0	75.00	61.604	6.26	6.26	0.00	3.604
191	GAAGGTCGTCTGCTGAGCTTT	249	21	0	52.38	60.607	8.40	2.28	36.71	3.607
192	AAGGTCGTCTGCTGAGCTTTC	248	21	0	52.38	60.607	4.25	0.00	34.76	3.607
193	GGCTCGTCGGGCTGG	161	15	0	80.00	59.383	8.47	0.00	0.00	3.617
194	GCACACCACGGACACAAA	304	20	0	55.00	61.649	0.00	0.00	0.00	3.649
195	CTCGTCGGGCTGGTAAATGTT	159	21	0	52.38	60.675	0.00	0.00	0.00	3.675
196	CACACAAAGGACAGGGTGTTC	292	21	0	52.38	59.324	0.00	0.00	42.16	3.676
197	GCTGCTGGAATGCTCGA	264	17	0	58.82	57.303	0.00	0.00	0.00	3.697
198	AGCTGCTGGAATGCTCG	265	17	0	58.82	57.295	0.00	0.00	30.41	3.705
199	GCTGCTGGAATGCTCGAAGG	264	20	0	60.00	61.713	0.00	0.00	0.00	3.713
200	CGTTTTCTTCCTCTGAGTCGC	114	21	0	52.38	59.278	0.00	0.00	0.00	3.722
201	GAATGCTCGAAGGTCGTCT	257	19	0	52.63	57.276	0.00	0.00	43.45	3.724
202	ACATTGTGTGACGTTGTGGT	183	20	0	45.00	58.255	0.00	0.00	0.00	3.745
203	CACACAAAGGACAGGGTGTT	292	20	0	50.00	58.237	0.00	0.00	42.16	3.763
204	GCTCGTCGGGCTGGTA	160	16	0	68.75	58.231	10.59	0.00	0.00	3.769
205	GACACAAAGGACAGGGT	294	19	0	52.63	57.199	0.00	0.00	0.00	3.801
206	GTCGGGCTGGTAAATGTTGA	156	20	0	50.00	58.184	0.00	0.00	0.00	3.816
207	GTGACGTTGTGGTTCGGCTC	176	20	0	60.00	61.828	0.00	0.00	35.95	3.828
208	GATGCACACCACGGACACAC	307	20	0	60.00	61.842	0.00	0.00	0.00	3.842
209	TTGTGGTTCGGCTCGTC	170	17	0	58.82	57.158	0.00	0.00	0.00	3.842
210	ACAAAGGACAGGGTGTTCAGA	289	21	0	47.62	59.156	0.00	0.00	0.00	3.844
211	AGGACAGGGTGTTCAGAAACA	285	21	0	47.62	59.156	0.01	0.00	43.06	3.844
212	TTGTGGTTCGGCTCGTCGG	170	19	0	63.16	62.852	0.00	0.00	0.00	3.852
213	GATGCACACCACGGACA	307	17	0	58.82	57.129	0.00	0.00	0.00	3.871
214	ACTGCTGGGATGCACAC	315	17	0	58.82	57.088	0.00	0.00	38.12	3.912
215	TCGTCGGGCTGGTAAATGTTG	158	21	0	52.38	60.942	0.00	0.00	0.00	3.942
216	CGTCGGGCTGGTAAATGTTGA	157	21	0	52.38	60.942	0.00	0.00	0.00	3.942
217	CACAAAGGACAGGGTGTTCAG	290	21	0	52.38	59.050	0.00	0.00	0.00	3.950

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218	GTTCAGAAACAGCTGCTGGAA	275 21	0 47	.62	59.050	20.26	11.82	35.51	3.950
219	TGAGTCGCTTAATTGCTCGTG	101 21	0 47	.62	59.007	0.00	0.00	0.00	3.993
220	AGTCGCTTAATTGCTCGTGAC	99 21	0 47	.62	59.007	4.62	4.62	0.00	3.993
221	CACAAAGGACAGGGTGTTCA	290 20	0 50	.00	57.949	0.00	0.00	0.00	4.051
222	ACACAAAGGACAGGGTGTTC	291 20	0 50	.00	57.947	0.00	0.00	41.36	4.053
223	CTGGAATGCTCGAAGGTCGTC	260 21	0 57	.14	61.064	0.00	0.00	43.45	4.064
224	GGAATGCTCGAAGGTCGTCTG	258 21	0 57	.14	61.064	0.00	0.00	43.45	4.064
225	TTCGGCTCGTCGGGC	164 15	0 73	.33	58.930	2.73	2.73	0.00	4.070
226	GGACACAAAGGACAGGGTG	295 21	0 57	.14	61.085	0.00	0.00	37.29	4.085
227	AACATTGTGTGACGTTGTGGT	184 21	0 42	.86	58.912	0.00	0.00	0.00	4.088
228	ACATTGTGTGACGTTGTGGTT	183 21	0 42	. 86	58.912	0.00	0.00	0.00	4.088
229	TCAGAAACAGCTGCTGGAATG	273 21			58.843	22.36			4.157
230	AGCTCAATTCTGGCTTCACAC	220 21			58.842	0.00	0.00	0.00	4.158
231	GCTCAATTCTGGCTTCACACT	219 21			58.842	0.00	0.00	0.00	4.158
232	TCGTCTGCTGAGCTTTCTACT	244 21			58.836	0.18	0.00	0.00	4.164
233	ACAGCTGCTGGAATGCTCGA	267 20			62.176	31.00		35.51	4.176
234		253 20			62.201	6.06		43.45	4.201
235	TCGAAGGTCGTCTGCTGAGC	251 20			62.201	0.00		43.45	4.201
236	CGAAGGTCGTCTGCTGAGCT	251 20			62.201	5.83		37.02	4.201
237	GGACAGGGTGTTCAGAAACAG	284 21			58.778	0.00		45.78	4.222
238	GTCGTCTGCTGAGCTTTCTAC	245 21			58.745	0.00	0.00	0.00	4.255
239	GAGTCGCTTAATTGCTCGTGA	100 21			58.740	0.00	0.00	0.00	4.260
240	CTGAGTCGCTTAATTGCTCGT	102 21			58.737	0.00	0.00	0.00	4.263
241	CTGCTGGAATGCTCGAAGGTC	263 21			61.338	0.00	0.00	0.00	4.338
242	GGACAGGGTGTTCAGAAACA	284 20			57.658	0.01		43.06	4.342
243	TTCAGAAACAGCTGCTGGAA	274 20			57.653		12.30		4.347
244	ACAAAGGACAGGGTGTTCAG	289 20			57.652	0.00	0.00	0.00	4.348
245	TGAGTCGCTTAATTGCTCGT	101 20	0 45	.00	57.636	0.00	0.00	0.00	4.364
	AGTCGCTTAATTGCTCGTGA	99 20			57.636	0.00	0.00	0.00	4.364
247	TCGGGCTGGTAAATGTTGATG	155 21			58.635	0.00	0.00		4.365
	CGGGCTGGTAAATGTTGATGA	154 21	0 47	.62	58.635	0.00	0.00	0.00	4.365
249	GTCGGGCTGGTAAATGTTGAT	156 21	0 47	.62	58.634	0.00	0.00	0.00	4.366
250	GTGTGACGTTGTGGTTCGGC	178 20			62.369	0.00	0.00	35.47	4.369
251	GTGGTTCGGCTCGTCG	168 16	0 68	.75	57.630	0.00	0.00	0.00	4.370
252	GTGTTCAGAAACAGCTGCTG	277 20	0 50	.00	57.604	27.02	27.02	45.78	4.396
253	CCACGGACACAAAGGACAG	299 21	0 57	.14	61.406	0.00	0.00	0.00	4.406
254	CACGGACACAAAGGACAGG	298 21	0 57	.14	61.406	0.00	0.00	0.00	4.406
255	AGGTCGTCTGCTGAGCTTTCT	247 21	0 52	.38	61.436	0.00	0.00	0.00	4.436
256	CAACATTGTGTGACGTTGTGG	185 21	0 47	.62	58.551	0.00	0.00	0.00	4.449
257	TGCTCGAAGGTCGTCTGCTG	254 20	0 60	.00	62.478	11.58	0.00	43.45	4.478
258	TGGAATGCTCGAAGGTCGTCT	259 21	0 52	.38	61.498	0.00	0.00	43.45	4.498
259	TCTGAGTCGCTTAATTGCTCG	103 21	0 47	.62	58.470	0.00	0.00	0.00	4.530
260	GGGATGCACCACGG	309 16	0 68	.75	57.451	12.79	0.00	0.00	4.549
261	GCTCGTCGGGCTGGT	160 15	0 73	.33	58.449	10.59	0.00	0.00	4.551
262	TCAGAAACAGCTGCTGGAAT	273 20	0 45	.00	57.426	31.00	17.58	35.51	4.574
263	AGCTCAATTCTGGCTTCACA	220 20	0 45	.00	57.426	0.00	0.00	0.00	4.574
264	TGTGACGTTGTGGTTCGGCT	177 20	0 55	.00	62.581	0.00	0.00	35.95	4.581

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Pri	mers data									
265	GGCTCGTCGGGCTGGTAAAT	161	20	0	60.00	62.589	8.47	0.00	0.00	4.589
266	TTGTGTGACGTTGTGGTTCGG	180	21	0	52.38	61.599	0.00	0.00	0.00	4.599
267	ATGCACACCACGGACACACA	306	20	0	55.00	62.601	0.00	0.00	0.00	4.601
268	GCTCAATTCTGGCTTCACAC	219	20	0	50.00	57.374	0.00	0.00	0.00	4.626
269	CAGAAACAGCTGCTGGAATG	272	20	0	50.00	57.369	19.76	10.50	35.51	4.631
270	CAAAGGACAGGGTGTTCAGA	288	20	0	50.00	57.362	0.00	0.00	0.00	4.638
271	GTCGTCTGCTGAGCTTTCTA	245	20	0	50.00	57.361	0.00	0.00	0.00	4.639
272	TCGTCTGCTGAGCTTTCTAC	244	20	0	50.00	57.361	0.00	0.00	0.00	4.639
273	AGGACAGGGTGTTCAGAAAC	285	20	0	50.00	57.360	0.00	0.00	35.85	4.640
274	CGTCTGCTGAGCTTTCTACT	243	20	0	50.00	57.354	0.18	0.00	0.00	4.646
275	GAATGCTCGAAGGTCGTCTGC	257	21	0	57.14	61.648	6.26	6.26	43.45	4.648
276	CACCACGGACACAAAGGAC	301	21	0	57.14	61.670	0.00	0.00	0.00	4.670
277	GAGTCGCTTAATTGCTCGTG	100	20	0	50.00	57.297	0.00	0.00	0.00	4.703
278	CATTGTGTGACGTTGTGGTTC	182	21	0	47.62	58.285	0.00	0.00	0.00	4.715
279	TCGGGCTGGTAAATGTTGAT	155	20	0	45.00	57.201	0.00	0.00	0.00	4.799
280	TGCACACCACGGACACAA	305	20	0	55.00	62.805	0.00	0.00	0.00	4.805
281	AAACAGCTGCTGGAATGCTCG	269	21	0	52.38	61.812	31.00	5.48	35.51	4.812
282	AACATTGTGTGACGTTGTGG	184	20	0	45.00	57.158	0.00	0.00	0.00	4.842
283	CATTGTGTGACGTTGTGGTT	182	20	0	45.00	57.158	0.00	0.00	0.00	4.842
284	CGGGCTGGTAAATGTTGATG	154	20	0	50.00	57.150	0.00	0.00	0.00	4.850
285	TTCAGAAACAGCTGCTGGAAT	274	21	0	42.86	58.132	21.26	9.35	35.51	4.868
286	TCCTCTGAGTCGCTTAATTGC	106	21	0	47.62	58.106	3.66	0.00	0.00	4.894
287	CCTCTGAGTCGCTTAATTGCT	105	21	0	47.62	58.101	3.66	0.00	0.00	4.899
288	CAGCTGCTGGAATGCTCGAAG	266	21	0	57.14	61.927	21.71	0.00	34.51	4.927
289	CAAAGGACAGGGTGTTCAGAA	288	21	0	47.62	58.069	0.00	0.00	0.00	4.931
290	AAGGACAGGGTGTTCAGAAAC	286	21	0	47.62	58.068	0.00	0.00	35.85	4.932
291	CACACCACGGACACAAAGG	303	21	0	57.14	61.935	0.00	0.00	0.00	4.935
292	CTGAGTCGCTTAATTGCTCG	102	20	0	50.00	57.013	0.00	0.00	0.00	4.987
293	TGGCTTCACACTTACAACACA	210	21	0	42.86	58.007	0.00	0.00	0.00	4.993
294	TTACTGCTGGGATGCACACCA	317	21	0	52.38	62.004	6.10	4.92	38.12	5.004
295	GTTTTCTTCCTCTGAGTCGCT	113	21	0	47.62	57.967	0.00	0.00	0.00	5.033
296	AATGCTCGAAGGTCGTCTGCT	256	21	0	52.38	62.099	11.58	0.55	43.45	5.099
297	ACACCACGGACACAAAGGA	302	21	0	52.38	62.124	0.00	0.00	0.00	5.124
298	ACCACGGACACAAAGGACA	300	21	0	52.38	62.124	0.00	0.00	0.00	5.124
299	ACAACATTGTGTGACGTTGTG	186	21	0	42.86	57.870	5.36	0.00	41.89	5.130
300	TCGTTTTCTTCCTCTGAGTCG	115	21	0	47.62	57.716	0.00	0.00	0.00	5.284
301	GCACACCACGGACACACAAAG	304	21	0	57.14	62.514	0.00	0.00	0.00	5.514
302	ACGGACACAAAGGACAGGG	297	21	0	57.14	62.522	0.00	0.00	0.00	5.522
303	CGGACACAAAGGACAGGGT	296	21	0	57.14	62.522	0.00	0.00	0.00	5.522
304	TCTGGCTTCACACTTACAACA	212	21	0	42.86	57.448	0.00	0.00	0.00	5.552
305	CTGGCTTCACACTTACAACAC	211	21	0	47.62	57.391	0.00	0.00	0.00	5.609
306	AACAGCTGCTGGAATGCTCGA	268	21	0	52.38	62.651	31.00	0.00	35.51	5.651
307	ACAGCTGCTGGAATGCTCGAA	267	21	0	52.38	62.651	31.00	0.00	35.51	5.651
308	TACTGCTGGGATGCACACCAC	316	21	0	57.14	62.660	3.46	0.78	38.12	5.660
309	AAAGGACAGGGTGTTCAGAAA	287	21	0	42.86	57.337	0.00	0.00	0.00	5.663
310	CGAAGGTCGTCTGCTGAGCTT	250	21	0	57.14	62.671	7.47	7.47	38.04	5.671
311	TAGCTCAATTCTGGCTTCACA	221	21	0	42.86	57.299	2.49	0.00	33.07	5.701

315 ACTAGCTCAATTCTGGCTTCA	223 21	0 42.86 57.011	0.00 0.00	33.07	5.989
21 - 2002 0000 2000000	222 21	0 42.86 57.011	0 00 0 00	22 07	F 000
314 CGTCTGCTGAGCTTTCTACTA	243 21	0 47.62 57.233	0.18 0.00	0.00	5.767
313 TTTTCTTCCTCTGAGTCGCTT	112 21	0 42.86 57.247	0.00 0.00	0.00	5.753

312 ATGCTCGAAGGTCGTCTGCTG 255 21 0 57.14 62.738 11.58 0.00 43.45 5.738

BLASTN 2.7.1+

Reference: Stephen F. Altschul, Thomas L. Madden, Alejandro A. Schaffer, Jinghui Zhang, Zheng Zhang, Webb Miller, and David J. Lipman (1997), "Gapped BLAST and PSI-BLAST: a new generation of protein database search programs", Nucleic Acids Res. 25:3389-3402.

RID: 09A39ENW014

Database: Nucleotide collection (nt)

45,172,210 sequences; 162,240,808,426 total letters

Query= Length=20

Score E

Sequences producing significant alignments: (Bits) Value						
MF288727.1	Human papillomavirus type 18 isolate 1494573_N-P,	37.4	2.8			
MF288726.1	Human papillomavirus type 18 isolate 1314304_N-P,	37.4	2.8			
MF288725.1	Human papillomavirus type 18 isolate C451606_R3_19	37.4	2.8			
MF288724.1	Human papillomavirus type 18 isolate C451606_R2_15	37.4	2.8			
MF288723.1	Human papillomavirus type 18 isolate C451606_R1_12	37.4	2.8			
MF288722.1	Human papillomavirus type 18 isolate C387957_R1_11	37.4	2.8			
MF288721.1	Human papillomavirus type 18 isolate 1209155_R1_P	37.4	2.8			
MF288720.1	Human papillomavirus type 18 isolate 1174820_N-P,	37.4	2.8			
MF288719.1	Human papillomavirus type 18 isolate C487627_R2_15	37.4	2.8			
MF288718.1	Human papillomavirus type 18 isolate C487627_R1_11	37.4	2.8			
MF288717.1	Human papillomavirus type 18 isolate 1533526_N-P,	37.4	2.8			
MF288716.1	Human papillomavirus type 18 isolate C458963_R2_15	37.4	2.8			
MF288715.1	Human papillomavirus type 18 isolate C458963_R1_12	37.4	2.8			
MF288714.1	Human papillomavirus type 18 isolate C533713_R3_17	37.4	2.8			
MF288713.1	Human papillomavirus type 18 isolate C533713_R2_14	37.4	2.8			
MF288712.1	Human papillomavirus type 18 isolate C578032_R3_17	37.4	2.8			
MF288711.1	Human papillomavirus type 18 isolate C578032_R2_14	37.4	2.8			
MF288710.1	Human papillomavirus type 18 isolate C335048_R3_19	37.4	2.8			
MF288709.1	Human papillomavirus type 18 isolate C412607_R2_15	37.4	2.8			
MF288708.1	Human papillomavirus type 18 isolate C412607_R1_12	37.4	2.8			
MF288707.1	Human papillomavirus type 18 isolate C440384_R1_13	37.4	2.8			
MF288706.1	Human papillomavirus type 18 isolate C425046_R2_16	37.4	2.8			
MF288705.1	Human papillomavirus type 18 isolate C425046_R1_13	37.4	2.8			
MF288704.1	Human papillomavirus type 18 isolate C627893_R2_16	37.4	2.8			
MF288703.1	Human papillomavirus type 18 isolate C627893_R1_13	37.4	2.8			
MF288702.1	Human papillomavirus type 18 isolate C644657_R4_21	37.4	2.8			
MF288701.1	Human papillomavirus type 18 isolate C644657_R3_18	37.4	2.8			

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MF288698.1 Human papillomavirus type 18 isolate C335048 R2 15... 37.4
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MF288697.1 Human papillomavirus type 18 isolate 1445230_R2_P_... 37.4
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MF288692.1 Human papillomavirus type 18 isolate C353934 R1 11... 37.4
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MF288691.1 Human papillomavirus type 18 isolate 1390622_R1_P_... 37.4
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MF288690.1 Human papillomavirus type 18 isolate C532476_R1_11... 37.4
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MF288686.1 Human papillomavirus type 18 isolate C575918_R2_14... 37.4
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MF288685.1 Human papillomavirus type 18 isolate C575918_R1_10... 37.4
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MF288684.1 Human papillomavirus type 18 isolate 1389481_N-P, ... 37.4
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MF288683.1 Human papillomavirus type 18 isolate 1310091_R1_P_... 37.4
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MF288682.1 Human papillomavirus type 18 isolate C385149_R2_15... 37.4
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MF288681.1 Human papillomavirus type 18 isolate C385149_R1_11... 37.4
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MF288680.1 Human papillomavirus type 18 isolate C340709_R1_12... 37.4
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MF288679.1 Human papillomavirus type 18 isolate C310065_R1_13... 37.4
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MF288678.1 Human papillomavirus type 18 isolate C498643_R2_15... 37.4
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ALIGNMENTS

>MF288727.1 Human papillomavirus type 18 isolate 1494573_N-P, complete genome Length=7844

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Score = 37.4 bits (40), Expect = 2.8
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
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>MF288726.1 Human papillomavirus type 18 isolate 1314304_N-P, complete genome Length=7857

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Score = 37.4 bits (40), Expect = 2.8
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
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complete genome
Length=7857
Score = 37.4 bits (40), Expect = 2.8
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
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Sbjct 591 TGCATGGACCTAAGGCAACA 610
>MF288724.1 Human papillomavirus type 18 isolate C451606_R2_1584906_(52w),
complete genome
Length=7857
Score = 37.4 bits (40), Expect = 2.8
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
Query 1 TGCATGGACCTAAGGCAACA 20
Sbjct 591 TGCATGGACCTAAGGCAACA 610
>MF288723.1 Human papillomavirus type 18 isolate C451606_R1_1201181_(0w),
complete genome
Length=7857
Score = 37.4 bits (40), Expect = 2.8
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
          TGCATGGACCTAAGGCAACA 20
Query 1
Sbjct 591 TGCATGGACCTAAGGCAACA 610
>MF288722.1 Human papillomavirus type 18 isolate C387957_R1_1124102_N-P,
complete genome
Length=7857
Score = 37.4 bits (40), Expect = 2.8
Identities = 20/20 (100%), Gaps = 0/20 (0%)
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Strand=Plus/Plus

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Query 1 TGCATGGACCTAAGGCAACA 20
Sbjct 591 TGCATGGACCTAAGGCAACA 610
>MF288721.1 Human papillomavirus type 18 isolate 1209155_R1_P_R2_FAIL, complete
genome
Length=7857
Score = 37.4 bits (40), Expect = 2.8
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
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Query 1
Sbjct 591 TGCATGGACCTAAGGCAACA 610
>MF288720.1 Human papillomavirus type 18 isolate 1174820_N-P, complete genome
Length=7857
Score = 37.4 bits (40), Expect = 2.8
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
Query 1
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Sbjct 591 TGCATGGACCTAAGGCAACA 610
>MF288719.1 Human papillomavirus type 18 isolate C487627_R2_1553725_(52w),
complete genome
Length=7857
Score = 37.4 bits (40), Expect = 2.8
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
Query 1
         TGCATGGACCTAAGGCAACA 20
Sbjct 591 TGCATGGACCTAAGGCAACA 610
>MF288718.1 Human papillomavirus type 18 isolate C487627_R1_1178751_(0w),
complete genome
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Length=7857

Score = 37.
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Score = 37.4 bits (40), Expect = 2.8
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
Query 1
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Sbjct 591 TGCATGGACCTAAGGCAACA 610
>MF288717.1 Human papillomavirus type 18 isolate 1533526_N-P, complete genome
Length=7857
Score = 37.4 bits (40), Expect = 2.8
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
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Query 1
111111111111111111111
Sbjct 591 TGCATGGACCTAAGGCAACA 610
>MF288716.1 Human papillomavirus type 18 isolate C458963_R2_1584969_(48w),
complete genome
Length=7857
Score = 37.4 bits (40), Expect = 2.8
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
Query 1
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Sbjct 591 TGCATGGACCTAAGGCAACA 610
>MF288715.1 Human papillomavirus type 18 isolate C458963_R1_1201270_(0w),
complete genome
Length=7857
Score = 37.4 bits (40), Expect = 2.8
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
Query 1 TGCATGGACCTAAGGCAACA 20
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Sbjct 591 TGCATGGACCTAAGGCAACA 610

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>MF288714.1 Human papillomavirus type 18 isolate C533713_R3_1794175_(52w),
complete genome
Length=7857
Score = 37.4 bits (40), Expect = 2.8
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
          TGCATGGACCTAAGGCAACA 20
Query 1
Sbjct 591 TGCATGGACCTAAGGCAACA 610
>MF288713.1 Human papillomavirus type 18 isolate C533713_R2_1453915_(0w),
complete genome
Length=7857
Score = 37.4 bits (40), Expect = 2.8
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
Query 1 TGCATGGACCTAAGGCAACA 20
Sbjct 591 TGCATGGACCTAAGGCAACA 610
>MF288712.1 Human papillomavirus type 18 isolate C578032_R3_1777990_(53w),
complete genome
Length=7857
Score = 37.4 bits (40), Expect = 2.8
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
          TGCATGGACCTAAGGCAACA 20
Query 1
Sbjct 591 TGCATGGACCTAAGGCAACA 610
>MF288711.1 Human papillomavirus type 18 isolate C578032_R2_1441689_(0w),
complete genome
Length=7857
Score = 37.4 bits (40), Expect = 2.8
Identities = 20/20 (100%), Gaps = 0/20 (0%)
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Strand=Plus/Plus
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Query 1 TGCATGGACCTAAGGCAACA 20
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>MF288710.1 Human papillomavirus type 18 isolate C335048_R3_1978895_(63w),
complete genome
Length=7857
Score = 37.4 bits (40), Expect = 2.8
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
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>MF288709.1 Human papillomavirus type 18 isolate C412607_R2_1595957_(56w),
complete genome
Length=7857
Score = 37.4 bits (40), Expect = 2.8
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
         TGCATGGACCTAAGGCAACA 20
Query 1
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>MF288708.1 Human papillomavirus type 18 isolate C412607_R1_1241497_(0w),
complete genome
Length=7857
Score = 37.4 bits (40), Expect = 2.8
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
Query 1 TGCATGGACCTAAGGCAACA 20
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```

>MF288707.1 Human papillomavirus type 18 isolate C440384_R1_1310123_N-P,

BLAST INFORMATION complete genome Length=7842 Score = 37.4 bits (40), Expect = 2.8 Identities = 20/20 (100%), Gaps = 0/20 (0%) Strand=Plus/Plus Query 1 TGCATGGACCTAAGGCAACA 20 Sbjct 591 TGCATGGACCTAAGGCAACA 610 >MF288706.1 Human papillomavirus type 18 isolate C425046_R2_1678154_(54w), complete genome Length=7857 Score = 37.4 bits (40), Expect = 2.8 Identities = 20/20 (100%), Gaps = 0/20 (0%) Strand=Plus/Plus Query 1 TGCATGGACCTAAGGCAACA 20 Sbjct 591 TGCATGGACCTAAGGCAACA 610 >MF288705.1 Human papillomavirus type 18 isolate C425046_R1_1388626_(0w), complete genome Length=7857 Score = 37.4 bits (40), Expect = 2.8 Identities = 20/20 (100%), Gaps = 0/20 (0%) Strand=Plus/Plus Query 1 TGCATGGACCTAAGGCAACA 20 Sbjct 591 TGCATGGACCTAAGGCAACA 610 >MF288704.1 Human papillomavirus type 18 isolate C627893_R2_1657900_(46w), complete genome

Score = 37.4 bits (40), Expect = 2.8
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus

Query 1 TGCATGGACCTAAGGCAACA 20

Length=7857

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Sbjct 591 TGCATGGACCTAAGGCAACA 610
>MF288703.1 Human papillomavirus type 18 isolate C627893_R1_1382535_(0w),
complete genome
Length=7857
Score = 37.4 bits (40), Expect = 2.8
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
Query 1
          TGCATGGACCTAAGGCAACA 20
Sbjct 591 TGCATGGACCTAAGGCAACA 610
>MF288702.1 Human papillomavirus type 18 isolate C644657_R4_2179348_(154w),
complete genome
Length=7857
Score = 37.4 bits (40), Expect = 2.8
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
          TGCATGGACCTAAGGCAACA 20
Query 1
Sbjct 591 TGCATGGACCTAAGGCAACA 610
>MF288701.1 Human papillomavirus type 18 isolate C644657_R3_1822640_(103w),
complete genome
Length=7857
Score = 37.4 bits (40), Expect = 2.8
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
Query 1 TGCATGGACCTAAGGCAACA 20
Sbjct 591 TGCATGGACCTAAGGCAACA 610
>MF288700.1 Human papillomavirus type 18 isolate C644657_R2_1488163_(55w),
complete genome
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Length=7857

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BLAST INFORMATION
Score = 37.4 bits (40), Expect = 2.8
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
          TGCATGGACCTAAGGCAACA 20
Query 1
Sbjct 591 TGCATGGACCTAAGGCAACA 610
>MF288699.1 Human papillomavirus type 18 isolate C644657_R1_1107720_(0w),
complete genome
Length=7857
Score = 37.4 bits (40), Expect = 2.8
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
Query 1 TGCATGGACCTAAGGCAACA 20
Sbjct 591 TGCATGGACCTAAGGCAACA 610
>MF288698.1 Human papillomavirus type 18 isolate C335048_R2_1591561_(0w),
complete genome
Length=7857
Score = 37.4 bits (40), Expect = 2.8
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
Query 1
          TGCATGGACCTAAGGCAACA 20
Sbjct 591 TGCATGGACCTAAGGCAACA 610
>MF288697.1 Human papillomavirus type 18 isolate 1445230_R2_P_R1_FAIL, complete
genome
Length=7857
Score = 37.4 bits (40), Expect = 2.8
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
         TGCATGGACCTAAGGCAACA 20
Query 1
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Sbjct 591 TGCATGGACCTAAGGCAACA 610

```
>MF288696.1 Human papillomavirus type 18 isolate 1581449_N-P, complete genome
Length=7857
Score = 37.4 bits (40), Expect = 2.8
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
Query 1 TGCATGGACCTAAGGCAACA 20
Sbjct 591 TGCATGGACCTAAGGCAACA 610
>MF288695.1 Human papillomavirus type 18 isolate C440737_R1_1103988_N-P,
complete genome
Length=7857
Score = 37.4 bits (40), Expect = 2.8
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
          TGCATGGACCTAAGGCAACA 20
Query 1
Sbjct 591 TGCATGGACCTAAGGCAACA 610
>MF288694.1 Human papillomavirus type 18 isolate C439921_R1_1139309_N-P,
complete genome
Length=7857
Score = 37.4 bits (40), Expect = 2.8
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
Query 1
          TGCATGGACCTAAGGCAACA 20
Sbjct 591 TGCATGGACCTAAGGCAACA 610
>MF288693.1 Human papillomavirus type 18 isolate C430899_R1_1153841_N-P,
complete genome
Length=7857
Score = 37.4 bits (40), Expect = 2.8
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
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```
BLAST INFORMATION
Query 1
          TGCATGGACCTAAGGCAACA 20
Sbjct 591 TGCATGGACCTAAGGCAACA 610
>MF288692.1 Human papillomavirus type 18 isolate C353934_R1_1193001_N-P,
complete genome
Length=7857
Score = 37.4 bits (40), Expect = 2.8
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
Query 1
          TGCATGGACCTAAGGCAACA 20
Sbjct 591 TGCATGGACCTAAGGCAACA 610
>MF288691.1 Human papillomavirus type 18 isolate 1390622_R1_P_R2_FAIL, complete
genome
Length=7857
Score = 37.4 bits (40), Expect = 2.8
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
Query 1 TGCATGGACCTAAGGCAACA 20
Sbjct 591 TGCATGGACCTAAGGCAACA 610
>MF288690.1 Human papillomavirus type 18 isolate C532476_R1_1127082_N-P,
complete genome
Length=7857
Score = 37.4 bits (40), Expect = 2.8
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
          TGCATGGACCTAAGGCAACA 20
Query 1
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>MF288689.1 Human papillomavirus type 18 isolate 1508690_N-P, complete genome Length=7857

Sbjct 591 TGCATGGACCTAAGGCAACA 610

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BLAST INFORMATION
Score = 37.4 bits (40), Expect = 2.8
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
          TGCATGGACCTAAGGCAACA 20
Query 1
Sbjct 591 TGCATGGACCTAAGGCAACA 610
>MF288688.1 Human papillomavirus type 18 isolate 1115002_N-P, complete genome
Length=7857
Score = 37.4 bits (40), Expect = 2.8
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
Query 1
         TGCATGGACCTAAGGCAACA 20
Sbjct 591 TGCATGGACCTAAGGCAACA 610
>MF288687.1 Human papillomavirus type 18 isolate C406921_R1_1177005_N-P,
complete genome
Length=7857
Score = 37.4 bits (40), Expect = 2.8
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
Query 1 TGCATGGACCTAAGGCAACA 20
Sbjct 591 TGCATGGACCTAAGGCAACA 610
>MF288686.1 Human papillomavirus type 18 isolate C575918_R2_1429982_(50w),
complete genome
Length=7857
Score = 37.4 bits (40), Expect = 2.8
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
Query 1
          TGCATGGACCTAAGGCAACA 20
```

Sbjct 591 TGCATGGACCTAAGGCAACA 610

```
BLAST INFORMATION
>MF288685.1 Human papillomavirus type 18 isolate C575918_R1_1071512_(0w),
complete genome
Length=7857
Score = 37.4 bits (40), Expect = 2.8
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
Query 1 TGCATGGACCTAAGGCAACA 20
Sbjct 591 TGCATGGACCTAAGGCAACA 610
>MF288684.1 Human papillomavirus type 18 isolate 1389481_N-P, complete genome
Length=7857
Score = 37.4 bits (40), Expect = 2.8
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
Query 1 TGCATGGACCTAAGGCAACA 20
Sbjct 591 TGCATGGACCTAAGGCAACA 610
>MF288683.1 Human papillomavirus type 18 isolate 1310091_R1_P_R2_FAIL, complete
genome
Length=7857
Score = 37.4 bits (40), Expect = 2.8
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
          TGCATGGACCTAAGGCAACA 20
Query 1
Sbjct 591 TGCATGGACCTAAGGCAACA 610
>MF288682.1 Human papillomavirus type 18 isolate C385149_R2_1569977_(54w),
complete genome
Length=7857
Score = 37.4 bits (40), Expect = 2.8
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
```

Query 1

TGCATGGACCTAAGGCAACA 20

```
Sbjct 591 TGCATGGACCTAAGGCAACA 610
>MF288681.1 Human papillomavirus type 18 isolate C385149_R1_1186655_(0w),
complete genome
Length=7857
Score = 37.4 bits (40), Expect = 2.8
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
Query 1
          TGCATGGACCTAAGGCAACA 20
Sbjct 591 TGCATGGACCTAAGGCAACA 610
>MF288680.1 Human papillomavirus type 18 isolate C340709_R1_1279365_N-P,
complete genome
Length=7857
Score = 37.4 bits (40), Expect = 2.8
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
          TGCATGGACCTAAGGCAACA 20
Query 1
Sbjct 591 TGCATGGACCTAAGGCAACA 610
>MF288679.1 Human papillomavirus type 18 isolate C310065_R1_1389573_N-P,
complete genome
Length=7857
Score = 37.4 bits (40), Expect = 2.8
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
Query 1 TGCATGGACCTAAGGCAACA 20
Sbjct 591 TGCATGGACCTAAGGCAACA 610
>MF288678.1 Human papillomavirus type 18 isolate C498643_R2_1564219_(54w),
complete genome
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Length=7857

BLAST INFORMATION Score = 37.4 bits (40), Expect = 2.8Identities = 20/20 (100%), Gaps = 0/20 (0%) Strand=Plus/Plus TGCATGGACCTAAGGCAACA 20 Query 1 Sbjct 591 TGCATGGACCTAAGGCAACA 610 Database: Nucleotide collection (nt) Posted date: Nov 3, 2017 6:38 PM Number of letters in database: 162,240,808,426 Number of sequences in database: 45,172,210 Lambda K 0.634 0.408 0.912 Gapped Lambda K 0.625 0.410 0.780 Matrix: blastn matrix:2 -3 Gap Penalties: Existence: 5, Extension: 2 Number of Sequences: 45172210 Number of Hits to DB: 231098 Number of extensions: 63 Number of successful extensions: 63 Number of sequences better than 10: 0 Number of HSP's better than 10 without gapping: 0 Number of HSP's gapped: 63 Number of HSP's successfully gapped: 0 Length of query: 20 Length of database: 162240808426 Length adjustment: 17 Effective length of query: 3 Effective length of database: 161472880856 Effective search space: 484418642568 Effective search space used: 484418642568 X1: 22 (20.1 bits) X2: 33 (29.8 bits) X3: 110 (99.2 bits) S1: 28 (26.5 bits) S2: 38 (35.6 bits)

BLASTN 2.7.1+

Reference: Stephen F. Altschul, Thomas L. Madden, Alejandro A. Schaffer, Jinghui Zhang, Zheng Zhang, Webb Miller, and David J. Lipman (1997), "Gapped BLAST and PSI-BLAST: a new generation of protein database search programs", Nucleic Acids Res. 25:3389-3402.

RID: 09A3T4B0015

Database: Nucleotide collection (nt)

45,172,210 sequences; 162,240,808,426 total letters

Query= Length=21

Score E

Sequences producing significant alignments:						(Bits)	Value	
MF288726.1	Human	papillomavirus	type	18	isolate	1314304_N-P,	39.2	0.79
MF288725.1	Human	papillomavirus	type	18	isolate	C451606_R3_19	39.2	0.79
MF288724.1	Human	papillomavirus	type	18	isolate	C451606_R2_15	39.2	0.79
MF288723.1	Human	papillomavirus	type	18	isolate	C451606_R1_12	39.2	0.79
MF288722.1	Human	papillomavirus	type	18	isolate	C387957_R1_11	39.2	0.79
MF288721.1	Human	papillomavirus	type	18	isolate	1209155_R1_P	39.2	0.79
MF288720.1	Human	papillomavirus	type	18	isolate	1174820_N-P,	39.2	0.79
MF288719.1	Human	papillomavirus	type	18	isolate	C487627_R2_15	39.2	0.79
MF288718.1	Human	papillomavirus	type	18	isolate	C487627_R1_11	39.2	0.79
MF288717.1	Human	papillomavirus	type	18	isolate	1533526_N-P,	39.2	0.79
MF288716.1	Human	papillomavirus	type	18	isolate	C458963_R2_15	39.2	0.79
MF288715.1	Human	papillomavirus	type	18	isolate	C458963_R1_12	39.2	0.79
MF288714.1	Human	papillomavirus	type	18	isolate	C533713_R3_17	39.2	0.79
MF288713.1	Human	papillomavirus	type	18	isolate	C533713_R2_14	39.2	0.79
MF288712.1	Human	papillomavirus	type	18	isolate	C578032_R3_17	39.2	0.79
MF288711.1	Human	papillomavirus	type	18	isolate	C578032_R2_14	39.2	0.79
MF288710.1	Human	papillomavirus	type	18	isolate	C335048_R3_19	39.2	0.79
MF288709.1	Human	papillomavirus	type	18	isolate	C412607_R2_15	39.2	0.79
MF288708.1	Human	papillomavirus	type	18	isolate	C412607_R1_12	39.2	0.79
MF288707.1	Human	papillomavirus	type	18	isolate	C440384_R1_13	39.2	0.79
MF288706.1	Human	papillomavirus	type	18	isolate	C425046_R2_16	39.2	0.79
MF288705.1	Human	papillomavirus	type	18	isolate	C425046_R1_13	39.2	0.79
MF288704.1	Human	papillomavirus	type	18	isolate	C627893_R2_16	39.2	0.79
MF288703.1	Human	papillomavirus	type	18	isolate	C627893_R1_13	39.2	0.79
MF288702.1	Human	papillomavirus	type	18	isolate	C644657_R4_21	39.2	0.79
MF288701.1	Human	papillomavirus	type	18	isolate	C644657_R3_18	39.2	0.79
MF288700.1	Human	papillomavirus	type	18	isolate	C644657_R2_14	39.2	0.79

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BLAST INFORMATION
MF288699.1 Human papillomavirus type 18 isolate C644657_R1_11... 39.2
                                                                        0.79
MF288698.1 Human papillomavirus type 18 isolate C335048_R2_15... 39.2
                                                                        0.79
MF288697.1 Human papillomavirus type 18 isolate 1445230 R2 P ... 39.2
                                                                        0.79
MF288696.1 Human papillomavirus type 18 isolate 1581449_N-P, ... 39.2
                                                                        0.79
MF288695.1 Human papillomavirus type 18 isolate C440737_R1_11... 39.2
                                                                       0.79
MF288694.1 Human papillomavirus type 18 isolate C439921 R1 11... 39.2
                                                                        0.79
MF288693.1 Human papillomavirus type 18 isolate C430899_R1_11... 39.2
                                                                        0.79
MF288692.1 Human papillomavirus type 18 isolate C353934_R1_11... 39.2
                                                                       0.79
MF288691.1 Human papillomavirus type 18 isolate 1390622_R1_P_... 39.2
                                                                        0.79
MF288690.1 Human papillomavirus type 18 isolate C532476_R1_11... 39.2
                                                                        0.79
MF288689.1 Human papillomavirus type 18 isolate 1508690_N-P, ... 39.2
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MF288688.1 Human papillomavirus type 18 isolate 1115002_N-P, ... 39.2
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MF288687.1 Human papillomavirus type 18 isolate C406921_R1_11... 39.2
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MF288686.1 Human papillomavirus type 18 isolate C575918_R2_14... 39.2
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MF288685.1 Human papillomavirus type 18 isolate C575918_R1_10... 39.2
                                                                        0.79
MF288684.1 Human papillomavirus type 18 isolate 1389481_N-P, ... 39.2
                                                                        0.79
MF288683.1 Human papillomavirus type 18 isolate 1310091_R1_P_... 39.2
                                                                       0.79
MF288682.1 Human papillomavirus type 18 isolate C385149_R2_15... 39.2
                                                                        0.79
MF288681.1 Human papillomavirus type 18 isolate C385149_R1_11... 39.2
                                                                        0.79
MF288680.1 Human papillomavirus type 18 isolate C340709_R1_12... 39.2
                                                                       0.79
MF288679.1 Human papillomavirus type 18 isolate C310065_R1_13... 39.2
                                                                        0.79
MF288678.1 Human papillomavirus type 18 isolate C498643_R2_15... 39.2
                                                                        0.79
MF288677.1 Human papillomavirus type 18 isolate C498643_R1_11... 39.2
                                                                        0.79
ALIGNMENTS
>MF288726.1 Human papillomavirus type 18 isolate 1314304_N-P, complete genome
Length=7857
Score = 39.2 \text{ bits } (42), Expect = 0.79
Identities = 21/21 (100%), Gaps = 0/21 (0%)
Strand=Plus/Plus
Query 1 TGCATGGACCTAAGGCAACAT 21
Sbjct 591 TGCATGGACCTAAGGCAACAT 611
```

>MF288725.1 Human papillomavirus type 18 isolate C451606 R3 1915584 (100w), complete genome Length=7857

Score = 39.2 bits (42), Expect = 0.79Identities = 21/21 (100%), Gaps = 0/21 (0%) Strand=Plus/Plus

TGCATGGACCTAAGGCAACAT 21

```
Sbjct 591 TGCATGGACCTAAGGCAACAT 611
```

```
>MF288724.1 Human papillomavirus type 18 isolate C451606_R2_1584906_(52w),
complete genome
Length=7857
Score = 39.2 \text{ bits } (42), Expect = 0.79
Identities = 21/21 (100%), Gaps = 0/21 (0%)
Strand=Plus/Plus
Query 1 TGCATGGACCTAAGGCAACAT 21
Sbjct 591 TGCATGGACCTAAGGCAACAT 611
>MF288723.1 Human papillomavirus type 18 isolate C451606_R1_1201181_(0w),
complete genome
Length=7857
Score = 39.2 \text{ bits } (42), Expect = 0.79
Identities = 21/21 (100%), Gaps = 0/21 (0%)
Strand=Plus/Plus
          TGCATGGACCTAAGGCAACAT 21
Query 1
Sbjct 591 TGCATGGACCTAAGGCAACAT 611
>MF288722.1 Human papillomavirus type 18 isolate C387957_R1_1124102_N-P,
complete genome
Length=7857
Score = 39.2 bits (42), Expect = 0.79
Identities = 21/21 (100%), Gaps = 0/21 (0%)
Strand=Plus/Plus
Query 1
          TGCATGGACCTAAGGCAACAT 21
Sbjct 591 TGCATGGACCTAAGGCAACAT 611
>MF288721.1 Human papillomavirus type 18 isolate 1209155_R1_P_R2_FAIL, complete
genome
Length=7857
Score = 39.2 bits (42), Expect = 0.79
```

```
BLAST INFORMATION
Identities = 21/21 (100%), Gaps = 0/21 (0%)
Strand=Plus/Plus
          TGCATGGACCTAAGGCAACAT 21
Query 1
Sbjct 591 TGCATGGACCTAAGGCAACAT 611
>MF288720.1 Human papillomavirus type 18 isolate 1174820_N-P, complete genome
Length=7857
Score = 39.2 \text{ bits } (42), Expect = 0.79
Identities = 21/21 (100%), Gaps = 0/21 (0%)
Strand=Plus/Plus
          TGCATGGACCTAAGGCAACAT 21
Query 1
Sbjct 591 TGCATGGACCTAAGGCAACAT 611
>MF288719.1 Human papillomavirus type 18 isolate C487627_R2_1553725_(52w),
complete genome
Length=7857
Score = 39.2 bits (42), Expect = 0.79
Identities = 21/21 (100%), Gaps = 0/21 (0%)
Strand=Plus/Plus
Query 1
          TGCATGGACCTAAGGCAACAT 21
Sbjct 591 TGCATGGACCTAAGGCAACAT 611
>MF288718.1 Human papillomavirus type 18 isolate C487627_R1_1178751_(0w),
complete genome
Length=7857
Score = 39.2 \text{ bits } (42), Expect = 0.79
Identities = 21/21 (100%), Gaps = 0/21 (0%)
Strand=Plus/Plus
Query 1 TGCATGGACCTAAGGCAACAT 21
Sbjct 591 TGCATGGACCTAAGGCAACAT 611
```

>MF288717.1 Human papillomavirus type 18 isolate 1533526_N-P, complete genome

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Length=7857
Score = 39.2 \text{ bits } (42), Expect = 0.79
Identities = 21/21 (100%), Gaps = 0/21 (0%)
Strand=Plus/Plus
Query 1
           TGCATGGACCTAAGGCAACAT 21
Sbjct 591 TGCATGGACCTAAGGCAACAT 611
>MF288716.1 Human papillomavirus type 18 isolate C458963_R2_1584969_(48w),
complete genome
Length=7857
Score = 39.2 \text{ bits } (42), Expect = 0.79
Identities = 21/21 (100%), Gaps = 0/21 (0%)
Strand=Plus/Plus
           TGCATGGACCTAAGGCAACAT 21
Query 1
Sbjct 591 TGCATGGACCTAAGGCAACAT 611
>MF288715.1 Human papillomavirus type 18 isolate C458963_R1_1201270_(0w),
complete genome
Length=7857
Score = 39.2 \text{ bits } (42), Expect = 0.79
Identities = 21/21 (100%), Gaps = 0/21 (0%)
Strand=Plus/Plus
Query 1 TGCATGGACCTAAGGCAACAT 21
Sbjct 591 TGCATGGACCTAAGGCAACAT 611
>MF288714.1 Human papillomavirus type 18 isolate C533713_R3_1794175_(52w),
complete genome
Length=7857
Score = 39.2 \text{ bits } (42), Expect = 0.79
Identities = 21/21 (100%), Gaps = 0/21 (0%)
Strand=Plus/Plus
Query 1
           TGCATGGACCTAAGGCAACAT 21
```

```
Sbjct 591 TGCATGGACCTAAGGCAACAT 611
```

```
>MF288713.1 Human papillomavirus type 18 isolate C533713_R2_1453915_(0w),
complete genome
Length=7857
Score = 39.2 \text{ bits } (42), Expect = 0.79
Identities = 21/21 (100%), Gaps = 0/21 (0%)
Strand=Plus/Plus
Query 1 TGCATGGACCTAAGGCAACAT 21
Sbjct 591 TGCATGGACCTAAGGCAACAT 611
>MF288712.1 Human papillomavirus type 18 isolate C578032_R3_1777990_(53w),
complete genome
Length=7857
Score = 39.2 \text{ bits } (42), Expect = 0.79
Identities = 21/21 (100%), Gaps = 0/21 (0%)
Strand=Plus/Plus
          TGCATGGACCTAAGGCAACAT 21
Query 1
Sbjct 591 TGCATGGACCTAAGGCAACAT 611
>MF288711.1 Human papillomavirus type 18 isolate C578032_R2_1441689_(0w),
complete genome
Length=7857
Score = 39.2 \text{ bits } (42), Expect = 0.79
Identities = 21/21 (100%), Gaps = 0/21 (0%)
Strand=Plus/Plus
Query 1
          TGCATGGACCTAAGGCAACAT 21
Sbjct 591 TGCATGGACCTAAGGCAACAT 611
>MF288710.1 Human papillomavirus type 18 isolate C335048_R3_1978895_(63w),
complete genome
Length=7857
Score = 39.2 bits (42), Expect = 0.79
```

```
BLAST INFORMATION
Identities = 21/21 (100%), Gaps = 0/21 (0%)
Strand=Plus/Plus
          TGCATGGACCTAAGGCAACAT 21
Query 1
Sbjct 591 TGCATGGACCTAAGGCAACAT 611
>MF288709.1 Human papillomavirus type 18 isolate C412607_R2_1595957_(56w),
complete genome
Length=7857
Score = 39.2 \text{ bits } (42), Expect = 0.79
Identities = 21/21 (100%), Gaps = 0/21 (0%)
Strand=Plus/Plus
Query 1
          TGCATGGACCTAAGGCAACAT 21
Sbjct 591 TGCATGGACCTAAGGCAACAT 611
>MF288708.1 Human papillomavirus type 18 isolate C412607_R1_1241497_(0w),
complete genome
Length=7857
Score = 39.2 \text{ bits } (42), Expect = 0.79
Identities = 21/21 (100%), Gaps = 0/21 (0%)
Strand=Plus/Plus
Query 1 TGCATGGACCTAAGGCAACAT 21
Sbjct 591 TGCATGGACCTAAGGCAACAT 611
>MF288707.1 Human papillomavirus type 18 isolate C440384_R1_1310123_N-P,
complete genome
Length=7842
Score = 39.2 \text{ bits } (42), Expect = 0.79
Identities = 21/21 (100%), Gaps = 0/21 (0%)
Strand=Plus/Plus
          TGCATGGACCTAAGGCAACAT 21
Query 1
```

Sbjct 591 TGCATGGACCTAAGGCAACAT 611

BLAST INFORMATION >MF288706.1 Human papillomavirus type 18 isolate C425046_R2_1678154_(54w), complete genome Length=7857 Score = 39.2 bits (42), Expect = 0.79Identities = 21/21 (100%), Gaps = 0/21 (0%) Strand=Plus/Plus Query 1 TGCATGGACCTAAGGCAACAT 21 Sbjct 591 TGCATGGACCTAAGGCAACAT 611 >MF288705.1 Human papillomavirus type 18 isolate C425046_R1_1388626_(0w), complete genome Length=7857 Score = 39.2 bits (42), Expect = 0.79Identities = 21/21 (100%), Gaps = 0/21 (0%) Strand=Plus/Plus TGCATGGACCTAAGGCAACAT 21 Query 1 Sbjct 591 TGCATGGACCTAAGGCAACAT 611 >MF288704.1 Human papillomavirus type 18 isolate C627893_R2_1657900_(46w), complete genome Length=7857 Score = 39.2 bits (42), Expect = 0.79Identities = 21/21 (100%), Gaps = 0/21 (0%) Strand=Plus/Plus Query 1 TGCATGGACCTAAGGCAACAT 21 Sbjct 591 TGCATGGACCTAAGGCAACAT 611 >MF288703.1 Human papillomavirus type 18 isolate C627893_R1_1382535_(0w), complete genome

Length=7857

Strand=Plus/Plus

Score = 39.2 bits (42), Expect = 0.79

Identities = 21/21 (100%), Gaps = 0/21 (0%)

```
BLAST INFORMATION
Query 1
          TGCATGGACCTAAGGCAACAT 21
Sbjct 591 TGCATGGACCTAAGGCAACAT 611
>MF288702.1 Human papillomavirus type 18 isolate C644657_R4_2179348_(154w),
complete genome
Length=7857
Score = 39.2 \text{ bits } (42), Expect = 0.79
Identities = 21/21 (100%), Gaps = 0/21 (0%)
Strand=Plus/Plus
Query 1
          TGCATGGACCTAAGGCAACAT 21
Sbjct 591 TGCATGGACCTAAGGCAACAT 611
>MF288701.1 Human papillomavirus type 18 isolate C644657_R3_1822640_(103w),
complete genome
Length=7857
Score = 39.2 \text{ bits } (42), Expect = 0.79
Identities = 21/21 (100%), Gaps = 0/21 (0%)
Strand=Plus/Plus
Query 1 TGCATGGACCTAAGGCAACAT 21
Sbjct 591 TGCATGGACCTAAGGCAACAT 611
>MF288700.1 Human papillomavirus type 18 isolate C644657_R2_1488163_(55w),
complete genome
Length=7857
Score = 39.2 bits (42), Expect = 0.79
Identities = 21/21 (100%), Gaps = 0/21 (0%)
Strand=Plus/Plus
          TGCATGGACCTAAGGCAACAT 21
Query 1
Sbjct 591 TGCATGGACCTAAGGCAACAT 611
>MF288699.1 Human papillomavirus type 18 isolate C644657_R1_1107720_(0w),
```

complete genome Length=7857

Score = 39.2 bits (42), Expect = 0.79

```
Identities = 21/21 (100%), Gaps = 0/21 (0%)
Strand=Plus/Plus
Query 1 TGCATGGACCTAAGGCAACAT 21
Sbjct 591 TGCATGGACCTAAGGCAACAT 611
>MF288698.1 Human papillomavirus type 18 isolate C335048_R2_1591561_(0w),
complete genome
Length=7857
Score = 39.2 \text{ bits } (42), Expect = 0.79
Identities = 21/21 (100%), Gaps = 0/21 (0%)
Strand=Plus/Plus
          TGCATGGACCTAAGGCAACAT 21
Query 1
Sbjct 591 TGCATGGACCTAAGGCAACAT 611
>MF288697.1 Human papillomavirus type 18 isolate 1445230_R2_P_R1_FAIL, complete
genome
Length=7857
Score = 39.2 \text{ bits } (42), Expect = 0.79
Identities = 21/21 (100%), Gaps = 0/21 (0%)
Strand=Plus/Plus
Query 1
          TGCATGGACCTAAGGCAACAT 21
Sbjct 591 TGCATGGACCTAAGGCAACAT 611
>MF288696.1 Human papillomavirus type 18 isolate 1581449_N-P, complete genome
Length=7857
Score = 39.2 \text{ bits } (42), Expect = 0.79
Identities = 21/21 (100%), Gaps = 0/21 (0%)
Strand=Plus/Plus
Query 1
          TGCATGGACCTAAGGCAACAT 21
Sbjct 591 TGCATGGACCTAAGGCAACAT 611
```

```
complete genome
Length=7857
Score = 39.2 \text{ bits } (42), Expect = 0.79
Identities = 21/21 (100%), Gaps = 0/21 (0%)
Strand=Plus/Plus
Query 1
           TGCATGGACCTAAGGCAACAT 21
Sbjct 591 TGCATGGACCTAAGGCAACAT 611
>MF288694.1 Human papillomavirus type 18 isolate C439921_R1_1139309_N-P,
complete genome
Length=7857
Score = 39.2 \text{ bits } (42), Expect = 0.79
Identities = 21/21 (100%), Gaps = 0/21 (0%)
Strand=Plus/Plus
           TGCATGGACCTAAGGCAACAT 21
Query 1
Sbjct 591 TGCATGGACCTAAGGCAACAT 611
>MF288693.1 Human papillomavirus type 18 isolate C430899_R1_1153841_N-P,
complete genome
Length=7857
Score = 39.2 \text{ bits } (42), Expect = 0.79
Identities = 21/21 (100%), Gaps = 0/21 (0%)
Strand=Plus/Plus
Query 1 TGCATGGACCTAAGGCAACAT 21
Sbjct 591 TGCATGGACCTAAGGCAACAT 611
>MF288692.1 Human papillomavirus type 18 isolate C353934_R1_1193001_N-P,
complete genome
Length=7857
Score = 39.2 \text{ bits } (42), Expect = 0.79
Identities = 21/21 (100%), Gaps = 0/21 (0%)
Strand=Plus/Plus
```

>MF288695.1 Human papillomavirus type 18 isolate C440737_R1_1103988_N-P,

Length=7857

```
TGCATGGACCTAAGGCAACAT 21
Query 1
Sbjct 591 TGCATGGACCTAAGGCAACAT 611
>MF288691.1 Human papillomavirus type 18 isolate 1390622_R1_P_R2_FAIL, complete
genome
Length=7857
Score = 39.2 \text{ bits } (42), Expect = 0.79
Identities = 21/21 (100%), Gaps = 0/21 (0%)
Strand=Plus/Plus
Query 1 TGCATGGACCTAAGGCAACAT 21
Sbjct 591 TGCATGGACCTAAGGCAACAT 611
>MF288690.1 Human papillomavirus type 18 isolate C532476_R1_1127082_N-P,
complete genome
Length=7857
Score = 39.2 \text{ bits } (42), Expect = 0.79
Identities = 21/21 (100%), Gaps = 0/21 (0%)
Strand=Plus/Plus
          TGCATGGACCTAAGGCAACAT 21
Query 1
Sbjct 591 TGCATGGACCTAAGGCAACAT 611
>MF288689.1 Human papillomavirus type 18 isolate 1508690_N-P, complete genome
Length=7857
Score = 39.2 \text{ bits } (42), Expect = 0.79
Identities = 21/21 (100%), Gaps = 0/21 (0%)
Strand=Plus/Plus
          TGCATGGACCTAAGGCAACAT 21
Query 1
Sbjct 591 TGCATGGACCTAAGGCAACAT 611
>MF288688.1 Human papillomavirus type 18 isolate 1115002_N-P, complete genome
```

```
BLAST INFORMATION
Score = 39.2 bits (42), Expect = 0.79
Identities = 21/21 (100%), Gaps = 0/21 (0%)
Strand=Plus/Plus
          TGCATGGACCTAAGGCAACAT 21
Query 1
Sbjct 591 TGCATGGACCTAAGGCAACAT 611
>MF288687.1 Human papillomavirus type 18 isolate C406921_R1_1177005_N-P,
complete genome
Length=7857
Score = 39.2 \text{ bits } (42), Expect = 0.79
Identities = 21/21 (100%), Gaps = 0/21 (0%)
Strand=Plus/Plus
Query 1 TGCATGGACCTAAGGCAACAT 21
Sbjct 591 TGCATGGACCTAAGGCAACAT 611
>MF288686.1 Human papillomavirus type 18 isolate C575918_R2_1429982_(50w),
complete genome
Length=7857
Score = 39.2 bits (42), Expect = 0.79
Identities = 21/21 (100%), Gaps = 0/21 (0%)
Strand=Plus/Plus
Query 1
          TGCATGGACCTAAGGCAACAT 21
Sbjct 591 TGCATGGACCTAAGGCAACAT 611
>MF288685.1 Human papillomavirus type 18 isolate C575918_R1_1071512_(0w),
complete genome
Length=7857
Score = 39.2 \text{ bits } (42), Expect = 0.79
Identities = 21/21 (100%), Gaps = 0/21 (0%)
Strand=Plus/Plus
         TGCATGGACCTAAGGCAACAT 21
Query 1
```

Sbjct 591 TGCATGGACCTAAGGCAACAT 611

```
>MF288684.1 Human papillomavirus type 18 isolate 1389481_N-P, complete genome
Length=7857
Score = 39.2 \text{ bits } (42), Expect = 0.79
Identities = 21/21 (100%), Gaps = 0/21 (0%)
Strand=Plus/Plus
Query 1 TGCATGGACCTAAGGCAACAT 21
Sbjct 591 TGCATGGACCTAAGGCAACAT 611
>MF288683.1 Human papillomavirus type 18 isolate 1310091_R1_P_R2_FAIL, complete
genome
Length=7857
Score = 39.2 \text{ bits } (42), Expect = 0.79
Identities = 21/21 (100%), Gaps = 0/21 (0%)
Strand=Plus/Plus
           TGCATGGACCTAAGGCAACAT 21
Query 1
Sbjct 591 TGCATGGACCTAAGGCAACAT 611
>MF288682.1 Human papillomavirus type 18 isolate C385149_R2_1569977_(54w),
complete genome
Length=7857
Score = 39.2 \text{ bits } (42), Expect = 0.79
Identities = 21/21 (100%), Gaps = 0/21 (0%)
Strand=Plus/Plus
Query 1
          TGCATGGACCTAAGGCAACAT 21
Sbjct 591 TGCATGGACCTAAGGCAACAT 611
>MF288681.1 Human papillomavirus type 18 isolate C385149_R1_1186655_(0w),
complete genome
Length=7857
Score = 39.2 \text{ bits } (42), Expect = 0.79
Identities = 21/21 (100%), Gaps = 0/21 (0%)
Strand=Plus/Plus
```

```
BLAST INFORMATION
          TGCATGGACCTAAGGCAACAT 21
Query 1
Sbjct 591 TGCATGGACCTAAGGCAACAT 611
>MF288680.1 Human papillomavirus type 18 isolate C340709_R1_1279365_N-P,
complete genome
Length=7857
Score = 39.2 \text{ bits } (42), Expect = 0.79
Identities = 21/21 (100%), Gaps = 0/21 (0%)
Strand=Plus/Plus
Query 1
          TGCATGGACCTAAGGCAACAT 21
Sbjct 591 TGCATGGACCTAAGGCAACAT 611
>MF288679.1 Human papillomavirus type 18 isolate C310065_R1_1389573_N-P,
complete genome
Length=7857
Score = 39.2 \text{ bits } (42), Expect = 0.79
Identities = 21/21 (100%), Gaps = 0/21 (0%)
Strand=Plus/Plus
Query 1 TGCATGGACCTAAGGCAACAT 21
Sbjct 591 TGCATGGACCTAAGGCAACAT 611
>MF288678.1 Human papillomavirus type 18 isolate C498643_R2_1564219_(54w),
complete genome
Length=7857
Score = 39.2 bits (42), Expect = 0.79
Identities = 21/21 (100%), Gaps = 0/21 (0%)
Strand=Plus/Plus
          TGCATGGACCTAAGGCAACAT 21
Query 1
Sbjct 591 TGCATGGACCTAAGGCAACAT 611
```

>MF288677.1 Human papillomavirus type 18 isolate C498643_R1_1192121_(0w), complete genome Length=7857

Score = 39.2 bits (42), Expect = 0.79

```
Identities = 21/21 (100%), Gaps = 0/21 (0%)
Strand=Plus/Plus
Query 1 TGCATGGACCTAAGGCAACAT 21
Sbjct 591 TGCATGGACCTAAGGCAACAT 611
Database: Nucleotide collection (nt)
Posted date: Nov 3, 2017 6:38 PM
Number of letters in database: 162,240,808,426
Number of sequences in database: 45,172,210
Lambda
          K
                    Η
       0.408 0.912
0.634
Gapped
Lambda
          K
                    Η
0.625
       0.410
                0.780
Matrix: blastn matrix:2 -3
Gap Penalties: Existence: 5, Extension: 2
Number of Sequences: 45172210
Number of Hits to DB: 314439
Number of extensions: 158
Number of successful extensions: 158
Number of sequences better than 10: 11
Number of HSP's better than 10 without gapping: 0
Number of HSP's gapped: 158
Number of HSP's successfully gapped: 11
Length of query: 21
Length of database: 162240808426
Length adjustment: 18
Effective length of query: 3
Effective length of database: 161427708646
Effective search space: 484283125938
Effective search space used: 484283125938
A: 0
X1: 22 (20.1 bits)
X2: 33 (29.8 bits)
X3: 110 (99.2 bits)
S1: 28 (26.5 bits)
S2: 38 (35.6 bits)
```