### CLUSTAL 2.1 multiple sequence alignment

7/0//7/010 1	
KC470218.1	ATGGCGCGCTTTGAGGATCCAACACGGCGACCCTACAAGCTACCTGATCT
KC470214.1	ATGGCGCGCTTTGAGGATCCAACACGGCGACCCTACAAGCTACCTGATCT
KC470215.1	ATGGCGCGCTTTGAGGATCCAACACGGCGACCCTACAAGCTACCTGATCT
KC470216.1	ATGGCGCGCTTTGAGGATCCAACACGGCGACCCTACAAGCTACCTGATCT
KC470217.1	ATGGCGCGCTTTGAGGATCCAACACGGCGACCCTACAAGCTACCTGATCT
KC470219.1	ATGGCGCGCTTTGAGGATCCAACACGGCGACCCTACAAGCTACCTGATCT
KC470220.1	ATGGCGCGCTTTGAGGATCCAACACGGCGACCCTACAAGCTACCTGATCT
KC470221.1	ATGGCGCGCTTTGAGGATCCAACACGGCGACCCTACAAGCTACCTGATCT
KC470222.1	ATGGCGCGCTTTGAGGATCCAACACGGCGACCCTACAAGCTACCTGATCT
KC470223.1	ATGGCGCGCTTTGAGGATCCAACACGGCGACCCTACAAGCTACCTGATCT
KC470225.1	ATGGCGCGCTTTGAGGATCCAACACGGCGACCCTACAAGCTACCTGATCT
KC470226.1	ATGGCGCGCTTTGAGGATCCAACACGGCGACCCTACAAGCTACCTGATCT
KC470227.1	$\tt ATGGCGCGCTTTGAGGATCCAACACGGCGACCCTACAAGCTACCTGATCT$
KC470228.1	$\tt ATGGCGCGCTTTGAGGATCCAACACGGCGACCCTACAAGCTACCTGATCT$
KC470230.1	$\tt ATGGCGCGCTTTGAGGATCCAACACGGCGACCCTACAAGCTACCTGATCT$
KC470229.1	$\tt ATGGCGCGCTTTGAGGATCCAACACGGCGACCCTACAAGCTACCTGATCT$
KC470224.1	$\tt ATGGCGCGCTTTGAGGATCCAACACGGCGACCCTACAAGCTACCTGATCT$
KX514433.1	ATGGCGCGCTTTGAGGATCCAACACGGCGACCCTACAAGCTACCTGATCT
KU298886.1	ATGGCGCGCTTTGAGGATCCAACACGGCGACCCTACAAGCTACCTGATCT
KC470213.1	ATGGCGCGCTTTGAGGATCCAACACGGCGACCCTACAAGCTACCTGATCT
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KC470218.1	GTGCACGGAACTGAACACTTCACTGCAAGACATAGAAATAACCTGTGTAT
KC470214.1	GTGCACGGAACTGAACACTTCACTGCAAGACATAGAAATAACCTGTGTAT
KC470215.1	GTGCACGGAACTGAACACTTCACTGCAAGACATAGAAATAACCTGTGTAT
KC470216.1	GTGCACGGAACTGAACACTTCACTGCAAGACATAGAAATAACCTGTGTAT
KC470217.1	GTGCACGGAACTGAACACTTCACTGCAAGACATAGAAATAACCTGTGTAT
KC470219.1	GTGCACGGAACTGAACACTTCACTGCAAGACATAGAAATAACCTGTGTAT
KC470219.1 KC470220.1	GTGCACGGAACTGAACACTTCACTGCAAGACATAGAAATAACCTGTGTAT GTGCACGGAACTGAACACTTCACTGCAAGACATAGAAATAACCTGTGTAT
KC470220.1	GTGCACGGAACTGAACACTTCACTGCAAGACATAGAAATAACCTGTGTAT
KC470220.1 KC470221.1	GTGCACGGAACTGAACACTTCACTGCAAGACATAGAAATAACCTGTGTAT GTGCACGGAACTGAACACTTCACTGCAAGACATAGAAATAACCTGTGTAT
KC470220.1 KC470221.1 KC470222.1	GTGCACGGAACTGAACACTTCACTGCAAGACATAGAAATAACCTGTGTAT GTGCACGGAACTGAACACTTCACTGCAAGACATAGAAATAACCTGTGTAT GTGCACGGAACTGAACACTTCACTGCAAGACATAGAAATAACCTGTGTAT
KC470220.1 KC470221.1 KC470222.1 KC470223.1	GTGCACGGAACTGAACACTTCACTGCAAGACATAGAAATAACCTGTGTAT GTGCACGGAACTGAACACTTCACTGCAAGACATAGAAATAACCTGTGTAT GTGCACGGAACTGAACACTTCACTGCAAGACATAGAAATAACCTGTGTAT GTGCACGGAACTGAACACTTCACTGCAAGACATAGAAATAACCTGTGTAT
KC470220.1 KC470221.1 KC470222.1 KC470223.1 KC470225.1	GTGCACGGAACTGAACACTTCACTGCAAGACATAGAAATAACCTGTGTAT GTGCACGGAACTGAACACTTCACTGCAAGACATAGAAATAACCTGTGTAT GTGCACGGAACTGAACACTTCACTGCAAGACATAGAAATAACCTGTGTAT GTGCACGGAACTGAACACTTCACTGCAAGACATAGAAATAACCTGTGTAT GTGCACGGAACTGAACACTTCACTGCAAGACATAGAAATAACCTGTGTAT
KC470220.1 KC470221.1 KC470222.1 KC470223.1 KC470225.1 KC470226.1	GTGCACGGAACTGAACACTTCACTGCAAGACATAGAAATAACCTGTGTAT GTGCACGGAACTGAACACTTCACTGCAAGACATAGAAATAACCTGTGTAT GTGCACGGAACTGAACACTTCACTGCAAGACATAGAAATAACCTGTGTAT GTGCACGGAACTGAACACTTCACTGCAAGACATAGAAATAACCTGTGTAT GTGCACGGAACTGAACACTTCACTGCAAGACATAGAAATAACCTGTGTAT GTGCACGGAACTGAACACTTCACTGCAAGACATAGAAATAACCTGTGTAT
KC470220.1 KC470221.1 KC470222.1 KC470223.1 KC470225.1 KC470226.1 KC470227.1	GTGCACGGAACTGAACACTTCACTGCAAGACATAGAAATAACCTGTGTAT GTGCACGGAACTGAACACTTCACTGCAAGACATAGAAATAACCTGTGTAT GTGCACGGAACTGAACACTTCACTGCAAGACATAGAAATAACCTGTGTAT GTGCACGGAACTGAACACTTCACTGCAAGACATAGAAATAACCTGTGTAT GTGCACGGAACTGAACACTTCACTGCAAGACATAGAAATAACCTGTGTAT GTGCACGGAACTGAACACTTCACTGCAAGACATAGAAATAACCTGTGTAT GTGCACGGAACTGAACACTTCACTGCAAGACATAGAAATAACCTGTGTAT
KC470220.1 KC470221.1 KC470222.1 KC470223.1 KC470225.1 KC470226.1 KC470227.1 KC470228.1	GTGCACGGAACTGAACACTTCACTGCAAGACATAGAAATAACCTGTGTAT GTGCACGGAACTGAACACTTCACTGCAAGACATAGAAATAACCTGTGTAT GTGCACGGAACTGAACACTTCACTGCAAGACATAGAAATAACCTGTGTAT GTGCACGGAACTGAACACTTCACTGCAAGACATAGAAATAACCTGTGTAT GTGCACGGAACTGAACACTTCACTGCAAGACATAGAAATAACCTGTGTAT GTGCACGGAACTGAACACTTCACTGCAAGACATAGAAATAACCTGTGTAT GTGCACGGAACTGAACACTTCACTGCAAGACATAGAAATAACCTGTGTAT GTGCACGGAACTGAACACTTCACTGCAAGACATAGAAATAACCTGTGTAT
KC470220.1 KC470221.1 KC470222.1 KC470223.1 KC470225.1 KC470226.1 KC470227.1 KC470228.1 KC470230.1	GTGCACGGAACTGAACACTTCACTGCAAGACATAGAAATAACCTGTGTAT GTGCACGGAACTGAACACTTCACTGCAAGACATAGAAATAACCTGTGTAT GTGCACGGAACTGAACACTTCACTGCAAGACATAGAAATAACCTGTGTAT GTGCACGGAACTGAACACTTCACTGCAAGACATAGAAATAACCTGTGTAT GTGCACGGAACTGAACACTTCACTGCAAGACATAGAAATAACCTGTGTAT GTGCACGGAACTGAACACTTCACTGCAAGACATAGAAATAACCTGTGTAT GTGCACGGAACTGAACACTTCACTGCAAGACATAGAAATAACCTGTGTAT GTGCACGGAACTGAACACTTCACTGCAAGACATAGAAATAACCTGTGTAT GTGCACGGAACTGAACACTTCACTGCAAGACATAGAAATAACCTGTGTAT
KC470220.1 KC470221.1 KC470222.1 KC470223.1 KC470225.1 KC470226.1 KC470227.1 KC470228.1 KC470230.1 KC470229.1	GTGCACGGAACTGAACACTTCACTGCAAGACATAGAAATAACCTGTGTAT
KC470220.1 KC470221.1 KC470222.1 KC470223.1 KC470225.1 KC470226.1 KC470227.1 KC470228.1 KC470230.1 KC470229.1 KC470224.1 KX514433.1	GTGCACGGAACTGAACACTTCACTGCAAGACATAGAAATAACCTGTGTAT
KC470220.1 KC470221.1 KC470222.1 KC470223.1 KC470225.1 KC470226.1 KC470227.1 KC470228.1 KC470230.1 KC470229.1 KC470224.1 KX514433.1 KU298886.1	GTGCACGGAACTGAACACTTCACTGCAAGACATAGAAATAACCTGTGTAT
KC470220.1 KC470221.1 KC470222.1 KC470223.1 KC470225.1 KC470226.1 KC470227.1 KC470228.1 KC470230.1 KC470229.1 KC470224.1 KX514433.1	GTGCACGGAACTGAACACTTCACTGCAAGACATAGAAATAACCTGTGTAT
KC470220.1 KC470221.1 KC470222.1 KC470223.1 KC470225.1 KC470226.1 KC470227.1 KC470228.1 KC470230.1 KC470229.1 KC470224.1 KX514433.1 KU298886.1	GTGCACGGAACTGAACACTTCACTGCAAGACATAGAAATAACCTGTGTAT

CLUSTAL	2.1	multiple	sequence	alignment
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KC470214.1	ATTGCAAGACAGTATTGGAACTTACAGAGGTATTTGAATTTGCATTCAAA
KC470215.1	ATTGCAAGACAGTATTGGAACTTACAGAGGTATTTGAATTTGCATTCAAA
KC470216.1	ATTGCAAGACAGTATTGGAACTTACAGAGGTATTTGAATTTGCATTCAAA
KC470217.1	ATTGCAAGACAGTATTGGAACTTACAGAGGTATTTGAATTTGCATTCAAA
KC470219.1	ATTGCAAGACAGTATTGGAACTTACAGAGGTATTTGAATTTGCATTCAAA
KC470220.1	ATTGCAAGACAGTATTGGAACTTACAGAGGTATTTGAATTTGCATTCAAA
KC470221.1	ATTGCAAGACAGTATTGGAACTTACAGAGGTATTTGAATTTGCATTCAAA
KC470222.1	ATTGCAAGACAGTATTGGAACTTACAGAGGTATTTGAATTTGCATTCAAA
KC470223.1	ATTGCAAGACAGTATTGGAACTTACAGAGGTATTTGAATTTGCATTCAAA
KC470225.1	ATTGCAAGACAGTATTGGAACTTACAGAGGTATTTGAATTTGCATTCAAA
KC470226.1	ATTGCAAGACAGTATTGGAACTTACAGAGGTATTTGAATTTGCATTCAAA
KC470227.1	ATTGCAAGACAGTATTGGAACTTACAGAGGTATTTGAATTTGCATTCAAA
KC470228.1	ATTGCAAGACAGTATTGGAACTTACAGAGGTATTTGAATTTGCATTCAAA
KC470230.1	ATTGCAAGACAGTATTGGAACTTACAGAGGTATTTGAATTTGCATTCAAA
KC470229.1	ATTGCAAGACAGTATTGGAACTTACAGAGGTATTTGAATTTGCATTCAAA
KC470224.1	ATTGCAAGACAGTATTGGAACTTACAGAGGTATTTGAATTTGCATTCAAA
KX514433.1	ATTGCAAGACAGTATTGGAACTTACAGAGGTATTTGAATTTGCATTTAAA
KU298886.1	ATTGCAAGACAGTATTGGAACTTACAGAGGTATTTGAATTTGCATTTAAA
KC470213.1	ATTGCAAGACAGTATTGGAACTTACAGAGGTATTTGAATTTGCATTTAAA
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KC470218.1	GATTTATTTGTAGTGTATAGAGACAGTATACCGCATGCTGCATGCCATAA
KC470214.1	GATTTATTTGTAGTGTATAGAGACAGTATACCGCATGCTGCATGCCATAA
KC470215.1	GATTTATTTGTAGTGTATAGAGACAGTATACCGCATGCTGCATGCCATAA
KC470216.1	GATTTATTTGTAGTGTATAGAGACAGTATACCGCATGCTGCATGCCATAA
KC470217.1	GATTTATTTGTAGTGTATAGAGACAGTATACCGCATGCTGCATGCCATAA
KC470219.1	GATTTATTTGTAGTGTATAGAGACAGTATACCGCATGCTGCATGCCATAA
KC470220.1	GATTTATTTGTAGTGTATAGAGACAGTATACCGCATGCTGCATGCCATAA
KC470221.1	GATTTATTTGTAGTGTATAGAGACAGTATACCGCATGCTGCATGCCATAA
KC470222.1	GATTTATTTGTAGTGTATAGAGACAGTATACCGCATGCTGCATGCCATAA
KC470223.1	GATTTATTTGTAGTGTATAGAGACAGTATACCGCATGCTGCATGCCATAA
KC470225.1	GATTTATTTGTAGTGTATAGAGACAGTATACCGCATGCTGCATGCCATAA
KC470226.1	GATTTATTTGTAGTGTATAGAGACAGTATACCGCATGCTGCATGCCATAA
KC470227.1	GATTTATTTGTAGTGTATAGAGACAGTATACCGCATGCTGCATGCCATAA
KC470228.1	GATTTATTTGTAGTGTATAGAGACAGTATACCGCATGCTGCATGCCATAA
KC470230.1	GATTTATTTGTAGTGTATAGAGACAGTATACCGCATGCTGCATGCCATAA
KC470229.1	GATTTATTTGTAGTGTATAGAGACAGTATACCGCATGCTGCATGCCATAA
KC470224.1	GATTTATTTGTAGTGTATAGAGACAGTATACCGCATGCTGCATGCCATAA
KX514433.1	GATTTATTTGTGGTGTATAGAGACAGTATACCGCATGCTGCATGCCATAA
KU298886.1	GATTTATTTGTGGTGTATAGAGACAGTATACCGCATGCTGCATGCCATAA
KC470213.1	GATTTATTTGTGGTGTATAGAGACAGTATACCGCATGCTGCATGCCATAA
	******
KC470218.1	ATGTATAGATTTCTATTCTAGAATTAGAGAATTAAGACATTATTCAGACT
KC470214.1	ATGTATAGATTTCTATTCTAGAATTAGAGAATTAAGACATTATTCAGACT
KC470215.1	ATGTATAGATTTCTATTCTAGAATTAGAGAATTAAGACATTATTCAGACT
KC470216.1	ATGTATAGATTTCTATTCTAGAATTAGAGAATTAAGACATTATTCAGACT

CLUSTAL	2.1	multiple	sequence	alignment
KC470215	7 1			<u>አጥር</u> ሞአጥአ <i>ር</i>

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KC470217.1	ATGTATAGATTTCTAGAATTAGAGAATTAAGACATTATTCAGACT
KC470219.1	ATGTATAGATTTCTAGAATTAGAGAATTAAGACATTATTCAGACT
KC470220.1	ATGTATAGATTTCTAGAATTAGAGAATTAAGACATTATTCAGACT
KC470221.1	ATGTATAGATTTCTAGAATTAGAGAATTAAGACATTATTCAGACT
KC470222.1	ATGTATAGATTTTATTCTAGAATTAGAGAATTAAGACATTATTCAGACT
KC470223.1	ATGTATAGATTTTATTCTAGAATTAGAGAATTAAGACATTATTCAGACT
KC470225.1	ATGTATAGATTTTATTCTAGAATTAGAGAATTAAGACATTATTCAGACT
KC470226.1	ATGTATAGATTTTTATTCTAGAATTAGAGAATTAAGACATTATTCAGACT
KC470227.1	ATGTATAGATTTTATTCTAGAATTAGAGAATTAAGACATTATTCAGACT
KC470228.1	ATGTATAGATTTTATTCTAGAATTAGAGAATTAAGACATTATTCAGACT
KC470230.1	ATGTATAGATTTTATTCTAGAATTAGAGAATTAAGACATTATTCAGACT
KC470229.1	ATGTATAGATTTTATTCTAGAATTAGAGAATTAAGACATTATTCAGACT
KC470224.1	ATGTATAGATTTTATTCTAGAATTAGAGAATTAAGACATTATTCAGACT
KX514433.1	ATGTATAGATTTTTATTCTAGAATTAGAGAATTAAGACATTATTCAGACT
КU298886.1	ATGTATAGATTTTATTCTAGAATTAGAGAATTAAGACATTATTCAGACT
KC470213.1	ATGTATAGATTTTATTCTAGAATTAGAGAATTAAGACATTATCAGACT
101,0213.1	**********
кС470218.1	CTGTGTATGGAGACACATTAGAAAAACTAACTAACACTGGGTTATACAAT
KC470214.1	CTGTGTATGGAGACACATTAGAAAAACTAACTAACACTGGGTTATACAAT
KC470214.1 KC470215.1	
	CTGTGTATGGAGACACATTAGAAAAACTAACTAACACTGGGTTATACAAT
KC470216.1	CTGTGTATGGAGACACATTAGAAAAACTAACTAACACTGGGTTATACAAT
KC470217.1	CTGTGTATGGAGACACATTAGAAAAACTAACTAACACTGGGTTATACAAT
KC470219.1	CTGTGTATGGAGACACATTAGAAAAACTAACTAACACTGGGTTATACAAT
KC470220.1	CTGTGTATGGAGACACATTAGAAAAACTAACTAACACTGGGTTATACAAT
KC470221.1	CTGTGTATGGAGACACATTAGAAAAACTAACTAACACTGGGTTATACAAT
KC470222.1	CTGTGTATGGAGACACATTAGAAAAACTAACTAACACTGGGTTATACAAT
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KC470225.1	CTGTGTATGGAGACACATTAGAAAAACTAACTAACACTGGGTTATACAAT
KC470226.1	CTGTGTATGGAGACACATTAGAAAAACTAACTAACACTGGGTTATACAAT
KC470227.1	CTGTGTATGGAGACACATTAGAAAAACTAACTAACACTGGGTTATACAAT
KC470228.1	CTGTGTATGGAGACACATTAGAAAAACTAACTAACACTGGGTTATACAAT
KC470230.1	CTGTGTATGGAGACACATTAGAAAAACTAACTAACACTGGGTTATACAAT
KC470229.1	$\tt CTGTGTATGGAGACACATTAGAAAAACTAACTAACACTGGGTTATACAAT$
KC470224.1	CTGTGTATGGAGACACATTAGAAAAACTAACTAACACTGGGTTATACAAT
KX514433.1	CTGTGTATGGAGACACATTGGAAAAACTAACTAACACTGGGTTATACAAT
KU298886.1	CTGTGTATGGAGACACATTGGAAAAACTAACTAACACTGGGTTATACAAT
KC470213.1	CTGTGTATGGAGACACATTGGAAAAACTAACTAACACTGGGTTATACAAT
	**********
KC470218.1	TTATTAATAAGGTGCCTGCGGTGCCAGAAACCGTTGAATCCAGCAGAAAA
KC470214.1	TTATTAATAAGGTGCCTGCGGTGCCAGAAACCGTTGAATCCAGCAGAAAA
KC470215.1	TTATTAATAAGGTGCCTGCGGTGCCAGAAACCGTTGAATCCAGCAGAAAA
KC470216.1	TTATTAATAAGGTGCCTGCGGTGCCAGAAACCGTTGAATCCAGCAGAAAA
KC470217.1	TTATTAATAAGGTGCCTGCGGTGCCAGAAACCGTTGAATCCAGCAGAAAA
KC470217.1	TTATTAATAAGGTGCCTGCGGTGCCAGAAACCGTTGAATCCAGCAGAAAA
KC470219.1 KC470220.1	TTATTAATAAGGTGCCTGCGGTGCCAGAAACCGTTGAATCCAGCAGAAAA
NCT/UZZU.1	AAAADAJOAJIAADI 10JJAAAADAJJOIOOJOIJOOI EDAAIAAIIAII

CLUSTAL	2.1	multiple	sequence	alignment
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Chosian 2.1 mulciple sequence aligi	menc
KC470221.1	TTATTAATAAGGTGCCTGCGGTGCCAGAAACCGTTGAATCCAGCAGAAAA
KC470222.1	TTATTAATAAGGTGCCTGCGGTGCCAGAAACCGTTGAATCCAGCAGAAAA
KC470223.1	TTATTAATAAGGTGCCTGCGGTGCCAGAAACCGTTGAATCCAGCAGAAAA
KC470225.1	TTATTAATAAGGTGCCTGCGGTGCCAGAAACCGTTGAATCCAGCAGAAAA
KC470226.1	TTATTAATAAGGTGCCTGCGGTGCCAGAAACCGTTGAATCCAGCAGAAAA
KC470227.1	TTATTAATAAGGTGCCTGCGGTGCCAGAAACCGTTGAATCCAGCAGAAAA
KC470228.1	TTATTAATAAGGTGCCTGCGGTGCCAGAAACCGTTGAATCCAGCAGAAAA
KC470230.1	TTATTAATAAGGTGCCTGCGGTGCCAGAAACCATTGAATCCAGCAGAAAA
KC470229.1	TTATTAATAAGGTGCCTGCGGTGCCAGAAACCATTGAATCCAGCAGAAAA
KC470224.1	TTATTAATAAGGTGCCTGCGGTGCCAGAAACCGTTGAATCCAGCAGAAAA
KX514433.1	TTATTAATAAGGTGCCTGCGGTGCCAGAAACCGTTGAATCCAGCAGAAAA
KU298886.1	TTATTAATAAGGTGCCTGCGGTGCCAGAAACCGTTGAATCCAGCAGAAAA
KC470213.1	TTATTAATAAGGTGCCTGCGGTGCCAGAAACCGTTGAATCCAGCAGAAAA
RC1/0213.1	*************
KC470218.1	ACTTAGACACCTTAATGAAAAACGACGATTCCACAAAATAGCTGGGCACT
KC470214.1	ACTTAGACACCTTAATGAAAAACGACGATTCCACAAAATAGCTGGGCACT
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KC470229.1	ACTTAGACACCTTAATGAAAAACGACGATTCCACAACATAGCTGGGCACT
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KX514433.1	ACTTAGACACCTTAATGAAAAACGACGATTCCACAACATAGCTGGGCACT
KU298886.1	ACTTAGACACCTTAATGAAAAACGACGATTCCACAACNNNGCTGGGCACT
KC470213.1	ACTTAGACACCTTAATGAAAAACGACGATTCCACAACATAGCTGGGCACT
	************************
KC470218.1	ATAGAGGCCAGTGCCATTCGTGCTGCAACCAAGCACGACAGGAGAGACTC
KC470214.1	ATAGAGGCCAGTGCCATTCGTGCTGCAACCGAGCACGACAGGAGAGACTC
KC470215.1	ATAGAGGCCAGTGCCATTCGTGCTGCAACCGAGCACGACAGGAGAGACTC
KC470216.1	ATAGAGGCCAGTGCCATTCGTGCTGCAACCGAGCACGACAGAGAGAG
KC470210.1	ATAGAGGCCAGTGCCATTCGTGCTGCAACCGAGCACGACAGGAGAGACTC
KC470219.1	ATAGAGGCCAGTGCCATTCGTGCTGCAACCGAGCACGACAGGAGAGACTC
KC470220.1	ATAGAGGCCAGTGCCATTCGTGCTGCAACCGAGCACGACAGGAGAGACTC
KC470221.1	ATAGAGGCCAGTGCCATTCGTGCTGCAACCGAGCACGACAGGAGAGACTC
KC470222.1	ATAGAGGCCAGTGCCATTCGTGCTGCAACCGAGCACGACAGGAGAGACTC
KC470223.1	ATAGAGGCCAGTGCCATTCGTGCTGCAACCGAGCACGACAGGAGAGACTC

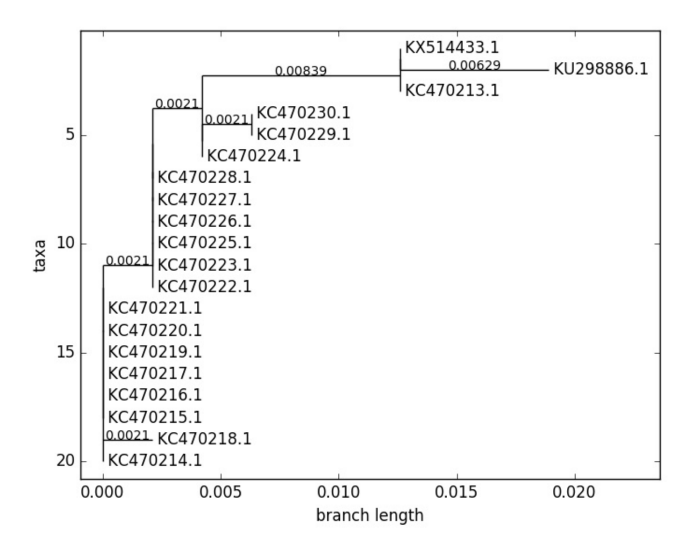
CLUSTAL 2	. 1	multiple	sequence	alignment
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KU298886.1

KC470213.1

CHOSTAN Z.I MUTCIPIE BEQUENCE ATTYM	menc
KC470225.1	ATAGAGGCCAGTGCCATTCGTGCTGCAACCGAGCACGACAGGAGAGACTC
KC470226.1	ATAGAGGCCAGTGCCATTCGTGCTGCAACCGAGCACGACAGGAGAGACTC
KC470227.1	ATAGAGGCCAGTGCCATTCGTGCTGCAACCGAGCACGACAGGAGAGACTC
KC470228.1	ATAGAGGCCAGTGCCATTCGTGCTGCAACCGAGCACGACAGGAGAGACTC
KC470230.1	ATAGAGGCCAGTGCCATTCGTGCTGCAACCGAGCACGACAGGAGAGACTC
KC470229.1	ATAGAGGCCAGTGCCATTCGTGCTGCAACCGAGCACGACAGGAGAGACTC
KC470224.1	ATAGAGGCCAGTGCCATTCGTGCTGCAACCGAGCACGACAGGAGAGACTC
KX514433.1	ATAGAGGCCAGTGCCATTCGTGCTGCAACCGAGCACGACAGGAAAGACTC
KU298886.1	ATAGAGGCCAGTGCCATTCGTGCTGCAACCGAGCACGACAGGAAAGACTC
KC470213.1	ATAGAGGCCAGTGCCATTCGTGCTGCAACCGAGCACGACAGGAAAGACTC
	********** ****** ****** ****** *****
KC470218.1	CAACGACGCAGAAACACAAGTATAA
KC470214.1	CAACGACGCAGAAACACAAGTATAA
KC470215.1	CAACGACGCAGAAACACAAGTATAA
KC470216.1	CAACGACGCAGAAACACAAGTATAA
KC470217.1	CAACGACGCAGAAACACAAGTATAA
KC470219.1	CAACGACGCAGAAACACAAGTATAA
KC470220.1	CAACGACGCAGAAACACAAGTATAA
KC470221.1	CAACGACGCAGAAACACAAGTATAA
KC470222.1	CAACGACGCAGAAACACAAGTATAA
KC470223.1	CAACGACGCAGAAACACAAGTATAA
KC470225.1	CAACGACGCAGAAACACAAGTATAA
KC470226.1	CAACGACGCAGAAACACAAGTATAA
KC470227.1	CAACGACGCAGAAACACAAGTATAA
KC470228.1	CAACGACGCAGAAACACAAGTATAA
KC470230.1	CAACGACGCAGAAACACAAGTATAA
KC470229.1	CAACGACGCAGAAACACAAGTATAA
KC470224.1	CAACGACGCAGAAACACAAGTATAA
KX514433.1	CAACGACGCAGAAACACAAGTATAA

CAACGACGCAGAGAAACACAAGTATAA



### Consensus sequence

ACCEPTABLE LEFT PRIMERS

	0-based		#			self	sel	f hair-	- qual-
# sequence	start	ln	N	GC%	Tm	any_th	end_t	h pir	n lity
0 GCGCTTTGAGGATCCAACAC	5	20	0	55.00	59.831	13.26	2.92	0.00	0.169
1 CGCTTTGAGGATCCAACACG	6	20	0	55.00	59.556	8.43	5.26	0.00	0.444
2 CTTTGAGGATCCAACACGGC	8	20	0	55.00	59.195	6.97	0.00	33.14	0.805
3 GCTTTGAGGATCCAACACGG	7	20	0	55.00	59.195	10.97	7.33	0.00	0.805
4 TTTGAGGATCCAACACGGCG	9	20	0	55.00	60.953	13.26	4.80	33.14	0.953
5 CGCGCTTTGAGGATCCAACA	4	20	0	55.00	61.298	13.26	9.79	0.00	1.298
6 TTGAGGATCCAACACGGCG	10	19	0	57.89	60.375	13.26	4.80	33.14	1.375
7 CGCGCTTTGAGGATCCAAC	4	19	0	57.89	59.575	8.27	2.56	0.00	1.425
8 GAGGATCCAACACGGCGAC	12	19	0	63.16	60.810	9.28	0.00	33.14	1.810
9 TTGAGGATCCAACACGGCGA	10	20	0	55.00	61.822	16.62	0.00	33.14	1.822
10 GATCCAACACGGCGACCCTA	15	20	0	60.00	61.952	0.00	0.00	33.14	1.952
11 ATCCAACACGGCGACCCTA	16	19	0	57.89	60.986	16.62	0.00	33.14	1.986
12 ATGGCGCGCTTTGAGGAT	0	18	0	55.56	60.125	14.45	0.00	0.00	2.125
13 ATGGCGCGCTTTGAGGATC	0	19	0	57.89	9 61.183	14.45	0.00	0.00	2.183
14 ATCCAACACGGCGACCCTAC	16	20	0	60.00	62.235	16.62	0.00	33.14	2.235
15 TGAGGATCCAACACGGCG	11	18	0	61.11	L 59.736	13.26	4.80	33.14	2.264
16 AGGATCCAACACGGCGAC	13	18	0	61.11	L 59.735	15.82	0.00	33.14	2.265
17 TGAGGATCCAACACGGCGA	11	19	0	57.89	61.280	16.62	0.00	33.14	2.280
18 CAACACGGCGACCCTACA	19	18	0	61.11	L 59.659	16.62	0.00	0.00	2.341
19 GCGCGCTTTGAGGATCCAA	3	19	0	57.89	61.399	26.67	0.00	0.00	2.399
20 CCAACACGGCGACCCTAC	18	18	0	66.67	7 60.434	16.62	0.00	0.00	2.434
21 TGAGGATCCAACACGGCGAC	11	20	0	60.00	62.501	13.26	0.00	33.14	2.501
22 GCGCTTTGAGGATCCAACA	5	19	0	52.63	3 58.445	13.26	9.79	0.00	2.555
23 GAGGATCCAACACGGCGA	12	18	0	61.11	L 59.423	16.62	0.00	33.14	2.577
24 GCGCGCTTTGAGGATCCAAC	3	20	0	60.00	62.584	26.67	2.56	0.00	2.584
25 TCCAACACGGCGACCCTA	17	18	0	61.11	L 60.601	16.62	0.00	33.14	2.601
26 GATCCAACACGGCGACCC	15	18	0	66.67	7 60.818	0.00	0.00	33.14	2.818
27 GGATCCAACACGGCGACC	14	18	0	66.67	7 60.818	6.95	0.00	33.14	2.818
28 GCGCGCTTTGAGGATCCA	3	18	0	61.11	L 60.820	26.67	0.00	0.00	2.820
29 TGGCGCGCTTTGAGGATC	1	18	0	61.11	L 60.820	32.29	11.34	0.00	2.820
30 CTTTGAGGATCCAACACGGCG	8	21	0	57.14	1 61.862	6.97	4.80	33.14	2.862
31 CGCTTTGAGGATCCAACACGG	6	21	0	57.14	1 61.862	2 10.97	7.33	0.00	2.862
32 TCCAACACGGCGACCCTAC	17	19	0	63.16	61.935	16.62	0.00	33.14	2.935
33 TTTGAGGATCCAACACGGC	9	19	0	52.63	3 58.059	13.26	0.00	33.14	2.941
34 GCTTTGAGGATCCAACACGGC	7	21	0	57.14	4 62.138	14.76	14.22	33.14	3.138
35 CCAACACGGCGACCCTACA	18	19	0	63.16	62.229	16.62	0.00	0.00	3.229
36 GATCCAACACGGCGACCCT	15	19	0	63.16	62.311	0.00	0.00	33.14	3.311
37 AGGATCCAACACGGCGACC	13	19	0	63.16	62.311	15.82	0.00	33.14	3.311
38 TTTGAGGATCCAACACGGCGA	9	21	0	52.38	62.314	16.62	0.00	33.14	3.314
39 ATCCAACACGGCGACCCT	16	18	0	61.11	61.319	16.62	0.00	33.14	3.319
40 GCGCTTTGAGGATCCAACACG	5	21	0	57.14	1 62.439	13.68	12.53	0.00	3.439
41 CGCGCTTTGAGGATCCAACAC	4	21	0	57.14	1 62.439	13.26	2.92	0.00	3.439
42 GGCGCGCTTTGAGGATCC	2	18	0	66.67	7 61.577	32.29	5.91	0.00	3.577
43 CGCGCTTTGAGGATCCAA	4	18	0	55.56	5 58.111	15.82	0.00	0.00	3.889
44 TTGAGGATCCAACACGGCGAC	10	21	0	57.14	1 62.955	13.26	0.00	33.14	3.955

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ACCEPTABLE RIGHT PRIMERS		
	0-based	# self self hair- qual-
# sequence	start ln	N GC% Tm any_th end_th pin lity
0 ACTGGCCTCTATAGTGCCCA	411 20	0 55.00 60.031 15.67 7.83 37.93 0.031
1 TTCTCTGCGTCGTTGGAGTC	464 20	0 55.00 60.039 9.66 0.00 0.00 0.039
2 TGTTTCTCTGCGTCGTTGGA	467 20	0 50.00 59.897 0.00 0.00 0.00 0.103
3 GTGTTTCTCTGCGTCGTTGG	468 20	0 55.00 59.768 0.00 0.00 0.00 0.232
4 GCACTGGCCTCTATAGTGCC	413 20	0 60.00 60.250 26.85 21.72 38.72 0.250
5 TTTCTCTGCGTCGTTGGAGT	465 20	0 50.00 59.612 9.66 0.00 0.00 0.388
6 CACTGGCCTCTATAGTGCCC	412 20	0 60.00 59.605 24.00 12.62 37.93 0.395
7 GAGTCTCTCCTGTCGTGCTC	449 20	0 60.00 59.550 0.00 0.00 0.00 0.450
8 CTCTGCGTCGTTGGAGTCTC	462 20	0 60.00 60.455 4.08 0.00 0.00 0.455
9 CTGCGTCGTTGGAGTCTCTC	460 20	0 60.00 60.455 0.00 0.00 0.00 0.455
10 CTGGCCTCTATAGTGCCCAG	410 20	0 60.00 59.316 18.76 14.40 37.93 0.684
11 GTTTCTCTGCGTCGTTGGAG	466 20	0 55.00 59.215 5.37 5.37 0.00 0.785
12 GCAGCATGCGGTATACTGTC	190 20	0 55.00 59.138 14.57 0.71 37.60 0.862
13 TCTCTGCGTCGTTGGAGTCT	463 20	0 55.00 60.890 9.66 0.00 0.00 0.890
14 TCTGCGTCGTTGGAGTCTCT	461 20	0 55.00 60.890 0.00 0.00 0.00 0.890
15 TGTGTTTCTCTGCGTCGTTG	469 20	0 50.00 59.072 0.00 0.00 0.00 0.928
16 ATTCAACGGTTTCTGGCACC	338 20	0 50.00 59.041 0.00 0.00 0.00 0.959
17 GGAGTCTCTCCTGTCGTGCT	450 20	0 60.00 60.963 8.38 0.00 46.41 0.963
18 GGATTCAACGGTTTCTGGCAC	340 21	0 52.38 60.069 0.00 0.00 0.00 1.069
19 GATTCAACGGTTTCTGGCACC	339 21	0 52.38 60.069 0.00 0.00 0.00 1.069
20 AGTGCCCAGCTATTTTGTGGA	399 21	0 47.62 59.925 0.00 0.00 0.00 1.075
21 GGCCTCTATAGTGCCCAGCT	408 20	0 60.00 61.124 19.28 0.00 34.65 1.124
22 CTCGGTTGCAGCACGAATG	432 19	0 57.89 59.867 13.22 2.38 0.00 1.133
23 GTCTCTCCTGTCGTGCTCG	447 19	0 63.16 59.864 0.81 0.81 38.55 1.136
24 GCTGGATTCAACGGTTTCTGG	343 21	0 52.38 59.800 0.00 0.00 0.00 1.200
25 CTGGATTCAACGGTTTCTGGC	342 21	0 52.38 59.800 0.00 0.00 0.00 1.200
26 ACTTGTGTTTCTCTGCGTCG	472 20	0 50.00 58.791 0.00 0.00 0.00 1.209
27 CTTGTGTTTCTCTGCGTCGT	471 20	0 50.00 58.791 0.00 0.00 0.00 1.209
28 TGGATTCAACGGTTTCTGGC	341 20	0 50.00 58.756 0.00 0.00 0.00 1.244
29 GGATTCAACGGTTTCTGGCA	340 20	0 50.00 58.756 0.00 0.00 0.00 1.244
30 AGTCTCTCTGTCGTGCTCG	448 20	0 60.00 61.302 0.81 0.81 38.55 1.302
31 TTGTGTTTCTCTGCGTCGTTG	470 21	0 47.62 59.674 0.00 0.00 0.00 1.326
32 GGCCTCTATAGTGCCCAGC	408 19	0 63.16 59.627 19.28 8.03 34.65 1.373
33 TGGCCTCTATAGTGCCCAGC	409 20	0 60.00 61.411 15.67 2.73 37.93 1.411 0 47.62 60.471 0.00 0.00 0.00 1.471
34 ACTTGTGTTTCTCTGCGTCGT	472 21	
35 TGGATTCAACGGTTTCTGGCA	341 21	0 47.62 60.478 0.00 0.00 0.00 1.478 0 63.16 59.491 8.38 0.00 46.41 1.509
36 GGAGTCTCTCCTGTCGTGC  37 GCAGCATGCGGTATACTGTCT	450 19 190 21	0 63.16 59.491 8.38 0.00 46.41 1.509 0 52.38 60.539 14.57 1.67 37.60 1.539
37 GCAGCATGCGGTATACTGTCT 38 GTGCCCAGCTATTTTGTGGA	398 20	
	398 20	
39 AGTGCCCAGCTATTTTGTGG	399 20	0 50.00 58.451 0.00 0.00 0.00 1.549

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Pr	imers data									
40	TCTCTGCGTCGTTGGAGTC	463	19	0	57.89	59.420	9.66	0.00	0.00	1.580
41	TCTGCGTCGTTGGAGTCTC	461	19	0	57.89	59.420	0.00	0.00	0.00	1.580
42	TGCGTCGTTGGAGTCTCTC	459	19	0	57.89	59.420	0.00	0.00	0.00	1.580
43	CTCTGCGTCGTTGGAGTCT	462	19	0	57.89	59.418	4.08	0.00	0.00	1.582
44	CTGCGTCGTTGGAGTCTCT	460	19	0	57.89	59.418	0.00	0.00	0.00	1.582
45	CTTGTGTTTCTCTGCGTCGTT	471	21	0	47.62	59.407	0.00	0.00	0.00	1.593
46	TTTCTCTGCGTCGTTGGAGTC	465	21	0	52.38	60.602	9.66	0.00	0.00	1.602
47	GCCTCTATAGTGCCCAGCTA	407	20	0	55.00	58.366	5.93	0.00	0.00	1.634
48	TTGTGTTTCTCTGCGTCGTT	470	20	0	45.00	58.353	0.00	0.00	0.00	1.647
49	GTTGCAGCACGAATGGCAC	428	19	0	57.89	60.731	0.00	0.00	0.00	1.731
50	CTCCTGTCGTGCTCGGTTG	443	19	0	63.16	60.734	0.00	0.00	37.69	1.734
51	GGCCTCTATAGTGCCCAGCTA	408	21				19.28	0.00	34.65	1.828
52	CTCTCCTGTCGTGCTCGGTT	445				61.868	0.00		37.69	1.868
53	TGTTTCTCTGCGTCGTTGGAG	467				60.868	5.37	5.37	0.00	1.868
54	GTTTCTCTGCGTCGTTGGAGT	466				60.869	9.66	0.00	0.00	1.869
55	TGCTGGATTCAACGGTTTCTG	344				59.121	0.00	0.00	0.00	1.879
56	GTGCCCAGCTATTTTGTGGAA	398				59.108	0.00	0.00	0.00	1.892
57	TTGCAGCACGAATGGCACT	427				60.898	0.00	0.00	0.00	1.898
58	TCTCCTGTCGTGCTCGGTT	444				60.902	0.00		37.69	1.902
59	CAGCATGCGGTATACTGTCTCT	189				59.965	6.85	0.00	0.00	2.035
60	ACTGGCCTCTATAGTGCCCAG	411				61.037				2.033
61	TTCTCTGCGTCGTTGGAGT	464				58.961	9.66	0.00	0.00	2.037
62	TCTCTCCTGTCGTGCTCGG	446				61.041	8.49		37.69	2.039
63	CTGCTGGATTCAACGGTTTCTG	345				60.096		0.00	0.00	2.041
							0.00			
64	GGTTGCAGCACGAATGC	429				60.126	2.05	0.00	0.00	2.126
65	GTGTTTCTCTGCGTCGTTGGA	468					0.00	0.00	0.00	2.132
66	TTGCAGCACGAATGGCACTG	427				62.133	0.00	0.00	0.00	2.133
67	GTTGCAGCACGAATGGCACT	428				62.134	0.00	0.00	0.00	2.134
	TCTCCTGTCGTGCTCGGTTG	444				62.142	0.00		37.69	2.142
	TCTGCTGGATTCAACGGTTTC		21			58.849				
	CTGCTGGATTCAACGGTTTCT	345				58.846	0.00	0.00	0.00	2.154
	CGGTTGCAGCACGAATGG	430				59.823	5.71	0.00	0.00	2.177
	GCAGCACGAATGGCACTG	425				59.822	0.00	0.00	0.00	2.178
	GCCTCTATAGTGCCCAGCTAT	407				58.816	5.93	0.00	0.00	2.184
	GCCCAGCTATTTTGTGGAATCG	396				60.224	0.00	0.00	0.00	2.224
75	TACTTGTGTTTCTCTGCGTCGT	473				60.224	0.00	0.00	0.00	2.224
76	TCTGCTGGATTCAACGGTTTCT	346				60.225	0.00	0.00	0.00	2.225
77	CTCGGTTGCAGCACGAATGG	432				62.244	0.00	0.00	0.00	2.244
78	GTCTCTCCTGTCGTGCTCGG	447	20			62.260	8.49	0.00	37.69	2.260
79	TTTCTGCTGGATTCAACGGT	348	20	0	45.00	57.730	0.00	0.00	0.00	2.270
80	TTCTGCTGGATTCAACGGTT	347	20	0	45.00	57.730	0.00	0.00	0.00	2.270
81	TCTGCTGGATTCAACGGTTT	346	20	0	45.00	57.730	0.00	0.00	0.00	2.270
82	TGCTGGATTCAACGGTTTCT	344	20	0	45.00	57.730	0.00	0.00	0.00	2.270
83	TGCAGCACGAATGGCACT	426	18	0	55.56	60.282	0.00	0.00	0.00	2.282
84	TCTCCTGTCGTGCTCGGT	444	18	0	61.11	60.282	0.00	0.00	37.69	2.282
85	TGCCCAGCTATTTTGTGGAA	397	20	0	45.00	57.699	0.00	0.00	0.00	2.301
86	TAGTGCCCAGCTATTTTGTGGA	400	22	0	45.45	59.693	0.00	0.00	0.00	2.307

Primers data				
87 CACTGGCCTCTATAGTGCCCA	412 21	0 57.14 61.311 1	5.67 7.83 37.93	2.311
88 AGCATGCGGTATACTGTCTCT	188 21	0 47.62 58.689 1	3.01 0.00 0.00	2.311
89 TCGGTTGCAGCACGAATGG	431 19	0 57.89 61.324	0.00 0.00 0.00	2.324
90 GATTCAACGGTTTCTGGCAC	339 20	0 50.00 57.668	0.00 0.00 0.00	2.332
91 CTCTCCTGTCGTGCTCGGT	445 19	0 63.16 61.336	0.00 0.00 37.69	2.336
92 TCCTGTCGTGCTCGGTTG	442 18	0 61.11 59.661	0.00 0.00 37.69	2.339
93 TACTTGTGTTTCTCTGCGTCG	473 21	0 47.62 58.609	0.00 0.00 0.00	2.391
94 TGTGTTTCTCTGCGTCGTTGG	469 21	0 52.38 61.396	0.00 0.00 0.00	2.396
95 CAGCATGCGGTATACTGTCTC	189 21	0 52.38 58.602	6.85 0.00 0.00	2.398
96 TTCAACGGTTTCTGGCACC	337 19	0 52.63 58.590	0.00 0.00 0.00	2.410
97 TTCTCTGCGTCGTTGGAGTCT	464 21	0 52.38 61.422	9.66 0.00 0.00	2.422
98 GCCTCTATAGTGCCCAGCT	407 19	0 57.89 58.553	5.93 0.00 0.00	2.447
99 CTCTCCTGTCGTGCTCGG	445 18	0 66.67 59.513	8.49 0.00 37.69	2.487
100 AGTGCCCAGCTATTTTGTGGAA	399 22	0 45.45 60.491	0.00 0.00 0.00	2.491
101 GTGCCCAGCTATTTTGTGGAAT	398 22	0 45.45 59.503	0.00 0.00 0.00	2.497
102 CAGCATGCGGTATACTGTCT	189 20	0 50.00 57.484	9.90 1.67 0.00	2.516
103 TTCTGCTGGATTCAACGGTTTC	347 22	0 45.45 59.448	0.00 0.00 0.00	2.552
104 GAGTCTCTCCTGTCGTGCT	449 19	0 57.89 58.438	0.00 0.00 0.00	2.562
105 AGTCTCTCCTGTCGTGCTC	448 19	0 57.89 58.438	0.00 0.00 0.00	2.562
106 GCCTCTATAGTGCCCAGCTATT	407 22	0 50.00 59.431	5.93 0.00 0.00	2.569
107 TTTTCTGCTGGATTCAACGGT	349 21	0 42.86 58.418	0.00 0.00 0.00	2.582
108 TTTCTGCTGGATTCAACGGTT	348 21	0 42.86 58.418	0.00 0.00 0.00	2.582
109 TTCTGCTGGATTCAACGGTTT	347 21	0 42.86 58.418	0.00 0.00 0.00	2.582
110 CTTGTGTTTCTCTGCGTCGTTG	471 22	0 50.00 60.598	0.00 0.00 0.00	2.598
111 TGTTTCTCTGCGTCGTTGG	467 19	0 52.63 58.400	0.00 0.00 0.00	2.600
112 CTGCTGGATTCAACGGTTTC	345 20	0 50.00 57.381	0.00 0.00 0.00	2.619
113 GCTGGATTCAACGGTTTCTG	343 20	0 50.00 57.381	0.00 0.00 0.00	2.619
114 TGCAGCACGAATGGCACTG	426 19	0 57.89 61.619	0.00 0.00 0.00	2.619
115 GTTGCAGCACGAATGGCA	428 18	0 55.56 59.356	0.00 0.00 0.00	2.644
116 TTGCAGCACGAATGGCAC	427 18	0 55.56 59.356	0.00 0.00 0.00	2.644
117 CTCCTGTCGTGCTCGGTT	443 18	0 61.11 59.349	0.00 0.00 37.69	2.651
118 CTGTCGTGCTCGGTTGCA	440 18	0 61.11 60.662	3.16 0.00 41.05	2.662
119 TGTCGTGCTCGGTTGCAG	439 18	0 61.11 60.662	3.16 0.00 44.20	2.662
120 CTGTCGTGCTCGGTTGCAG	440 19	0 63.16 61.662	7.90 7.90 44.20	2.662
121 GCTCGGTTGCAGCACGAAT	433 19	0 57.89 61.681	13.22 0.00 0.00	2.681
122 TAGTGCCCAGCTATTTTGTGG	400 21	0 47.62 58.278	0.00 0.00 0.00	2.722
123 TCTCTCCTGTCGTGCTCGGT	446 20	0 60.00 62.750	0.00 0.00 37.69	2.750
124 TGCCCAGCTATTTTGTGGAATC	397 22	0 45.45 59.240	0.00 0.00 0.00	2.760
125 AGCATGCGGTATACTGTCTC	188 20	0 50.00 57.197	13.01 0.00 0.00	2.803
126 GCATGCGGTATACTGTCTCT	187 20	0 50.00 57.197	3.72 0.00 0.00	2.803
127 TCTCTGCGTCGTTGGAGTCTC	463 21	0 57.14 61.806	9.66 0.00 0.00	2.806
128 TCTGCGTCGTTGGAGTCTCTC	461 21	0 57.14 61.806	0.00 0.00 0.00	2.806
129 CTCTGCGTCGTTGGAGTCTCT	462 21	0 57.14 61.810	4.08 0.00 0.00	2.810
130 TGCCCAGCTATTTTGTGGAAT	397 21	0 42.86 58.181	0.00 0.00 0.00	2.819
131 GCTCGGTTGCAGCACGAATG	433 20	0 60.00 62.839	0.00 0.00 0.00	2.839
132 CTGGCCTCTATAGTGCCCA	410 19	0 57.89 58.160	15.67 7.83 37.93	2.840
133 TGGCCTCTATAGTGCCCAG	409 19	0 57.89 58.160	15.67 0.00 37.93	2.840

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Prıı	mers data								
134	ACTGGCCTCTATAGTGCCC	411 19	0	57.89	58.158	24.00	12.62	37.93	2.842
135	CAGCACGAATGGCACTGG	424 18	0	61.11	59.127	0.00	0.00	0.00	2.873
136	GGAGTCTCTCCTGTCGTGCTC	450 21	0	61.90	61.878	8.38	0.00	46.41	2.878
137	GTTTCTCTGCGTCGTTGGA	466 19	0	52.63	58.105	0.00	0.00	0.00	2.895
138	GGTTGCAGCACGAATGGCA	429 19	0	57.89	61.918	2.05	0.00	0.00	2.918
139	TTTTTCTGCTGGATTCAACGGT	350 22	0	40.91	59.044	0.00	0.00	0.00	2.956
140	TTTTCTGCTGGATTCAACGGTT	349 22	0	40.91	59.044	0.00	0.00	0.00	2.956
141	TTTCTGCTGGATTCAACGGTTT	348 22	0	40.91	59.044	0.00	0.00	0.00	2.956
142	ATACTTGTGTTTCTCTGCGTCG	474 22	0	45.45	59.013	0.00	0.00	0.00	2.987
143	GCAGCATGCGGTATACTGT	190 19	0	52.63	58.008	14.57	0.00	37.60	2.992
	ACTTGTGTTTCTCTGCGTCGTT	472 22			60.994	0.00	0.00	0.00	2.994
145	TTTCTGCTGGATTCAACGGTTTC	348 23			59.996	0.00	0.00	0.00	3.004
146	AGTTTTCTGCTGGATTCAACGG	352 23			59.995	0.00	0.00	0.00	3.005
147	GCCTCTATAGTGCCCAGCTATTT	407 23			59.992	5.93	0.00	0.00	3.008
148	CCCAGCTATTTTGTGGAATCGT	395 22	0		58.983	0.00	0.00	0.00	3.017
149	ATAGTGCCCAGCTATTTTGTGGA	401 23	-		60.054	0.00	0.00	0.00	3.054
150	AGCACGAATGGCACTGGC	423 18			61.058	0.00		40.13	3.058
151		395 23			59.935	0.00	0.00	0.00	3.065
152	GGCCTCTATAGTGCCCAGCTAT	408 22			61.156			34.65	3.156
153	GAGTCTCTCCTGTCGTGCTCG	449 21			62.184	0.81		38.55	3.184
154	TTTCTCTGCGTCGTTGGAG	465 19			57.805	5.37	5.37	0.00	3.195
155	CCAGCTATTTGTGGAATCGTCG	394 23			60.242	0.00	0.00	0.00	3.242
156	GTTTTCTGCTGGATTCAACGGT	351 23			60.243	0.00	0.00	0.00	3.242
157	TAGTGCCCAGCTATTTTGTGGAA	400 23			60.245	0.00	0.00	0.00	3.245
158	TCGGTTGCAGCACGAATG	431 18			58.747	0.00	0.00	0.00	3.243
159	CAGCATGCGGTATACTGTCTCTA	189 23			59.747	6.85	0.00	0.00	3.253
160	ATAGTGCCCAGCTATTTTGTGG	401 22			58.709	0.00	0.00	0.00	3.291
	GCACTGGCCTCTATAGTGC	413 19			57.703			32.60	3.291
	GTTTTCTGCTGGATTCAACGG	351 22			58.691	0.00	0.00		3.309
	CTGGCCTCTATAGTGCCCAGC	410 21			62.318			0.00	3.318
		425 19			62.332	3.18	0.00		
	GCAGCACGAATGGCACTGG							0.00	3.332
	CAGCACGAATGGCACTGGC	424 19			62.332	0.00		40.13	3.332
	GCTCGGTTGCAGCACGAA	433 18			61.346		1.56	0.00	3.346
	TTGTGTTTCTCTGCGTCGT	470 19			57.644	0.00	0.00	0.00	3.356
	TGTGTTTCTCTGCGTCGTT	469 19			57.644	0.00	0.00	0.00	3.356
	CAGCTATTTTGTGGAATCGTCGT	393 23			59.629	0.00	0.00	0.00	3.371
	TTTTTCTGCTGGATTCAACGGTT	350 23			59.617	0.00	0.00	0.00	3.383
	TTTTCTGCTGGATTCAACGGTTT	349 23			59.617	0.00	0.00	0.00	3.383
	TGCTGGATTCAACGGTTTCTGG	344 22			61.392	0.00	0.00	0.00	3.392
	CTGGATTCAACGGTTTCTGGCA	342 22			61.392	0.00	0.00	0.00	3.392
	CCTGTCGTGCTCGGTTGC	441 18			61.415	0.00		37.69	3.415
	GTGCCCAGCTATTTTGTGGAATC	398 23			60.429	0.00	0.00	0.00	3.429
	GCAGCATGCGGTATACTGTCTC	190 22			61.430			37.60	3.430
	GCCCAGCTATTTTGTGGAATC	396 21			57.557	0.00	0.00	0.00	3.443
	AGCATGCGGTATACTGTCTCTA	188 22			58.513		0.00	0.00	3.487
	ATACTTGTGTTTCTCTGCGTCGT	474 23			60.552	0.00	0.00	0.00	3.552
180	CTCGGTTGCAGCACGAAT	432 18	0	55.56	58.434	13.22	0.00	0.00	3.566

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181	TCTCTCCTGTCGTGCTCG	446 18	0	61.11	58.420	0.81	0.81	38.55	3.580
182	GCACTGGCCTCTATAGTGCCC	413 21	0	61.90	62.584	26.85	12.62	37.93	3.584
183	AGCTATTTTGTGGAATCGTCGT	392 22	0	40.91	58.404	0.00	0.00	0.00	3.596
184	TTTTTCTGCTGGATTCAACGG	350 21	0	42.86	57.359	0.00	0.00	0.00	3.641
185	TGGATTCAACGGTTTCTGGCAC	341 22	0	50.00	61.648	0.00	0.00	0.00	3.648
186	CCCAGCTATTTTGTGGAATCG	395 21	0	47.62	57.302	0.00	0.00	0.00	3.698
187	GTGTTTCTCTGCGTCGTTG	468 19	0	52.63	57.290	0.00	0.00	0.00	3.710
188	GTTTCTCTGCGTCGTTGGAGTC	466 22	0	54.55	61.732	2.30	0.00	0.00	3.732
189	TACTTGTGTTTCTCTGCGTCGTT	473 23	0	43.48	60.736	0.00	0.00	0.00	3.736
190	TTCTGCTGGATTCAACGGTTTCT	347 23	0	43.48	60.747	0.00	0.00	0.00	3.747
191	TCTCTGCGTCGTTGGAGT	463 18	0	55.56	58.240	9.66	0.00	0.00	3.760
192	TCTGCGTCGTTGGAGTCT	461 18	0	55.56	58.240	0.00	0.00	0.00	3.760
193	TGCGTCGTTGGAGTCTCT	459 18	0	55.56	58.240	0.00	0.00	0.00	3.760
194	TGGCCTCTATAGTGCCCAGCT	409 21	0	57.14	62.801	15.67	0.00	37.93	3.801
195	AGTGCCCAGCTATTTTGTGGAAT	399 23	0	43.48	60.819	0.00	0.00	0.00	3.819
196	TCCTGTCGTGCTCGGTTGC	442 19	0	63.16	62.859	0.00	0.00	37.69	3.859
197	TTGTGTTTCTCTGCGTCGTTGG	470 22	0	50.00	61.873	0.00	0.00	0.00	3.873
198	TTTCTCTGCGTCGTTGGAGTCT	465 22	0	50.00	61.907	9.66	0.00	0.00	3.907
199	GCATGCGGTATACTGTCTCTA	187 21	0	47.62	57.083	3.72	0.00	0.00	3.917
200	CAGCTATTTTGTGGAATCGTCG	393 22	0	45.45	58.072	0.00	0.00	0.00	3.928
201	CGGTTGCAGCACGAATGGC	430 19	0	63.16	62.952	5.71	0.00	0.00	3.952
202	CTCTCCTGTCGTGCTCGGTTG	445 21	0	61.90	62.980	0.00	0.00	37.69	3.980
203	GTGTTTCTCTGCGTCGTTGGAG	468 22	0	54.55	61.982	5.37	5.37	0.00	3.982
204	CACTGGCCTCTATAGTGCC	412 19	0	57.89	57.002	21.72	21.72	38.72	3.998
205	AGCTATTTTGTGGAATCGTCGTT	392 23	0	39.13	58.999	0.00	0.00	0.00	4.001
206	TGTCTCCATACACAGAGTCTGA	264 22	0	45.45	57.966	26.86	4.71	35.18	4.034
207	GTCGTGCTCGGTTGCAGC	438 18	0	66.67	62.077	0.00	0.00	44.20	4.077
208	CAGCATGCGGTATACTGTCTCTAT	189 24	0	45.83	60.083	6.85	0.00	0.00	4.083
	AGCATGCGGTATACTGTCTCTAT	188 23			58.917		0.00	0.00	4.083
210	GGTGTCTAAGTTTTTCTGCTGGA	360 23	0	43.48	58.865	0.00	0.00	0.00	4.135
211	AGGTGTCTAAGTTTTTCTGCTGG	361 23			58.862	0.00	0.00	0.00	4.138
212	AGGTGTCTAAGTTTTTCTGCTGGA	361 24			60.141	0.00	0.00	0.00	4.141
213	TTTTTCTGCTGGATTCAACGGTTT	350 24			60.142	0.00	0.00	0.00	4.142
	CAGCTATTTTGTGGAATCGTCGTT	393 24			60.143	0.00	0.00	0.00	4.143
215	TCAACGGTTTCTGGCACC	336 18			57.849	0.00	0.00	0.00	4.151
	CTCTGCGTCGTTGGAGTC	462 18			57.845	4.08	0.00	0.00	4.155
217	CTGCGTCGTTGGAGTCTC	460 18			57.845	0.00	0.00	0.00	4.155
218	TATACTTGTGTTTCTCTGCGTCG	475 23			58.836	0.00	0.00	0.00	4.164
219	TATAGTGCCCAGCTATTTTGTGGA	402 24			59.835	0.00	0.00	0.00	4.165
	CACTGGCCTCTATAGTGCCCAG	412 22			62.185				4.185
	AGTTTTTCTGCTGGATTCAACG	352 22			57.771	0.00	0.00	0.00	4.229
	TTCTCTGCGTCGTTGGAGTCTC	464 22			62.268	9.66	0.00	0.00	4.268
	GGATTCAACGGTTTCTGGCACC	340 22			62.279	0.00	0.00	0.00	4.279
	CCAGCTATTTTGTGGAATCGTC	394 22			57.721	0.00	0.00	0.00	4.279
	TATACTTGTGTTTCTCTGCGTCGT	475 24			60.321	0.00	0.00	0.00	4.321
	GCTATTTGTGGAATCGTCGTT	391 22			57.663	0.00	0.00	0.00	4.337
227	TCTGCTGGATTCAACGGTTTCTG	346 23	0	47.83	61.360	0.00	0.00	0.00	4.360

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228	CGTGCTCGGTTGCAGCAC	436	18	0	66.67	62.375	15.86	15.86	44.20	4.375
229	GTGCTCGGTTGCAGCACG	435	18	0	66.67	62.375	13.30	9.35	44.20	4.375
230	TGTCTCCATACACAGAGTCTGAA	264	23	0	43.48	58.593	26.86	11.56	35.18	4.407
231	TGTTTCTCTGCGTCGTTGGAGT	467	22	0	50.00	62.417	9.66	0.00	0.00	4.417
232	TGGCCTCTATAGTGCCCAGCTA	409	22	0	54.55	62.440	15.67	1.80	37.93	4.440
233	GCATGCGGTATACTGTCTCTAT	187	22	0	45.45	57.558	3.72	0.00	0.00	4.442
234	AGCTATTTGTGGAATCGTCGTTT	392	24	0	37.50	59.545	0.00	0.00	0.00	4.455
235	TATAGTGCCCAGCTATTTTGTGG	402	23	0	43.48	58.540	0.00	0.00	0.00	4.460
236	GCGTCGTTGGAGTCTCTC	458	18	0	61.11	57.538	0.00	0.00	0.00	4.462
237	GGTGTCTAAGTTTTTCTGCTGG	360	22	0	45.45	57.510	0.00	0.00	0.00	4.490
238	TTTTCTGCTGGATTCAACGGTTTC	349	24	0	41.67	60.499	0.00	0.00	0.00	4.499
239	AAGTTTTTCTGCTGGATTCAACGG	353	24	0	41.67	60.500	0.00	0.00	0.00	4.500
240	GCCTCTATAGTGCCCAGCTATTTT	407	24	0	45.83	60.506	5.93	0.00	0.00	4.506
241	CTATAGTGCCCAGCTATTTTGTGG	403	24	0	45.83	59.483	0.00	0.00	0.00	4.517
242	ATAGTGCCCAGCTATTTTGTGGAA	401	24	0	41.67	60.570	0.00	0.00	0.00	4.570
243	TAGTGCCCAGCTATTTTGTGGAAT	400	24	0	41.67	60.570	0.00	0.00	0.00	4.570
244	AAGGTGTCTAAGTTTTTCTGCTGG	362	24	0	41.67	59.419	0.00	0.00	0.00	4.581
245	GCTGGATTCAACGGTTTCTGGC	343	22			62.585	0.00	0.00	0.00	4.585
246	TCGTGCTCGGTTGCAGCA	437	18	0	61.11	62.601	8.49	0.00	44.20	4.601
247	TGCTCGGTTGCAGCACGA	434	18	0	61.11	62.601	12.06	12.06	38.12	4.601
248	AAGTTTTTCTGCTGGATTCAACG	353	23	0	39.13	58.389	0.00	0.00	0.00	4.611
249	TTTGTGGAATCGTCGTTTTTCA	386	22	0	36.36	57.382	7.33	0.88	0.00	4.618
250	TTATACTTGTGTTTCTCTGCGTCG	476	24	0	41.67	59.380	0.00	0.00	0.00	4.620
251	CTCTGCGTCGTTGGAGTCTCTC	462	22	0	59.09	62.624	4.08	0.00	0.00	4.624
252	GGCCTCTATAGTGCCCAGCTATT	408	23	0	52.17	61.644	19.28	0.00	34.65	4.644
253	TGTGTTTCTCTGCGTCGTTGGA	469	22	0	50.00	62.667	0.00	0.00	0.00	4.667
254	GCTATTTTGTGGAATCGTCGTTT	391	23	0	39.13	58.281	0.00	0.00	0.00	4.719
255	GTTTTTCTGCTGGATTCAACGGTT	351	24	0	41.67	60.736	0.00	0.00	0.00	4.736
	TGCCCAGCTATTTTGTGGAATCG	397	_			61.737	0.00	0.00	0.00	4.737
257	GCCCAGCTATTTTGTGGAATCGT	396	23	0	47.83	61.738	0.00	0.00	0.00	4.738
258	GGTGTCTAAGTTTTTCTGCTGGAT	360	24	0	41.67	59.238	0.00	0.00	0.00	4.762
259	TTGTGGAATCGTCGTTTTTCAT	385	22	0	36.36	57.178	1.86	0.00	0.00	4.822
260	TGTGGAATCGTCGTTTTTCATT	384	22	0	36.36	57.178	1.86	0.00	0.00	4.822
261	AGTCTCTCCTGTCGTGCT	448	18	0	55.56	57.167	0.00	0.00	0.00	4.833
262	CCTCTATAGTGCCCAGCTATTT	406	22	0	45.45	57.101	0.00	0.00	0.00	4.899
263	TTCTCTGCGTCGTTGGAG	464	18	0	55.56	57.034	5.37	5.37	0.00	4.966
264	TTTTGTGGAATCGTCGTTTTTCA	387	23	0	34.78	58.017	7.33	0.88	0.00	4.983
265	CCTCTATAGTGCCCAGCTATTTTG	406	24	0	45.83	59.000	0.00	0.00	0.00	5.000
266	TGTCTCCATACACAGAGTCTGAAT	264	24	0	41.67	58.981	13.25	4.49	35.18	5.019
267	ATACTTGTGTTTCTCTGCGTCGTT	474	24	0	41.67	61.032	0.00	0.00	0.00	5.032
268	AGCTATTTGTGGAATCGTCGTTTT	392	25	0	36.00	60.049	0.00	0.00	0.00	5.049
269	TCGTCGTTTTTCATTAAGGTGTCT	377	24			58.948	0.00	0.00	0.00	5.052
270	ACTTGTGTTTCTCTGCGTCGTTG	472	23	0	47.83	62.062	0.00	0.00	0.00	5.062
271	GCATGCGGTATACTGTCTCTATACA	187	25	0	44.00	60.106	3.72	0.00	31.23	5.106
272	CAGCATGCGGTATACTGTCTCTATA	189	25	0	44.00	59.876	6.85	0.00	0.00	5.124
273	AGCATGCGGTATACTGTCTCTATAC	188	25	0	44.00	59.876	13.01	0.00	0.00	5.124
274	GCTATTTGTGGAATCGTCGTTTT	391	24	0	37.50	58.849	0.00	0.00	0.00	5.151

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275	ACTGTCTTGCAATATACACAGGT	112 23	0 39.13	57.834	18.84	9.32 0.00	5.166
276	TTTGTGGAATCGTCGTTTTTCAT	386 23	0 34.78	57.824	1.86	0.00 0.00	5.176
277	TTGTGGAATCGTCGTTTTTCATT	385 23	0 34.78	57.824	1.86	0.00 0.00	5.176
278	CTGCTGGATTCAACGGTTTCTGG	345 23	0 52.17	62.211	0.00	0.00 0.00	5.211
279	TTTCTGCTGGATTCAACGGTTTCT	348 24	0 41.67	61.227	0.00	0.00 0.00	5.227
280	CCTCTATAGTGCCCAGCTATTTT	406 23	0 43.48	57.766	0.00	0.00 0.00	5.234
281	GGTGTCTAAGTTTTTCTGCTGGATT	360 25	0 40.00	59.758	0.00	0.00 0.00	5.242
282	AGCATGCGGTATACTGTCTCTATA	188 24	0 41.67	58.746	13.01	0.00 0.00	5.254
283	TAAGTTTTTCTGCTGGATTCAACGG	354 25	0 40.00	60.278	0.00	0.00 0.00	5.278
284	TCGTCGTTTTCATTAAGGTGTC	377 23	0 39.13	57.667	0.00	0.00 0.00	5.333
285	GCATGCGGTATACTGTCTCTATAC	187 24	0 45.83	58.666	3.72	0.00 0.00	5.334
286	CGTCGTTTTCATTAAGGTGTCT	376 23	0 39.13	57.661	0.00	0.00 0.00	5.339
287	TATAGTGCCCAGCTATTTTGTGGAA	402 25	0 40.00	60.340	0.00	0.00 0.00	5.340
288	TCTATAGTGCCCAGCTATTTTGTG	404 24	0 41.67	58.632	0.00	0.00 0.00	5.368
289	CTGGATTCAACGGTTTCTGGCAC	342 23	0 52.17	62.451	0.00	0.00 0.00	5.451
290	CCTCTATAGTGCCCAGCTATTTTGT	406 25	0 44.00	60.455	0.00	0.00 0.00	5.455
291	AGGTGTCTAAGTTTTTCTGCTGGAT	361 25	0 40.00	60.457	0.00	0.00 0.00	5.457
292	AGTTTTCTGCTGGATTCAACGGT	352 24	0 41.67	61.468	0.00	0.00 0.00	5.468
293	CTCTATAGTGCCCAGCTATTTTGTG	405 25	0 44.00	59.532	0.00	0.00 0.00	5.468
294	TGGAATCGTCGTTTTTCATTAAGGT	382 25	0 36.00	59.529	7.33	0.00 0.00	5.471
295	ACTGTCTTGCAATATACACAGGTT	112 24	0 37.50	58.439	18.84	4.80 0.00	5.561
296	GCATGCGGTATACTGTCTCTATA	187 23	0 43.48	57.437	3.72	0.00 0.00	5.563
297	ATTTTGTGGAATCGTCGTTTTTCA	388 24	0 33.33	58.417	7.33	0.88 0.00	5.583
298	TTTTGTGGAATCGTCGTTTTTCAT	387 24	0 33.33	58.417	1.86	0.00 0.00	5.583
299	TTTGTGGAATCGTCGTTTTTCATT	386 24	0 33.33	58.417	1.86	0.00 0.00	5.583
300	GTCTCCATACACAGAGTCTGAAT	263 23	0 43.48		13.25	4.49 35.18	5.591
301	CTCTATAGTGCCCAGCTATTTTGT	405 24	0 41.67		0.00	0.00 0.00	5.615
302	TGTAAGTTCCAATACTGTCTTGCA	125 24	0 37.50		0.00	0.00 0.00	5.616
	CAGCTATTTTGTGGAATCGTCGTTT	393 25	0 40.00		0.00	0.00 0.00	5.617
	TGCGGTATACTGTCTCTATACACT		0 41.67		9.90	0.00 31.23	
	GCTATTTTGTGGAATCGTCGTTTTT	391 25	0 36.00		0.00	0.00 0.00	5.628
306	TCTATAGTGCCCAGCTATTTTGT	404 23	0 39.13		0.00	0.00 0.00	5.630
	AAGGTGTCTAAGTTTTTCTGCTGGA	362 25	0 40.00		0.00	0.00 0.00	5.631
	CTTGTGTTTCTCTGCGTCGTTGG	471 23	0 52.17		0.00	0.00 0.00	5.649
	AGTGCCCAGCTATTTTGTGGAATC	399 24	0 45.83		0.00	0.00 0.00	5.650
	GCAGCATGCGGTATACTGTCTCT	190 23	0 52.17			0.00 37.60	5.652
	CTATAGTGCCCAGCTATTTTGTG	403 23	0 43.48		0.00	0.00 0.00	5.680
	CCAGCTATTTTGTGGAATCGTCGT	394 24	0 45.83		0.00	0.00 0.00	5.681
	TCTATAGTGCCCAGCTATTTTGTGG	404 25	0 44.00		0.00	0.00 0.00	5.687
	CTATAGTGCCCAGCTATTTTGTGGA	403 25	0 44.00		0.00	0.00 0.00	5.687
	TTTCTCTGCGTCGTTGGAGTCTC	465 23	0 52.17		9.66	0.00 0.00	5.691
	TGGCCTCTATAGTGCCCAGCTAT	409 23	0 52.17			0.00 37.93	5.692
	ATCGTCGTTTTTCATTAAGGTGTCT	378 25	0 36.00		0.00	0.00 0.00	5.699
	CTGTAAGTTCCAATACTGTCTTGCA	126 25	0 40.00		0.00	0.00 0.00	5.701
	CCGCAGGCACCTTATTAATAAAT	320 23	0 39.13		9.15	0.00 46.14	5.706
	AAGGTGTCTAAGTTTTTCTGCTG	362 23	0 39.13		0.00	0.00 0.00	5.758
<b>3</b> ∠1	TAAGTTTTCTGCTGGATTCAACG	354 24	0 37.50	JO. 242	0.00	0.00 0.00	5.758

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322	TAAGGTGTCTAAGTTTTTCTGCTGG	363 25	0	40.00	59.237	0.00	0.00 0	.00	5.763
323	TACTTGTGTTTCTCTGCGTCGTTG	473 24	0	45.83	61.777	0.00	0.00 0	.00	5.777
324	ACCTCTGTAAGTTCCAATACTGTCT	130 25	0	40.00	59.219	0.00	0.00 0	.00	5.781
325	TTATACTTGTGTTTCTCTGCGTCGT	476 25	0	40.00	60.791	0.00	0.00 0	.00	5.791
326	TATACTTGTGTTTCTCTGCGTCGTT	475 25	0	40.00	60.791	0.00	0.00 0	.00	5.791
327	GTGGAATCGTCGTTTTTCATTAAGG	383 25	0	40.00	59.207	1.86	0.00 0	.00	5.793
328	GGAATCGTCGTTTTTCATTAAGGTG	381 25	0	40.00	59.207	7.33	0.00 0	.00	5.793
329	TTCTGCTGGATTCAACGGTTTCTG	347 24	0	45.83	61.808	0.00	0.00 0	.00	5.808
330	CTAAGTTTTCTGCTGGATTCAACG	355 25	0	40.00	59.147	0.00	0.00 0	.00	5.853
331	ATAGTGCCCAGCTATTTTGTGGAAT	401 25	0	40.00	60.870	0.00	0.00 0	.00	5.870
332	TCTATACATTTATGGCATGCAGC	208 23	0	39.13	57.115	7.99	0.11 44	.65	5.885
333	CCGCAGGCACCTTATTAATAAATTG	320 25			59.080	9.79	9.79 46	.14	5.920
334	ATCGTCGTTTTTCATTAAGGTGTC	378 24	0	37.50	58.076	0.00	0.00 0	.00	5.924
335	TGTGGAATCGTCGTTTTTCATTA	384 23	0	34.78	57.074	1.86	0.00 0	.00	5.926
336	TCTGTAAGTTCCAATACTGTCTTGC	127 25	0	40.00	59.068	0.75		.00	5.932
337		382 24			58.066	7.33		.00	5.934
338		466 23			62.935	4.98		.00	5.935
339		381 24			58.065	7.33		.00	5.935
340		378 23			57.065	0.00		.00	5.935
341		184 23			57.063	9.90		.23	5.937
342		359 23			57.051	0.00		.00	5.949
343		350 25			60.963	0.00		.00	5.963
344		359 26			60.017	0.00		.00	6.017
345		387 25			58.963	1.86		.00	6.037
346		130 24			57.961	0.00		.00	6.039
347		125 25			58.941	0.00		.00	6.059
348		358 25			58.939	0.00		.00	6.061
349		320 24			57.916	9.15		.14	6.084
350		408 24			62.091		0.00 40		6.091
	CTCTGTAAGTTCCAATACTGTCTTGC				59.909			.00	6.091
	CATGCGGTATACTGTCTCTATACACT	186 26			59.850	9.90	0.00 0		6.150
	CTGTAAGTTCCAATACTGTCTTGC				57.829			.00	
		126 24				0.75			6.171 6.183
	GCTATTTGTGGAATCGTCGTTTTTC	391 26			60.183 58.812	0.00		.00	
	TGTCTCCATACACAGAGTCTGAATA	264 25					9.54 35		6.188
	GTTTTTCTGCTGGATTCAACGGTTT	351 25			61.191	0.00		.00	6.191
	AATCGTCGTTTTTCATTAAGGTGTCT	379 26			59.796	0.00		.00	6.204
	CTGTAAGTTCCAATACTGTCTTGCAA	126 26			59.795	0.00		.00	6.205
	ATTTTGTGGAATCGTCGTTTTTCAT	388 25			58.787	1.86		.00	6.213
360		377 25			58.785	0.00		.00	6.215
	ATGCGGTATACTGTCTCTATACACT	185 25			58.761	9.90	0.00 31		6.239
	CCCAGCTATTTTGTGGAATCGTCG	395 24			62.256	0.00		.00	6.256
	TTAAGGTGTCTAAGTTTTTCTGCTGG	364 26			59.738	0.00		.00	6.262
	ACCTCTGTAAGTTCCAATACTGTCTT	130 26			59.730	0.00		.00	6.270
	CCTCTGTAAGTTCCAATACTGTCT	129 24			57.712	0.00		.00	6.288
	TCTAAGTTTTTCTGCTGGATTCAACG	356 26			60.293	0.00		.00	6.293
367	TACTGTCTTGCAATATACACAGGT	113 24			57.706			.00	6.294
368	TTGTGGAATCGTCGTTTTTCATTA	385 24	0	33.33	57.698	1.86	0.00 0	.00	6.302

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Pri	ners data								
369	TGTGGAATCGTCGTTTTTCATTAA	384 2	24 0	33.33	57.698	1.86	0.00	0.00	6.302
370	AATCGTCGTTTTTCATTAAGGTGT	379 2	24 0	33.33	57.692	0.00	0.00	0.00	6.308
371	CATGCGGTATACTGTCTCTATACAC	186 2	25 0	44.00	58.684	9.90	0.00 3	1.23	6.316
372	GTGTCTAAGTTTTTCTGCTGGATT	359 2	24 0	37.50	57.681	0.00	0.00	0.00	6.319
373	GCAGCATGCGGTATACTGTCTCTA	190 2	24 0	50.00	62.339	14.57	0.00 3	7.60	6.339
374	TGTGGAATCGTCGTTTTTCATTAAG	384 2	25 0	36.00	58.633	0.15	0.00	0.00	6.367
375	GAATCGTCGTTTTTCATTAAGGTGT	380 2	25 0	36.00	58.632	2.12	0.00	0.00	6.368
376	AATCGTCGTTTTTCATTAAGGTGTC	379 2	25 0	36.00	58.632	0.00	0.00	0.00	6.368
377	GTGTCTAAGTTTTTCTGCTGGATTC	359 2	25 0	40.00	58.625	0.00	0.00	0.00	6.375
378	TAGTGCCCAGCTATTTTGTGGAATC	400 2			61.382	0.00	0.00	0.00	6.382
379	TAAGGTGTCTAAGTTTTTCTGCTGGA	363 2	26 0	38.46	60.409	0.00	0.00	0.00	6.409
380	ATCTATACATTTATGGCATGCAGC	209 2	24 0	37.50	57.553	7.99	0.11 4	4.65	6.447
381	CGTCGTTTTCATTAAGGTGTCTA				57.544	0.00		0.00	6.456
382	CCGCAGGCACCTTATTAATAAATTGT				60.461			6.14	6.461
383	TCTGTAAGTTCCAATACTGTCTTGCA		-		60.464	0.00		0.00	6.464
384	CATGCGGTATACTGTCTCTATACA				57.504	9.90		1.23	6.496
385	ATGCGGTATACTGTCTCTATACAC				57.502	9.90		1.23	6.498
386	GCCCAGCTATTTTGTGGAATCGTC				62.502	0.00		0.00	6.502
387	CGCAGGCACCTTATTAATAAATTGT				58.496	13.46		0.00	6.504
388	AGCTATTTTGTGGAATCGTCGTTTTT				60.514	0.00	0.00	0.00	6.514
389	GAATCGTCGTTTTTCATTAAGGTGTC				59.481	2.12		0.00	6.519
390	GCCTCTATAGTGCCCAGCTATTTTG				61.544	5.93	0.00	0.00	6.544
					59.454				6.546
391	TGTCTAAGTTTTTCTGCTGGATTCAA					0.00	0.00	0.00	
392	GTAAGTTCCAATACTGTCTTGCAA				57.446	0.00	0.00	0.00	6.554
393	TGTCTAAGTTTTTCTGCTGGATTC				57.440	0.00		0.00	6.560
394	GTCTAAGTTTTTCTGCTGGATTCA				57.440	0.00	0.00	0.00	6.560
395	TGTGGAATCGTCGTTTTTCATTAAGG				60.567	0.15	0.00	0.00	6.567
396	TGGAATCGTCGTTTTTCATTAAGGTG				60.567	7.33	0.00	0.00	6.567
397	GTGGAATCGTCGTTTTTCATTAAGGT				60.567	1.86		0.00	6.567
	GGAATCGTCGTTTTTCATTAAGGTGT	381 2			60.567	7.33		0.00	6.567
	GGTGTCTAAGTTTTTCTGCTGGATTC	360 2			60.569	0.00		0.00	6.569
	CCTCTGTAAGTTCCAATACTGTCTTG	129 2			59.408	0.00		0.00	6.592
	TATAGTGCCCAGCTATTTTGTGGAAT	402 2			60.637	0.00		0.00	6.637
	TGCGGTATACTGTCTCTATACACTAC	184 2			59.352	9.90	0.00 3		6.648
	GCGGTATACTGTCTCTATACACTACA	183 2			59.352	6.27	0.00 3		6.648
	TGTCTCCATACACAGAGTCTGAATAA	264 2			59.336	7.70	0.00 3		6.664
	TTTTCTGCTGGATTCAACGGTTTCT	349 2			61.669	0.00		0.00	6.669
406	ATTTTGTGGAATCGTCGTTTTTCATT	388 2			59.300	1.86	0.00	0.00	6.700
407	TCGTCGTTTTTCATTAAGGTGTCTAA	377 2			59.299	0.00	0.00	0.00	6.701
408	GTCTCCATACACAGAGTCTGAATA	263 2	24 0	41.67	57.299	16.96	9.54 3	5.18	6.701
409	CCTCTGTAAGTTCCAATACTGTCTT	129 2	25 0	40.00	58.298	0.00	0.00	0.00	6.702
410	TACTGTCTTGCAATATACACAGGTT	113 2	25 0	36.00	58.293	18.84	8.21	0.00	6.707
411	ACTGTCTTGCAATATACACAGGTTA	112 2	25 0	36.00	58.293	18.84	0.00	0.00	6.707
412	TGTAAGTTCCAATACTGTCTTGCAAT	125 2	26 0	34.62	59.285	0.00	0.00	0.00	6.715
413	TATTTTGTGGAATCGTCGTTTTTCA	389 2	25 0	32.00	58.274	7.33	0.88	0.00	6.726
414	TTTGTGGAATCGTCGTTTTTCATTA	386 2	25 0	32.00	58.274	1.86	0.00	0.00	6.726
415	TTGTGGAATCGTCGTTTTTCATTAA	385 2	25 0	32.00	58.274	1.45	0.00	0.00	6.726

Pri	mers data									
416	GTGCCCAGCTATTTTGTGGAATCG	398	24	0	50.00	62.736	0.00	0.00	0.00	6.736
417	TGCGGTATACTGTCTCTATACACTA	184	25	0	40.00	58.238	9.90	0.00	31.23	6.762
418	GTGGAATCGTCGTTTTTCATTAAG	383	24	0	37.50	57.167	1.86	0.00	0.00	6.833
419	GAATCGTCGTTTTTCATTAAGGTG	380	24	0	37.50	57.167	2.12	0.00	0.00	6.833
420	CTATTTTGTGGAATCGTCGTTTTTCA	390	26	0	34.62	59.146	7.33	0.88	0.00	6.854
421	TTGTGGAATCGTCGTTTTTCATTAAG	385	26	0	34.62	59.146	2.73	0.00	0.00	6.854
422	GTCTAAGTTTTTCTGCTGGATTCAAC	357	26	0	38.46	59.142	0.00	0.00	0.00	6.858
423	AATCTATACATTTATGGCATGCAGC	210	25	0	36.00	58.140	7.99	0.11	44.65	6.860
424	TAAGGTGTCTAAGTTTTTCTGCTG	363	24	0	37.50	57.139	0.00	0.00	0.00	6.861
425	ATCGTCGTTTTTCATTAAGGTGTCTA	378	26	0	34.62	59.130	0.00	0.00	0.00	6.870
426	CGTCGTTTTCATTAAGGTGTCTAA	376	25	0	36.00	58.120	0.00	0.00	0.00	6.880
427	ATACTGTCTTGCAATATACACAGGT	114	25			58.111		5.21	0.00	6.889
428	CAGCATGCGGTATACTGTCTCTATAC	189	26			60.894	6.85	0.00	0.00	6.894
429	AAGTTTTTCTGCTGGATTCAACGGT	353	25			61.899	0.00	0.00	0.00	6.899
430	AGTTTTCTGCTGGATTCAACGGTT	352				61.899	0.00	0.00	0.00	6.899
431	AAGGTGTCTAAGTTTTTCTGCTGGAT	362				60.917	0.00	0.00	0.00	6.917
	AGGTGTCTAAGTTTTTCTGCTGGATT	361				60.917	0.00	0.00	0.00	6.917
433	TACCTCTGTAAGTTCCAATACTGTCT	131	26			59.048	0.00	0.00	0.00	6.952
434	GTCTAAGTTTTTCTGCTGGATTCAA	357	25	-		58.030	0.00	0.00	0.00	6.970
435	TCTAAGTTTTCTGCTGGATTCAAC	356	25			58.030	0.00	0.00	0.00	6.970
436	CGCAGGCACCTTATTAATAAATTG	319	24			57.009	9.79	9.79	0.00	6.991
437	CGTCGTTTTTCATTAAGGTGTCTAAG	376	26			58.992	0.00	0.00	0.00	7.008
438	ATACTTGTGTTTCTCTGCGTCGTTG	474				62.020	0.00	0.00	0.00	7.008
439	ATTAAGGTGTCTAAGTTTTTCTGCTGG	365	27			60.040	0.00		0.00	7.020
440	CAGCTATTTTGTGGAATCGTCGTTTT	393	26			61.055	0.00	0.00	0.00	7.040
441						61.055				
		355 183	26 25			57.941	0.00 6.27	0.00	0.00	7.057 7.059
442	GCGGTATACTGTCTCTATACACTAC		25					0.00	31.23	
443	CTGTAAGTTCCAATACTGTCTTGCAAT	126	27			60.095	0.00	0.00	0.00	7.095
	TCGTCGTTTTTCATTAAGGTGTCTAAG	377				60.096	0.00	0.00	0.00	7.096
	CCAGCTATTTTGTGGAATCGTCGTT						0.00			7.096
	GTCTCCATACACAGAGTCTGAATAA	263				57.898	7.70		35.18	7.102
	GCATGCGGTATACTGTCTCTATACAC	187				61.111			31.23	7.111
	CTATAGTGCCCAGCTATTTTGTGGAA	403				61.135	0.00	0.00	0.00	7.135
	GTAAGTTCCAATACTGTCTTGCAAT	124				57.854	0.24	0.00	0.00	7.146
	TACCTCTGTAAGTTCCAATACTGTC	131				57.833	0.00	0.00	0.00	7.167
	GCGGTATACTGTCTCTATACACTACAA	183				59.828	6.27		31.23	7.172
	TTTTGTGGAATCGTCGTTTTTCATTA	387				58.806	1.86	0.00	0.00	7.194
	TTTGTGGAATCGTCGTTTTTCATTAA	386				58.806	1.62	0.00	0.00	7.194
454	TTTCTGCTGGATTCAACGGTTTCTG	348				62.221	0.00	0.00	0.00	7.221
455	TTATACTTGTGTTTCTCTGCGTCGTT	476	26			61.227		0.00	0.00	7.227
	TGAATGCAAATTCAAATACCTCTGT	147				57.769			37.59	7.231
457	AGCATGCGGTATACTGTCTCTATACA	188	26			61.247	13.01	0.00	31.23	7.247
458	CTATTTTGTGGAATCGTCGTTTTTC	390	25	0	36.00	57.752	0.00	0.00	0.00	7.248
459	CCAATACTGTCTTGCAATATACACAGG	117	27			60.255		8.97	0.00	7.255
460	CCGCAGGCACCTTATTAATAAATTGTA	320	27	0	37.04	60.255	13.46	0.00	46.14	7.255
461	TTAAGGTGTCTAAGTTTTTCTGCTG	364	25	0	36.00	57.741	0.00	0.00	0.00	7.259
462	CAATACTGTCTTGCAATATACACAGGT	116	27	0	37.04	59.719	11.41	5.21	0.00	7.281

Primers data					
463 CGTCGTTTTTCATTAAGGTGTCTAAGT	376 27	0 37.04 60.307	0.00 0.00	0.00	7.307
464 AAATCTATACATTTATGGCATGCAGC	211 26	0 34.62 58.682	7.99 0.13	L 44.65	7.318
465 CATGCGGTATACTGTCTCTATACACTA	186 27	0 40.74 59.666	9.90 0.00	31.23	7.334
466 ATGCGGTATACTGTCTCTATACACTAC	185 27	0 40.74 59.666	9.90 0.00	31.23	7.334
467 TCCAATACTGTCTTGCAATATACACA	118 26	0 34.62 58.665	9.08 3.86	5 0.00	7.335
468 AATACTGTCTTGCAATATACACAGGT	115 26	0 34.62 58.661			7.339
469 ATACTGTCTTGCAATATACACAGGTT	114 26	0 34.62 58.661			7.339
	112 26				7.339
		0 34.62 58.661			
471 TGTCTCCATACACAGAGTCTGAATAAT	264 27	0 37.04 59.657		35.18	7.343
472 TCTCCATACACAGAGTCTGAATAATGT	262 27	0 37.04 59.657			7.343
473 TCCATACACAGAGTCTGAATAATGTCT	260 27	0 37.04 59.657	16.73 5.64	1 35.18	7.343
474 TATTTTGTGGAATCGTCGTTTTTCAT	389 26	0 30.77 58.636	1.86 0.00	0.00	7.364
475 TTTGTGGAATCGTCGTTTTTCATTAAG	386 27	0 33.33 59.623	2.89 0.00	0.00	7.377
476 AATCGTCGTTTTTCATTAAGGTGTCTA	379 27	0 33.33 59.614	0.00 0.00	0.00	7.386
477 ATCGTCGTTTTTCATTAAGGTGTCTAA	378 27	0 33.33 59.614	0.00 0.00	0.00	7.386
478 ATGCGGTATACTGTCTCTATACACTA	185 26	0 38.46 58.608	9.90 0.00	31.23	7.392
479 TCCATACACAGAGTCTGAATAATGT	260 25	0 36.00 57.576	16.73 13.16	35.18	7.424
480 TACCTCTGTAAGTTCCAATACTGTCTT	131 27	0 37.04 59.546	0.00 0.00	0.00	7.454
481 CCTCTATAGTGCCCAGCTATTTTGTG	406 26	0 46.15 61.459	0.00 0.00		7.459
482 CTCTATAGTGCCCAGCTATTTTGTGG	405 26	0 46.15 61.459	0.00 0.00		7.459
483 GTGTCTAAGTTTTTCTGCTGGATTCAA	359 27	0 37.04 60.469	0.00 0.00		7.469
484 TGTCTAAGTTTTTCTGCTGGATTCAAC		0 37.04 60.469	0.00 0.00		7.469
485 TCCATACACAGAGTCTGAATAATGTC	260 26	0 38.46 58.507		35.18	7.493
486 TCCAATACTGTCTTGCAATATACACAG	118 27	0 37.04 59.504			7.496
487 CTCCATACACAGAGTCTGAATAATGT	261 26	0 38.46 58.504		35.18	7.496
488 CCATACACAGAGTCTGAATAATGTCT	259 26	0 38.46 58.504	16.73 5.64	1 35.18	7.496
489 GGCCTCTATAGTGCCCAGCTATTTT	408 25	0 48.00 62.503	19.28 0.00	34.65	7.503
490 CTATTTTGTGGAATCGTCGTTTTTCAT	390 27	0 33.33 59.463	1.86 0.00	0.00	7.537
491 CCAATACTGTCTTGCAATATACACA	117 25	0 36.00 57.448	9.08 3.86	0.00	7.552
492 GCAGCATGCGGTATACTGTCTCTAT	190 25	0 48.00 62.567	14.57 0.00	37.60	7.567
493 GAATCGTCGTTTTTCATTAAGGTGTCT	380 27	0 37.04 60.571	2.12 0.00	0.00	7.571
494 CTCTGTAAGTTCCAATACTGTCTTG	128 25	0 40.00 57.412	0.00 0.00	0.00	7.588
495 ATACCTCTGTAAGTTCCAATACTGTCT	132 27	0 37.04 59.380	0.00 0.00	0.00	7.620
496 TAAGTTTTCTGCTGGATTCAACGGT	354 26	0 38.46 61.633		0.00	7.633
497 CCAATACTGTCTTGCAATATACACAG	117 26	0 38.46 58.366		1 0.00	7.634
498 CAATACTGTCTTGCAATATACACAGG	116 26	0 38.46 58.366			7.634
499 ATAGTGCCCAGCTATTTTGTGGAATC	401 26	0 42.31 61.638	0.00 0.00		7.631
500 CGCAGGCACCTTATTAATAATTGTA	319 26	0 34.62 58.356			7.644
501 GTCTCCATACACAGAGTCTGAATAATG	263 27	0 40.74 59.349			7.651
502 CTCCATACACAGAGTCTGAATAATGTC	261 27	0 40.74 59.349			7.651
503 TTGAATGCAAATTCAAATACCTCTGT	148 26	0 30.77 58.331		44.64	7.669
504 TGCGGTATACTGTCTCTATACACTACA	184 27	0 40.74 60.688	9.90 0.00	31.23	7.688
505 TAAGGTGTCTAAGTTTTTCTGCTGGAT	363 27	0 37.04 60.692	0.00 0.00	0.00	7.692
506 TTTTGTGGAATCGTCGTTTTTCATTAA	387 27	0 29.63 59.299	1.62 0.00	0.00	7.701
507 CCATACACAGAGTCTGAATAATGTC	259 25	0 40.00 57.294	16.73 0.00	35.18	7.706
508 TCATTAAGGTGTCTAAGTTTTTCTGCT	367 27	0 33.33 59.280	0.00 0.00	0.00	7.720
509 TCTCCATACACAGAGTCTGAATAATG	262 26	0 38.46 58.280	15.00 12.15	35.18	7.720

PLII	mers data									
510	GTCTCCATACACAGAGTCTGAATAAT	263	26	0	38.46	58.279	8.51	0.00	35.18	7.721
511	ATACCTCTGTAAGTTCCAATACTGT	132	25	0	36.00	57.265	0.00	0.00	0.00	7.735
512	ACCTCTGTAAGTTCCAATACTGTCTTG	130	27	0	40.74	60.741	0.00	0.00	0.00	7.741
513	GCAAATTCAAATACCTCTGTAAGTTCC	142	27	0	37.04	59.245	0.00	0.00	0.00	7.755
514	TATACTTGTGTTTCTCTGCGTCGTTG	475	26	0	42.31	61.756	0.00	0.00	0.00	7.756
515	TGCAAATTCAAATACCTCTGTAAGT	143	25			57.241	0.00	0.00	0.00	7.759
	ATACCTCTGTAAGTTCCAATACTGTC	132				58.217	0.00	0.00	0.00	7.783
517	TCCAATACTGTCTTGCAATATACAC	118	25			57.214	9.08	0.02	0.00	7.786
518	AATACTGTCTTGCAATATACACAGG	115	25			57.208		8.89	0.00	7.792
519	CTGTCTTGCAATATACACAGGTTAT	111	25	0	36.00	57.208	11.78	0.00	0.00	7.792
520	CATTAAGGTGTCTAAGTTTTTCTGCTG	366	27	0	37.04	59.195	0.00	0.00	0.00	7.805
521	AAAATCTATACATTTATGGCATGCAGC	212	27	0	33.33	59.184	7.99	0.11	44.65	7.816
522	TCTATAGTGCCCAGCTATTTTGTGGA	404	26	0	42.31	61.826	0.00	0.00	0.00	7.826
523	TTCCAATACTGTCTTGCAATATACACA	119	27	0	33.33	59.173	9.08	3.86	0.00	7.827
524	AATACTGTCTTGCAATATACACAGGTT	115	27	0	33.33	59.171	13.29	8.25	0.00	7.829
525	ACTGTCTTGCAATATACACAGGTTATT	112	27	0	33.33	59.171	18.84	3.84	0.00	7.829
526	TACTGTCTTGCAATATACACAGGTTA	113	26	0	34.62	58.157	18.84	7.57	0.00	7.843
527	TTAAGGTGTCTAAGTTTTTCTGCTGGA	364	2.7	0	37.04	60.853	0.00	0.00	0.00	7.853
528	GTCGTTTTCATTAAGGTGTCTAAGT	375	26			58.141	0.00	0.00	0.00	7.859
529	TATTTGTGGAATCGTCGTTTTCATT	389	27			59.137	0.00	0.00	0.00	7.863
						59.137				7.863
	ATTTTGTGGAATCGTCGTTTTTCATTA		27				0.00	0.00	0.00	
	TCATTAAGGTGTCTAAGTTTTTCTGC	367	26			58.128	0.00	0.00	0.00	7.872
	CATTAAGGTGTCTAAGTTTTTCTGCT	366	26			58.124	0.00	0.00	0.00	7.876
533	ATTAAGGTGTCTAAGTTTTTCTGCTG	365	26			58.124	0.00	0.00	0.00	7.876
534	TGTAAGTTCCAATACTGTCTTGCAATA	125	27	0	33.33	59.119	0.00	0.00	0.00	7.881
535	TCTGTAAGTTCCAATACTGTCTTGCAA	127	27	0	37.04	60.904	0.00	0.00	0.00	7.904
536	TCTCCATACACAGAGTCTGAATAAT	262	25	0	36.00	57.095	8.51	0.00	35.18	7.905
537	GTTTTTCTGCTGGATTCAACGGTTTC	351	26	0	42.31	61.921	0.00	0.00	0.00	7.921
538	CTCCATACACAGAGTCTGAATAATG	261	25	0	40.00	57.057	15.00	12.15	35.18	7.943
539	AAGTTCCAATACTGTCTTGCAATAT	122	25	0	32.00	57.055	0.00	0.00	0.00	7.945
540	CCATACACAGAGTCTGAATAATGTCTT	259	27	0	37.04	59.016	16.73	4.96	35.18	7.984
541	TTGTGGAATCGTCGTTTTTCATTAAGG	385	27	0	37.04	60.994	2.73	0.00	0.00	7.994
542	AGTTCCAATACTGTCTTGCAATATACA	121		0	33.33	58.953	9.08	9.08	0.00	8.047
	ATACACAGGTTATTTCTATGTCTTGCA		27			58.953	0.00	0.00	0.00	8.047
	TTTTCTGCTGGATTCAACGGTTTCT	350				62.076	0.00	0.00	0.00	8.076
	GTTCCAATACTGTCTTGCAATATACAC	120				58.877	9.08	0.02	0.00	8.123
	AATACCTCTGTAAGTTCCAATACTGT	133				57.853	0.00	0.00	0.00	8.147
	TGCAAATTCAAATACCTCTGTAAGTT	143				57.823	0.00	0.00	0.00	8.177
548	GTTCCAATACTGTCTTGCAATATACA	120	26	0	34.62	57.791	9.08	9.08	0.00	8.209
549	TTCCAATACTGTCTTGCAATATACAC	119	26	0	34.62	57.791	9.08	0.02	0.00	8.209
550	CTGTCTTGCAATATACACAGGTTATT	111	26	0	34.62	57.786	11.78	0.00	0.00	8.214
551	CTCTGTAAGTTCCAATACTGTCTTGCA	128	27	0	40.74	61.218	0.00	0.00	0.00	8.218
552	GTCTAAGTTTTTCTGCTGGATTCAACG	357	27	0	40.74	61.250	0.00	0.00	0.00	8.250
553	AGCTATTTTGTGGAATCGTCGTTTTTC	392	27	0	37.04	61.253	0.00	0.00	0.00	8.253
554	AATACCTCTGTAAGTTCCAATACTGTC	133	27	0	37.04	58.741	0.00	0.00	0.00	8.259
555	GTAAGTTCCAATACTGTCTTGCAATA	124	26	0	34.62	57.736	0.00	0.00	0.00	8.264
	GCAGCATGCGGTATACTGTCTCTATA	190				62.278			37.60	8.278
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Primers data					
557 CGCAGGCACCTTATTAATAAATTGTAT	319 27	0 33.33 58.703	13.46	4.54 0.00	8.297
558 TGCAAATTCAAATACCTCTGTAAGTTC	143 27	0 33.33 58.703	0.00	0.00 0.00	8.297
559 AAGTTTTTCTGCTGGATTCAACGGTT	353 26	0 38.46 62.298	0.00	0.00 0.00	8.298
560 AGTTTTTCTGCTGGATTCAACGGTTT	352 26	0 38.46 62.298	0.00	0.00 0.00	8.298
561 GGAATCGTCGTTTTTCATTAAGGTGTC	381 27	0 40.74 61.299	7.33	0.00 0.00	8.299
562 GTCGTTTTTCATTAAGGTGTCTAAGTT	375 27	0 33.33 58.660	0.00	0.00 0.00	8.340
563 AAGGTGTCTAAGTTTTTCTGCTGGATT	362 27	0 37.04 61.343	0.00	0.00 0.00	8.343
564 TGAATGCAAATTCAAATACCTCTGTA	147 26	0 30.77 57.653	2.83	0.00 37.59	8.347
565 TTCATTAAGGTGTCTAAGTTTTTCTGC	368 27	0 33.33 58.650	0.00	0.00 0.00	8.350
566 ATGCAAATTCAAATACCTCTGTAAGT	144 26	0 30.77 57.647	0.00	0.00 0.00	8.353
567 TATAGTGCCCAGCTATTTTGTGGAATC	402 27	0 40.74 61.389	0.00	0.00 0.00	8.389
568 CTATAGTGCCCAGCTATTTTGTGGAAT	403 27	0 40.74 61.392	0.00	0.00 0.00	8.392
569 AGTTCCAATACTGTCTTGCAATATAC	121 26	0 34.62 57.561	0.98	0.98 0.00	8.439
570 ATACACAGGTTATTTCTATGTCTTGC	99 26	0 34.62 57.561	0.00	0.00 0.00	8.439
571 TCGTTTTCATTAAGGTGTCTAAGTT	374 26	0 30.77 57.559	0.00	0.00 0.00	8.441
572 GCTATTTTGTGGAATCGTCGTTTTTCA	391 27	0 37.04 61.460	7.33	0.88 0.00	8.460
573 CAGCTATTTTGTGGAATCGTCGTTTTT	393 27	0 37.04 61.460	0.00		8.462
574 CCAGCTATTTTGTGGAATCGTCGTTT	394 26	0 42.31 62.481	0.00	0.00 0.00	8.481
575 ATACTGTCTTGCAATATACACAGGTTA	114 27	0 33.33 58.517	4.29	0.00 0.00	8.483
576 TACTGTCTTGCAATATACACAGGTTAT	113 27	0 33.33 58.517	3.21	0.00 0.00	8.483
577 GTGGAATCGTCGTTTTTCATTAAGGTG	383 27	0 40.74 61.507	1.86	0.00 0.00	8.507
578 TGTCTTGCAATATACACAGGTTATTT	110 26	0 30.77 57.426	9.60	0.00 0.00	8.574
579 CGGTATACTGTCTCTATACACTACAA	182 26	0 38.46 57.401	3.31	0.00 31.23	8.599
580 TTTTCTGCTGGATTCAACGGTTTCTG	349 26	0 42.31 62.603	0.00	0.00 0.00	8.603
581 AAATACCTCTGTAAGTTCCAATACTGT	134 27	0 33.33 58.397	0.00	0.00 0.00	8.603
582 AGGTGTCTAAGTTTTTCTGCTGGATTC	361 27	0 40.74 61.653	0.00	0.00 0.00	8.653
583 TGTCTTGCAATATACACAGGTTATTTC	110 27	0 33.33 58.326	9.60	0.00 0.00	8.674
584 GCAAATTCAAATACCTCTGTAAGTTC	142 26	0 34.62 57.325	0.00	0.00 0.00	8.675
585 CTGTCTTGCAATATACACAGGTTATTT	111 27	0 33.33 58.323	11.78	0.00 0.00	8.677
586 CCTCTGTAAGTTCCAATACTGTCTTGC	129 27	0 44.44 61.743	0.75	0.00 0.00	8.743
587 CAAATTCAAATACCTCTGTAAGTTCCA	141 27	0 33.33 58.207	0.00	0.00 0.00	8.793
588 TTGAATGCAAATTCAAATACCTCTGTA	148 27	0 29.63 58.199	0.00	0.00 44.64	8.801
589 TGAATGCAAATTCAAATACCTCTGTAA	147 27	0 29.63 58.199	2.83	0.00 37.59	8.801
590 AATGCAAATTCAAATACCTCTGTAAGT	145 27	0 29.63 58.194	0.00	0.00 0.00	8.806
591 ATGCAAATTCAAATACCTCTGTAAGTT	144 27	0 29.63 58.194	0.00	0.00 0.00	8.806
592 TCAAATACCTCTGTAAGTTCCAATACT	136 27	0 33.33 58.177	0.00	0.00 0.00	8.823
593 TGTGGAATCGTCGTTTTTCATTAAGGT	384 27	0 37.04 61.856	0.15	0.00 0.00	8.856
594 TGGAATCGTCGTTTTTCATTAAGGTGT	382 27	0 37.04 61.856	7.33	0.00 0.00	8.856
595 GGTGTCTAAGTTTTTCTGCTGGATTCA	360 27	0 40.74 61.863	0.00	0.00 0.00	8.863
596 GCCTCTATAGTGCCCAGCTATTTTGT	407 26	0 46.15 62.869	5.93	0.00 0.00	8.869
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598 GTAAGTTCCAATACTGTCTTGCAATAT	124 27	0 33.33 58.106	0.00	0.00 0.00	8.894
599 AAGTTCCAATACTGTCTTGCAATATAC	122 27	0 33.33 58.106	0.98	0.98 0.00	8.894
600 GTCTTGCAATATACACAGGTTATTTCT	109 27	0 33.33 58.106	2.76	0.00 0.00	8.894
601 TCGTTTTCATTAAGGTGTCTAAGTTT	374 27	0 29.63 58.104	0.00	0.00 0.00	8.896
602 AAATTCAAATACCTCTGTAAGTTCCA	140 26	0 30.77 57.066	0.00	0.00 0.00	8.934
603 AATTCAAATACCTCTGTAAGTTCCAA	139 26	0 30.77 57.066	0.00	0.00 0.00	8.934

604	CATACACAGAGTCTGAATAATGTCTT	258	26	0	34.62	57.061	14.59	2.70	35.18	8.939
605	TAAGTTTTTCTGCTGGATTCAACGGTT	354	27	0	37.04	62.027	0.00	0.00	0.00	9.027
606	CGGTATACTGTCTCTATACACTACAAA	182	27	0	37.04	57.950	3.31	0.00	31.23	9.050
607	ACCCAGTGTTAGTTAGTTTTCTAATG	291	27	0	33.33	57.936	0.00	0.00	40.23	9.064
608	CCCAGTGTTAGTTAGTTTTTCTAATGT	290	27	0	33.33	57.936	0.00	0.00	40.23	9.064
609	TCTAAGTTTTTCTGCTGGATTCAACGG	356	27	0	40.74	62.117	0.00	0.00	0.00	9.117
610	TTATACTTGTGTTTCTCTGCGTCGTTG	476	27	0	40.74	62.136	0.00	0.00	0.00	9.136
611	CAGCATGCGGTATACTGTCTCTATACA	189	27	0	44.44	62.172	6.85	0.00	31.23	9.172
612	AGCATGCGGTATACTGTCTCTATACAC	188	27	0	44.44	62.173	13.01	0.00	31.23	9.173
613	GCATGCGGTATACTGTCTCTATACACT	187	27	0	44.44	62.173	3.72	0.00	31.23	9.173
614	TCTATAGTGCCCAGCTATTTTGTGGAA	404	27	0	40.74	62.220	0.00	0.00	0.00	9.220
615	GAATGCAAATTCAAATACCTCTGTAAG	146	27	0	33.33	57.706	0.00	0.00	0.00	9.294
616	CTAAGTTTTTCTGCTGGATTCAACGGT	355	27	0	40.74	62.330	0.00	0.00	0.00	9.330
617	AAATTCAAATACCTCTGTAAGTTCCAA	140	27	0	29.63	57.638	0.00	0.00	0.00	9.362
618	TTCAAATACCTCTGTAAGTTCCAATAC	137	27	0	33.33	57.555	0.00	0.00	0.00	9.445
619	CGTTTTTCATTAAGGTGTCTAAGTTTT	373	27	0	29.63	57.489	0.00	0.00	0.00	9.511
620	CTCTATAGTGCCCAGCTATTTTGTGGA	405	27	0	44.44	62.525	0.00	0.00	0.00	9.525
621	AATTCAAATACCTCTGTAAGTTCCAAT	139	27	0	29.63	57.466	0.00	0.00	0.00	9.534
622	TATACACAGGTTATTTCTATGTCTTGC	100	27	0	33.33	57.458	0.00	0.00	0.00	9.542
623	AACCCAGTGTTAGTTAGTTTTCTAAT	292	27	0	29.63	57.363	0.00	0.00	36.90	9.637
624	AAGTTTTTCTGCTGGATTCAACGGTTT	353	27	0	37.04	62.668	0.00	0.00	0.00	9.668
625	AAAAATCTATACATTTATGGCATGCAG	213	27	0	29.63	57.299	7.99	0.00	0.00	9.701
626	CCAGCTATTTTGTGGAATCGTCGTTTT	394	27	0	40.74	62.838	0.00	0.00	0.00	9.838
627	TGCAATATACACAGGTTATTTCTATGT	105	27	0	29.63	57.154	0.00	0.00	0.00	9.846
628	TTGTATAACCCAGTGTTAGTTAGTTTT	298	27	0	29.63	57.150	0.00	0.00	42.50	9.850
629	TGTATAACCCAGTGTTAGTTAGTTTTT	297	27	0	29.63	57.150	0.00	0.00	42.50	9.850
630	CCAGTGTTAGTTAGTTTTCTAATGTG	289	27	0	33.33	57.121	0.00	0.00	40.23	9.879
631	ACACAGAGTCTGAATAATGTCTTAATT	255	27	0	29.63	57.050	17.85	0.00	35.18	9.950
632	TTTTTCTGCTGGATTCAACGGTTTCTG	350	27	0	40.74	62.957	0.00	0.00	0.00	9.957
633	AGTTTTCTGCTGGATTCAACGGTTTC	352	27	0	40.74	62.959	0.00	0.00	0.00	9.959
634	GTTTTTCTGCTGGATTCAACGGTTTCT	351	27	0	40.74	62.959	0.00	0.00	0.00	9.959
635	ACACTACAAATAAATCTTTGAATGCAA	165	27	0	25.93	57.020	2.43	0.00	38.17	9.980

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### BLASTN 2.6.1+

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: PD7SMY6J014

Database: Nucleotide collection (nt)

43,336,626 sequences; 150,108,365,041 total letters

Query= Length=20

### Score E

Sequences p	(Bits)	Value	
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KY457839.1	Human papillomavirus type 18 strain 18CNTZ35 E6 (E	37.4	2.6
KY457838.1	Human papillomavirus type 18 strain 18CNTZ34 E6 (E	37.4	2.6
KY457837.1	Human papillomavirus type 18 strain 18CNTZ33 E6 (E	37.4	2.6
KY457836.1	Human papillomavirus type 18 strain 18CNTZ32 E6 (E	37.4	2.6
KY457835.1	Human papillomavirus type 18 strain 18CNTZ31 E6 (E	37.4	2.6
KY457834.1	Human papillomavirus type 18 strain 18CNTZ30 E6 (E	37.4	2.6
KY457833.1	Human papillomavirus type 18 strain 18CNTZ29 E6 (E	37.4	2.6
KY457832.1	Human papillomavirus type 18 strain 18CNTZ28 E6 (E	37.4	2.6
KY457831.1	Human papillomavirus type 18 strain 18CNTZ27 E6 (E	37.4	2.6
KY457830.1	Human papillomavirus type 18 strain 18CNTZ26 E6 (E	37.4	2.6
KY457829.1	Human papillomavirus type 18 strain 18CNTZ25 E6 (E	37.4	2.6
КҮ457828.1	Human papillomavirus type 18 strain 18CNTZ24 E6 (E	37.4	2.6
KY457827.1	Human papillomavirus type 18 strain 18CNTZ23 E6 (E	37.4	2.6
KY457826.1	Human papillomavirus type 18 strain 18CNTZ22 E6 (E	37.4	2.6
KY457825.1	Human papillomavirus type 18 strain 18CNTZ21 E6 (E	37.4	2.6
KY457824.1	Human papillomavirus type 18 strain 18CNTZ20 E6 (E	37.4	2.6
KY457823.1	Human papillomavirus type 18 strain 18CNTZ19 E6 (E	37.4	2.6
KY457822.1	Human papillomavirus type 18 strain 18CNTZ18 E6 (E	37.4	2.6
KY457821.1	Human papillomavirus type 18 strain 18CNTZ17 E6 (E	37.4	2.6
KY457820.1	Human papillomavirus type 18 strain 18CNTZ16 E6 (E	37.4	2.6
KY457819.1	Human papillomavirus type 18 strain 18CNTZ15 E6 (E	37.4	2.6
KY457818.1	Human papillomavirus type 18 strain 18CNTZ14 E6 (E	37.4	2.6
KY457817.1	Human papillomavirus type 18 strain 18CNTZ13 E6 (E	37.4	2.6
KY457816.1	Human papillomavirus type 18 strain 18CNTZ12 E6 (E	37.4	2.6
KY457815.1	Human papillomavirus type 18 strain 18CNTZ11 E6 (E	37.4	2.6
KY457814.1	Human papillomavirus type 18 strain 18CNTZ10 E6 (E	37.4	2.6

# BLAST INFORMATION KY457813.1 Human

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KY457807.1 Human papillomavirus type 18 strain 18CNTZ03 E6 (E... 37.4
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KY502086.1 Human papillomavirus isolate HPV18 P1-10 transform... 37.4
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KY502085.1 Human papillomavirus isolate HPV18 Pl transforming... 37.4
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and L1 (L1) genes, complete cds
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Score = 37.4 bits (40), Expect = 2.6
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and L1 (L1) genes, complete cds
Length=2502
Score = 37.4 \text{ bits } (40), Expect = 2.6
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
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Query 1 GCGCTTTGAGGATCCAACAC 20

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BLAST INFORMATION
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Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
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Sbjct 6
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and L1 (L1) genes, complete cds
Length=2502
Score = 37.4 bits (40), Expect = 2.6
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
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Query 1
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Identities = 20/20 (100%), Gaps = 0/20 (0%)
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and L1 (L1) genes, complete cds

Length=2502

## BLAST INFORMATION Score = 37.4 bits (40), Expect = 2.6Identities = 20/20 (100%), Gaps = 0/20 (0%) Strand=Plus/Plus GCGCTTTGAGGATCCAACAC 20 Query 1 Sbjct 6 GCGCTTTGAGGATCCAACAC 25 >KY457834.1 Human papillomavirus type 18 strain 18CNTZ30 E6 (E6), E7 (E7), and L1 (L1) genes, complete cds Length=2502 Score = 37.4 bits (40), Expect = 2.6Identities = 20/20 (100%), Gaps = 0/20 (0%) Strand=Plus/Plus Query 1 GCGCTTTGAGGATCCAACAC 20 Sbjct 6 GCGCTTTGAGGATCCAACAC 25 >KY457833.1 Human papillomavirus type 18 strain 18CNTZ29 E6 (E6), E7 (E7), and L1 (L1) genes, complete cds Length=2502 Score = 37.4 bits (40), Expect = 2.6 Identities = 20/20 (100%), Gaps = 0/20 (0%) Strand=Plus/Plus GCGCTTTGAGGATCCAACAC 20 Query 1 GCGCTTTGAGGATCCAACAC 25 Sbjct 6 >KY457832.1 Human papillomavirus type 18 strain 18CNTZ28 E6 (E6), E7 (E7), and L1 (L1) genes, complete cds Length=2502 Score = 37.4 bits (40), Expect = 2.6

Identities = 20/20 (100%), Gaps = 0/20 (0%)

GCGCTTTGAGGATCCAACAC 20

Sbjct 6 GCGCTTTGAGGATCCAACAC 25

Strand=Plus/Plus

Query 1

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and L1 (L1) genes, complete cds
Length=2502
Score = 37.4 bits (40), Expect = 2.6
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
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Query 1
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Sbjct 6
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Strand=Plus/Plus
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Query 1
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Score = 37.4 \text{ bits } (40), Expect = 2.6
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
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Query 1
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Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
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Strand=Plus/Plus
Query 1
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Length=2502
Score = 37.4 bits (40), Expect = 2.6
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Strand=Plus/Plus
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Query 1
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Sbict 6
>KY457825.1 Human papillomavirus type 18 strain 18CNTZ21 E6 (E6), E7 (E7),
and L1 (L1) genes, complete cds
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Strand=Plus/Plus
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Length=2502
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Identities = 20/20 (100%), Gaps = 0/20 (0%)
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Query 1
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Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
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Score = 37.4 bits (40), Expect = 2.6
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Strand=Plus/Plus
Query 1
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Score = 37.4 bits (40), Expect = 2.6
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
Query 1
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Sbjct 6 GCGCTTTGAGGATCCAACAC 25

Score = 37.4 bits (40), Expect = 2.6

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Score = 37.4 bits (40), Expect = 2.6
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
Query 1
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Length=2502
Score = 37.4 bits (40), Expect = 2.6
Identities = 20/20 (100%), Gaps = 0/20 (0%)
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>KY457817.1 Human papillomavirus type 18 strain 18CNTZ13 E6 (E6), E7 (E7),
and L1 (L1) genes, complete cds
Length=2502
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BLAST INFORMATION
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
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Query 1
          GCGCTTTGAGGATCCAACAC 25
Sbjct 6
>KY457816.1 Human papillomavirus type 18 strain 18CNTZ12 E6 (E6), E7 (E7),
and L1 (L1) genes, complete cds
Length=2502
Score = 37.4 bits (40), Expect = 2.6
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
Query 1
         GCGCTTTGAGGATCCAACAC 20
          Sbjct 6 GCGCTTTGAGGATCCAACAC 25
>KY457815.1 Human papillomavirus type 18 strain 18CNTZ11 E6 (E6), E7 (E7),
and L1 (L1) genes, complete cds
Length=2502
Score = 37.4 bits (40), Expect = 2.6
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
Query 1
        GCGCTTTGAGGATCCAACAC 20
          Sbjct 6 GCGCTTTGAGGATCCAACAC 25
>KY457814.1 Human papillomavirus type 18 strain 18CNTZ10 E6 (E6), E7 (E7),
and L1 (L1) genes, complete cds
Length=2502
Score = 37.4 bits (40), Expect = 2.6
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
         GCGCTTTGAGGATCCAACAC 20
Query 1
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Sbjct 6 GCGCTTTGAGGATCCAACAC 25

```
and L1 (L1) genes, complete cds
Length=2502
Score = 37.4 bits (40), Expect = 2.6
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
Query 1
        GCGCTTTGAGGATCCAACAC 20
          Sbjct 6 GCGCTTTGAGGATCCAACAC 25
>KY457812.1 Human papillomavirus type 18 strain 18CNTZ08 E6 (E6), E7 (E7),
and L1 (L1) genes, complete cds
Length=2502
Score = 37.4 bits (40), Expect = 2.6
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
Query 1 GCGCTTTGAGGATCCAACAC 20
          Sbjct 6
        GCGCTTTGAGGATCCAACAC 25
>KY457811.1 Human papillomavirus type 18 strain 18CNTZ07 E6 (E6), E7 (E7),
and L1 (L1) genes, complete cds
Length=2502
Score = 37.4 bits (40), Expect = 2.6
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
         GCGCTTTGAGGATCCAACAC 20
Query 1
          Sbjct 6 GCGCTTTGAGGATCCAACAC 25
>KY457810.1 Human papillomavirus type 18 strain 18CNTZ06 E6 (E6), E7 (E7),
and L1 (L1) genes, complete cds
Length=2502
Score = 37.4 bits (40), Expect = 2.6
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
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>KY457813.1 Human papillomavirus type 18 strain 18CNTZ09 E6 (E6), E7 (E7),

## BLAST INFORMATION GCGCTTTGAGGATCCAACAC 20 Query 1 GCGCTTTGAGGATCCAACAC 25 Sbjct 6 >KY457809.1 Human papillomavirus type 18 strain 18CNTZ05 E6 (E6), E7 (E7), and L1 (L1) genes, complete cds Length=2502 Score = 37.4 bits (40), Expect = 2.6 Identities = 20/20 (100%), Gaps = 0/20 (0%) Strand=Plus/Plus Query 1 GCGCTTTGAGGATCCAACAC 20 Sbjct 6 GCGCTTTGAGGATCCAACAC 25 >KY457808.1 Human papillomavirus type 18 strain 18CNTZ04 E6 (E6), E7 (E7), and L1 (L1) genes, complete cds Length=2502 Score = 37.4 bits (40), Expect = 2.6Identities = 20/20 (100%), Gaps = 0/20 (0%) Strand=Plus/Plus Query 1 GCGCTTTGAGGATCCAACAC 20 Sbjct 6 GCGCTTTGAGGATCCAACAC 25 >KY457807.1 Human papillomavirus type 18 strain 18CNTZ03 E6 (E6), E7 (E7), and L1 (L1) genes, complete cds Length=2502 Score = 37.4 bits (40), Expect = 2.6Identities = 20/20 (100%), Gaps = 0/20 (0%) Strand=Plus/Plus

```
>KY457806.1 Human papillomavirus type 18 strain 18CNTZ02 E6 (E6), E7 (E7), and L1 (L1) genes, complete cds Length=2502
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GCGCTTTGAGGATCCAACAC 20

Sbjct 6 GCGCTTTGAGGATCCAACAC 25

Query 1

Score = 37.4 bits (40), Expect = 2.6

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Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
Query 1
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         Sbjct 6 GCGCTTTGAGGATCCAACAC 25
>KY457805.1 Human papillomavirus type 18 strain 18CNTZ01 E6 (E6), E7 (E7),
and L1 (L1) genes, complete cds
Length=2502
Score = 37.4 bits (40), Expect = 2.6
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
Query 1
         GCGCTTTGAGGATCCAACAC 20
         Sbjct 6
        GCGCTTTGAGGATCCAACAC 25
>KX514433.1 Human papillomavirus type 18 isolate pam9, complete genome
Length=7857
Score = 37.4 bits (40), Expect = 2.6
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
         GCGCTTTGAGGATCCAACAC 20
Query 1
         Sbjct 110 GCGCTTTGAGGATCCAACAC 129
>KY502096.1 Human papillomavirus isolate HPV18 P2-50 transforming protein
E6 (E6) gene, partial cds
Length=474
Score = 37.4 bits (40), Expect = 2.6
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
        GCGCTTTGAGGATCCAACAC 20
Query 1
         Sbjct 6 GCGCTTTGAGGATCCAACAC 25
```

```
>KY502095.1 Human papillomavirus isolate HPV18 P2-40 transforming protein
E6 (E6) gene, partial cds
Length=474
Score = 37.4 bits (40), Expect = 2.6
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
        GCGCTTTGAGGATCCAACAC 20
Query 1
          Sbjct 6
        GCGCTTTGAGGATCCAACAC 25
>KY502094.1 Human papillomavirus isolate HPV18 P2-30 transforming protein
E6 (E6) gene, partial cds
Length=474
Score = 37.4 bits (40), Expect = 2.6
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
         GCGCTTTGAGGATCCAACAC 20
Query 1
          Sbjct 6 GCGCTTTGAGGATCCAACAC 25
>KY502093.1 Human papillomavirus isolate HPV18 P2-20 transforming protein
E6 (E6) gene, partial cds
Length=474
Score = 37.4 bits (40), Expect = 2.6
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
        GCGCTTTGAGGATCCAACAC 20
Query 1
          Sbjct 6 GCGCTTTGAGGATCCAACAC 25
>KY502092.1 Human papillomavirus isolate HPV18 P2-10 transforming protein
E6 (E6) gene, partial cds
Length=474
Score = 37.4 bits (40), Expect = 2.6
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
```

```
Query 1
         GCGCTTTGAGGATCCAACAC 20
         Sbjct 6 GCGCTTTGAGGATCCAACAC 25
>KY502091.1 Human papillomavirus isolate HPV18 P2 transforming protein E6
(E6) gene, partial cds
Length=474
Score = 37.4 bits (40), Expect = 2.6
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
Query 1
        GCGCTTTGAGGATCCAACAC 20
         Sbjct 6 GCGCTTTGAGGATCCAACAC 25
>KY502090.1 Human papillomavirus isolate HPV18 P1-50 transforming protein
E6 (E6) gene, partial cds
Length=474
Score = 37.4 bits (40), Expect = 2.6
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
        GCGCTTTGAGGATCCAACAC 20
Query 1
         GCGCTTTGAGGATCCAACAC 25
Sbict 6
>KY502089.1 Human papillomavirus isolate HPV18 P1-40 transforming protein
E6 (E6) gene, partial cds
Length=474
Score = 37.4 bits (40), Expect = 2.6
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
Query 1
         GCGCTTTGAGGATCCAACAC 20
         Sbjct 6 GCGCTTTGAGGATCCAACAC 25
```

>KY502088.1 Human papillomavirus isolate HPV18 P1-30 transforming protein E6 (E6) gene, partial cds

Length=474

```
Score = 37.4 bits (40), Expect = 2.6
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
         GCGCTTTGAGGATCCAACAC 20
Query 1
          GCGCTTTGAGGATCCAACAC 25
>KY502086.1 Human papillomavirus isolate HPV18 P1-10 transforming protein
E6 (E6) gene, partial cds
Length=474
Score = 37.4 bits (40), Expect = 2.6
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
         GCGCTTTGAGGATCCAACAC 20
Query 1
          Sbjct 6 GCGCTTTGAGGATCCAACAC 25
>KY502085.1 Human papillomavirus isolate HPV18 P1 transforming protein E6
(E6) gene, partial cds
Length=474
Score = 37.4 bits (40), Expect = 2.6
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
Query 1
         GCGCTTTGAGGATCCAACAC 20
          Sbjct 6 GCGCTTTGAGGATCCAACAC 25
>KX545359.1 Human papillomavirus type 18 isolate NGSk274-18 E6 protein (E6)
gene, complete cds; and E7 protein (E7) gene, partial cds
Length=516
Score = 37.4 bits (40), Expect = 2.6
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
          GCGCTTTGAGGATCCAACAC 20
Query 1
```

```
>KX545354.1 Human papillomavirus type 18 isolate NGSk256-18 E6 protein (E6)
gene, complete cds; and E7 protein (E7) gene, partial cds
Length=579
Score = 37.4 bits (40), Expect = 2.6
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
Query 1
          GCGCTTTGAGGATCCAACAC 20
          Sbjct 6 GCGCTTTGAGGATCCAACAC 25
Database: Nucleotide collection (nt)
Posted date: Jul 10, 2017 6:27 PM
Number of letters in database: 150,108,365,041
Number of sequences in database: 43,336,626
Lambda
           K
                    Н
0.634
       0.408 0.912
Gapped
Lambda
           K
                    Н
0.625
       0.410
                 0.780
Matrix: blastn matrix:2 -3
Gap Penalties: Existence: 5, Extension: 2
Number of Sequences: 43336626
Number of Hits to DB: 823797
Number of extensions: 350
Number of successful extensions: 350
Number of sequences better than 10: 33
Number of HSP's better than 10 without gapping: 0
Number of HSP's gapped: 350
Number of HSP's successfully gapped: 33
Length of query: 20
Length of database: 150108365041
Length adjustment: 17
Effective length of query: 3
Effective length of database: 149371642399
Effective search space: 448114927197
Effective search space used: 448114927197
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)

### BLASTN 2.6.1+

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: PD7T5GZZ014

Database: Nucleotide collection (nt)

43,336,626 sequences; 150,108,365,041 total letters

Query= Length=20

### Score E

Sequences p	roducing significant alignments:	(Bits)	Value
KY457840.1	Human papillomavirus type 18 strain 18CNTZ36 E6 (E	37.4	2.6
KY457839.1	Human papillomavirus type 18 strain 18CNTZ35 E6 (E	37.4	2.6
KY457838.1	Human papillomavirus type 18 strain 18CNTZ34 E6 (E	37.4	2.6
KY457837.1	Human papillomavirus type 18 strain 18CNTZ33 E6 (E	37.4	2.6
KY457836.1	Human papillomavirus type 18 strain 18CNTZ32 E6 (E	37.4	2.6
KY457835.1	Human papillomavirus type 18 strain 18CNTZ31 E6 (E	37.4	2.6
KY457834.1	Human papillomavirus type 18 strain 18CNTZ30 E6 (E	37.4	2.6
KY457833.1	Human papillomavirus type 18 strain 18CNTZ29 E6 (E	37.4	2.6
KY457832.1	Human papillomavirus type 18 strain 18CNTZ28 E6 (E	37.4	2.6
KY457831.1	Human papillomavirus type 18 strain 18CNTZ27 E6 (E	37.4	2.6
KY457830.1	Human papillomavirus type 18 strain 18CNTZ26 E6 (E	37.4	2.6
KY457829.1	Human papillomavirus type 18 strain 18CNTZ25 E6 (E	37.4	2.6
KY457828.1	Human papillomavirus type 18 strain 18CNTZ24 E6 (E	37.4	2.6
KY457827.1	Human papillomavirus type 18 strain 18CNTZ23 E6 (E	37.4	2.6
KY457826.1	Human papillomavirus type 18 strain 18CNTZ22 E6 (E	37.4	2.6
KY457825.1	Human papillomavirus type 18 strain 18CNTZ21 E6 (E	37.4	2.6
KY457824.1	Human papillomavirus type 18 strain 18CNTZ20 E6 (E	37.4	2.6
KY457823.1	Human papillomavirus type 18 strain 18CNTZ19 E6 (E	37.4	2.6
KY457822.1	Human papillomavirus type 18 strain 18CNTZ18 E6 (E	37.4	2.6
KY457821.1	Human papillomavirus type 18 strain 18CNTZ17 E6 (E	37.4	2.6
KY457820.1	Human papillomavirus type 18 strain 18CNTZ16 E6 (E	37.4	2.6

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KY457819.1
           Human papillomavirus type 18 strain 18CNTZ15 E6 (E... 37.4
                                                                          2.6
KY457818.1 Human papillomavirus type 18 strain 18CNTZ14 E6 (E... 37.4
                                                                          2.6
KY457817.1 Human papillomavirus type 18 strain 18CNTZ13 E6 (E... 37.4
                                                                          2.6
KY457816.1 Human papillomavirus type 18 strain 18CNTZ12 E6 (E... 37.4
                                                                          2.6
KY457815.1 Human papillomavirus type 18 strain 18CNTZ11 E6 (E... 37.4
                                                                          2.6
KY457814.1 Human papillomavirus type 18 strain 18CNTZ10 E6 (E... 37.4
                                                                          2.6
KY457813.1 Human papillomavirus type 18 strain 18CNTZ09 E6 (E... 37.4
                                                                          2.6
KY457812.1 Human papillomavirus type 18 strain 18CNTZ08 E6 (E... 37.4
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KY457811.1 Human papillomavirus type 18 strain 18CNTZ07 E6 (E... 37.4
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KY457810.1 Human papillomavirus type 18 strain 18CNTZ06 E6 (E... 37.4
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KY457809.1 Human papillomavirus type 18 strain 18CNTZ05 E6 (E... 37.4
                                                                          2.6
KY457808.1 Human papillomavirus type 18 strain 18CNTZ04 E6 (E... 37.4
                                                                          2.6
KY457807.1 Human papillomavirus type 18 strain 18CNTZ03 E6 (E... 37.4
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KY457806.1 Human papillomavirus type 18 strain 18CNTZ02 E6 (E... 37.4
                                                                          2.6
KY457805.1 Human papillomavirus type 18 strain 18CNTZ01 E6 (E... 37.4
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KX514433.1 Human papillomavirus type 18 isolate pam9, complet... 37.4
                                                                          2.6
KY502096.1 Human papillomavirus isolate HPV18 P2-50 transform... 37.4
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KY502095.1 Human papillomavirus isolate HPV18 P2-40 transform... 37.4
                                                                          2.6
KY502094.1 Human papillomavirus isolate HPV18 P2-30 transform... 37.4
                                                                          2.6
KY502093.1 Human papillomavirus isolate HPV18 P2-20 transform... 37.4
                                                                          2.6
KY502092.1 Human papillomavirus isolate HPV18 P2-10 transform... 37.4
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KY502091.1 Human papillomavirus isolate HPV18 P2 transforming... 37.4
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KY502090.1 Human papillomavirus isolate HPV18 P1-50 transform... 37.4
                                                                          2.6
KY502089.1 Human papillomavirus isolate HPV18 P1-40 transform... 37.4
                                                                          2.6
KY502088.1 Human papillomavirus isolate HPV18 P1-30 transform... 37.4
                                                                          2.6
KY502086.1 Human papillomavirus isolate HPV18 P1-10 transform... 37.4
                                                                          2.6
KY502085.1 Human papillomavirus isolate HPV18 Pl transforming... 37.4
                                                                          2.6
KX545359.1 Human papillomavirus type 18 isolate NGSk274-18 E6... 37.4
                                                                          2.6
KX545354.1 Human papillomavirus type 18 isolate NGSk256-18 E6... 37.4
                                                                          2.6
ALIGNMENTS
>KY457840.1 Human papillomavirus type 18 strain 18CNTZ36 E6 (E6), E7 (E7),
Length=2502
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and L1 (L1) genes, complete cds

Score = 37.4 bits (40), Expect = 2.6Identities = 20/20 (100%), Gaps = 0/20 (0%) Strand=Plus/Plus

CGCTTTGAGGATCCAACACG 20 Query 1 Sbjct 7 CGCTTTGAGGATCCAACACG 26

>KY457839.1 Human papillomavirus type 18 strain 18CNTZ35 E6 (E6), E7 (E7), and L1 (L1) genes, complete cds Length=2502

Score = 37.4 bits (40), Expect = 2.6

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Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
Query 1
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         Sbjct 7 CGCTTTGAGGATCCAACACG 26
>KY457838.1 Human papillomavirus type 18 strain 18CNTZ34 E6 (E6), E7 (E7),
and L1 (L1) genes, complete cds
Length=2502
Score = 37.4 bits (40), Expect = 2.6
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
         CGCTTTGAGGATCCAACACG 20
Query 1
         Sbjct 7
        CGCTTTGAGGATCCAACACG 26
>KY457837.1 Human papillomavirus type 18 strain 18CNTZ33 E6 (E6), E7 (E7),
and L1 (L1) genes, complete cds
Length=2502
Score = 37.4 bits (40), Expect = 2.6
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
        CGCTTTGAGGATCCAACACG 20
Query 1
         Sbjct 7 CGCTTTGAGGATCCAACACG 26
>KY457836.1 Human papillomavirus type 18 strain 18CNTZ32 E6 (E6), E7 (E7),
and L1 (L1) genes, complete cds
Length=2502
Score = 37.4 bits (40), Expect = 2.6
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
Query 1
        CGCTTTGAGGATCCAACACG 20
         Sbjct 7 CGCTTTGAGGATCCAACACG 26
```

```
>KY457835.1 Human papillomavirus type 18 strain 18CNTZ31 E6 (E6), E7 (E7),
and L1 (L1) genes, complete cds
Length=2502
Score = 37.4 bits (40), Expect = 2.6
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
          CGCTTTGAGGATCCAACACG 20
Query 1
          Sbjct 7 CGCTTTGAGGATCCAACACG 26
>KY457834.1 Human papillomavirus type 18 strain 18CNTZ30 E6 (E6), E7 (E7),
and L1 (L1) genes, complete cds
Length=2502
Score = 37.4 bits (40), Expect = 2.6
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
Query 1
          CGCTTTGAGGATCCAACACG 20
          Sbjct 7 CGCTTTGAGGATCCAACACG 26
>KY457833.1 Human papillomavirus type 18 strain 18CNTZ29 E6 (E6), E7 (E7),
and L1 (L1) genes, complete cds
Length=2502
Score = 37.4 bits (40), Expect = 2.6
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
Query 1
          CGCTTTGAGGATCCAACACG 20
          CGCTTTGAGGATCCAACACG 26
Sbjct 7
>KY457832.1 Human papillomavirus type 18 strain 18CNTZ28 E6 (E6), E7 (E7),
and L1 (L1) genes, complete cds
Length=2502
Score = 37.4 bits (40), Expect = 2.6
Identities = 20/20 (100%), Gaps = 0/20 (0%)
```

Strand=Plus/Plus

```
Query 1
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         Sbjct 7 CGCTTTGAGGATCCAACACG 26
>KY457831.1 Human papillomavirus type 18 strain 18CNTZ27 E6 (E6), E7 (E7),
and L1 (L1) genes, complete cds
Length=2502
Score = 37.4 bits (40), Expect = 2.6
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
        CGCTTTGAGGATCCAACACG 20
Query 1
         Sbjct 7
        CGCTTTGAGGATCCAACACG 26
>KY457830.1 Human papillomavirus type 18 strain 18CNTZ26 E6 (E6), E7 (E7),
and L1 (L1) genes, complete cds
Length=2502
Score = 37.4 bits (40), Expect = 2.6
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
        CGCTTTGAGGATCCAACACG 20
Query 1
         Sbjct 7 CGCTTTGAGGATCCAACACG 26
>KY457829.1 Human papillomavirus type 18 strain 18CNTZ25 E6 (E6), E7 (E7),
and L1 (L1) genes, complete cds
Length=2502
Score = 37.4 bits (40), Expect = 2.6
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
Query 1
         CGCTTTGAGGATCCAACACG 20
         Sbjct 7 CGCTTTGAGGATCCAACACG 26
>KY457828.1 Human papillomavirus type 18 strain 18CNTZ24 E6 (E6), E7 (E7),
```

# BLAST INFORMATION and L1 (L1) genes, complete cds Length=2502 Score = 37.4 bits (40), Expect = 2.6Identities = 20/20 (100%), Gaps = 0/20 (0%) Strand=Plus/Plus Query 1 CGCTTTGAGGATCCAACACG 20 Sbjct 7 CGCTTTGAGGATCCAACACG 26 >KY457827.1 Human papillomavirus type 18 strain 18CNTZ23 E6 (E6), E7 (E7), and L1 (L1) genes, complete cds Length=2502 Score = 37.4 bits (40), Expect = 2.6Identities = 20/20 (100%), Gaps = 0/20 (0%) Strand=Plus/Plus Query 1 CGCTTTGAGGATCCAACACG 20 Sbjct 7 CGCTTTGAGGATCCAACACG 26 >KY457826.1 Human papillomavirus type 18 strain 18CNTZ22 E6 (E6), E7 (E7), and L1 (L1) genes, complete cds Length=2502 Score = 37.4 bits (40), Expect = 2.6 Identities = 20/20 (100%), Gaps = 0/20 (0%) Strand=Plus/Plus Query 1 CGCTTTGAGGATCCAACACG 20 CGCTTTGAGGATCCAACACG 26 Sbjct 7 >KY457825.1 Human papillomavirus type 18 strain 18CNTZ21 E6 (E6), E7 (E7), and L1 (L1) genes, complete cds Length=2502 Score = 37.4 bits (40), Expect = 2.6 Identities = 20/20 (100%), Gaps = 0/20 (0%) Strand=Plus/Plus

Query 1 CGCTTTGAGGATCCAACACG 20

```
BLAST INFORMATION
         Sbjct 7 CGCTTTGAGGATCCAACACG 26
>KY457824.1 Human papillomavirus type 18 strain 18CNTZ20 E6 (E6), E7 (E7),
and L1 (L1) genes, complete cds
Length=2502
Score = 37.4 bits (40), Expect = 2.6
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
Query 1 CGCTTTGAGGATCCAACACG 20
         CGCTTTGAGGATCCAACACG 26
Sbjct 7
>KY457823.1 Human papillomavirus type 18 strain 18CNTZ19 E6 (E6), E7 (E7),
and L1 (L1) genes, complete cds
Length=2502
Score = 37.4 bits (40), Expect = 2.6
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
        CGCTTTGAGGATCCAACACG 20
Query 1
         Sbjct 7 CGCTTTGAGGATCCAACACG 26
>KY457822.1 Human papillomavirus type 18 strain 18CNTZ18 E6 (E6), E7 (E7),
and L1 (L1) genes, complete cds
Length=2502
Score = 37.4 bits (40), Expect = 2.6
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
Query 1
        CGCTTTGAGGATCCAACACG 20
         Sbjct 7 CGCTTTGAGGATCCAACACG 26
```

>KY457821.1 Human papillomavirus type 18 strain 18CNTZ17 E6 (E6), E7 (E7),

and L1 (L1) genes, complete cds

Length=2502

# BLAST INFORMATION Score = 37.4 bits (40), Expect = 2.6Identities = 20/20 (100%), Gaps = 0/20 (0%) Strand=Plus/Plus Query 1 CGCTTTGAGGATCCAACACG 20 Sbjct 7 CGCTTTGAGGATCCAACACG 26 >KY457820.1 Human papillomavirus type 18 strain 18CNTZ16 E6 (E6), E7 (E7), and L1 (L1) genes, complete cds Length=2502 Score = 37.4 bits (40), Expect = 2.6Identities = 20/20 (100%), Gaps = 0/20 (0%) Strand=Plus/Plus Query 1 CGCTTTGAGGATCCAACACG 20 Sbjct 7 CGCTTTGAGGATCCAACACG 26 >KY457819.1 Human papillomavirus type 18 strain 18CNTZ15 E6 (E6), E7 (E7), and L1 (L1) genes, complete cds Length=2502 Score = 37.4 bits (40), Expect = 2.6 Identities = 20/20 (100%), Gaps = 0/20 (0%) Strand=Plus/Plus CGCTTTGAGGATCCAACACG 20 Query 1 Sbjct 7 CGCTTTGAGGATCCAACACG 26 >KY457818.1 Human papillomavirus type 18 strain 18CNTZ14 E6 (E6), E7 (E7), and L1 (L1) genes, complete cds Length=2502 Score = 37.4 bits (40), Expect = 2.6

Identities = 20/20 (100%), Gaps = 0/20 (0%)

CGCTTTGAGGATCCAACACG 20

Sbjct 7 CGCTTTGAGGATCCAACACG 26

Strand=Plus/Plus

Query 1

```
and L1 (L1) genes, complete cds
Length=2502
Score = 37.4 bits (40), Expect = 2.6
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
          CGCTTTGAGGATCCAACACG 20
Query 1
          CGCTTTGAGGATCCAACACG 26
Sbjct 7
>KY457816.1 Human papillomavirus type 18 strain 18CNTZ12 E6 (E6), E7 (E7),
and L1 (L1) genes, complete cds
Length=2502
Score = 37.4 bits (40), Expect = 2.6
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
          CGCTTTGAGGATCCAACACG 20
Query 1
          Sbjct 7 CGCTTTGAGGATCCAACACG 26
>KY457815.1 Human papillomavirus type 18 strain 18CNTZ11 E6 (E6), E7 (E7),
and L1 (L1) genes, complete cds
Length=2502
Score = 37.4 \text{ bits } (40), Expect = 2.6
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
          CGCTTTGAGGATCCAACACG 20
Query 1
          Sbjct 7 CGCTTTGAGGATCCAACACG 26
>KY457814.1 Human papillomavirus type 18 strain 18CNTZ10 E6 (E6), E7 (E7),
and L1 (L1) genes, complete cds
Length=2502
Score = 37.4 bits (40), Expect = 2.6
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
```

>KY457817.1 Human papillomavirus type 18 strain 18CNTZ13 E6 (E6), E7 (E7),

```
Query 1
         CGCTTTGAGGATCCAACACG 20
         Sbjct 7 CGCTTTGAGGATCCAACACG 26
>KY457813.1 Human papillomavirus type 18 strain 18CNTZ09 E6 (E6), E7 (E7),
and L1 (L1) genes, complete cds
Length=2502
Score = 37.4 bits (40), Expect = 2.6
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
Query 1
         CGCTTTGAGGATCCAACACG 20
         Sbjct 7 CGCTTTGAGGATCCAACACG 26
>KY457812.1 Human papillomavirus type 18 strain 18CNTZ08 E6 (E6), E7 (E7),
and L1 (L1) genes, complete cds
Length=2502
Score = 37.4 bits (40), Expect = 2.6
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
        CGCTTTGAGGATCCAACACG 20
Query 1
         Sbjct 7 CGCTTTGAGGATCCAACACG 26
>KY457811.1 Human papillomavirus type 18 strain 18CNTZ07 E6 (E6), E7 (E7),
and L1 (L1) genes, complete cds
Length=2502
Score = 37.4 bits (40), Expect = 2.6
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
Query 1
         CGCTTTGAGGATCCAACACG 20
         Sbjct 7 CGCTTTGAGGATCCAACACG 26
>KY457810.1 Human papillomavirus type 18 strain 18CNTZ06 E6 (E6), E7 (E7),
and L1 (L1) genes, complete cds
```

```
Length=2502
```

```
Score = 37.4 bits (40), Expect = 2.6
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
         CGCTTTGAGGATCCAACACG 20
Query 1
          CGCTTTGAGGATCCAACACG 26
>KY457809.1 Human papillomavirus type 18 strain 18CNTZ05 E6 (E6), E7 (E7),
and L1 (L1) genes, complete cds
Length=2502
Score = 37.4 bits (40), Expect = 2.6
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
Query 1
         CGCTTTGAGGATCCAACACG 20
          Sbjct 7 CGCTTTGAGGATCCAACACG 26
>KY457808.1 Human papillomavirus type 18 strain 18CNTZ04 E6 (E6), E7 (E7),
and L1 (L1) genes, complete cds
Length=2502
Score = 37.4 bits (40), Expect = 2.6
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
Query 1
         CGCTTTGAGGATCCAACACG 20
          Sbjct 7 CGCTTTGAGGATCCAACACG 26
>KY457807.1 Human papillomavirus type 18 strain 18CNTZ03 E6 (E6), E7 (E7),
and L1 (L1) genes, complete cds
Length=2502
Score = 37.4 bits (40), Expect = 2.6
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
Query 1
         CGCTTTGAGGATCCAACACG 20
```

```
Sbjct 7 CGCTTTGAGGATCCAACACG 26
>KY457806.1 Human papillomavirus type 18 strain 18CNTZ02 E6 (E6), E7 (E7),
and L1 (L1) genes, complete cds
Length=2502
Score = 37.4 bits (40), Expect = 2.6
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
Query 1
        CGCTTTGAGGATCCAACACG 20
          Sbjct 7 CGCTTTGAGGATCCAACACG 26
>KY457805.1 Human papillomavirus type 18 strain 18CNTZ01 E6 (E6), E7 (E7),
and L1 (L1) genes, complete cds
Length=2502
Score = 37.4 bits (40), Expect = 2.6
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
Query 1
         CGCTTTGAGGATCCAACACG 20
          Sbjct 7
        CGCTTTGAGGATCCAACACG 26
>KX514433.1 Human papillomavirus type 18 isolate pam9, complete genome
Length=7857
Score = 37.4 bits (40), Expect = 2.6
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
Query 1
          CGCTTTGAGGATCCAACACG 20
          Sbjct 111 CGCTTTGAGGATCCAACACG 130
>KY502096.1 Human papillomavirus isolate HPV18 P2-50 transforming protein
E6 (E6) gene, partial cds
Length=474
Score = 37.4 bits (40), Expect = 2.6
```

Identities = 20/20 (100%), Gaps = 0/20 (0%)

Strand=Plus/Plus

Query 1 CGCTTTGAGGATCCAACACG 20 Sbjct 7 CGCTTTGAGGATCCAACACG 26 >KY502095.1 Human papillomavirus isolate HPV18 P2-40 transforming protein E6 (E6) gene, partial cds Length=474 Score = 37.4 bits (40), Expect = 2.6Identities = 20/20 (100%), Gaps = 0/20 (0%) Strand=Plus/Plus Query 1 CGCTTTGAGGATCCAACACG 20 Sbjct 7 CGCTTTGAGGATCCAACACG 26 >KY502094.1 Human papillomavirus isolate HPV18 P2-30 transforming protein E6 (E6) gene, partial cds Length=474 Score = 37.4 bits (40), Expect = 2.6 Identities = 20/20 (100%), Gaps = 0/20 (0%) Strand=Plus/Plus CGCTTTGAGGATCCAACACG 20 Query 1 Sbjct 7 CGCTTTGAGGATCCAACACG 26 >KY502093.1 Human papillomavirus isolate HPV18 P2-20 transforming protein E6 (E6) gene, partial cds Length=474 Score = 37.4 bits (40), Expect = 2.6Identities = 20/20 (100%), Gaps = 0/20 (0%) Strand=Plus/Plus CGCTTTGAGGATCCAACACG 20 Query 1 Sbjct 7 CGCTTTGAGGATCCAACACG 26

>KY502092.1 Human papillomavirus isolate HPV18 P2-10 transforming protein

# BLAST INFORMATION E6 (E6) gene, partial cds Length=474 Score = 37.4 bits (40), Expect = 2.6 Identities = 20/20 (100%), Gaps = 0/20 (0%) Strand=Plus/Plus CGCTTTGAGGATCCAACACG 20 Query 1 Sbjct 7 CGCTTTGAGGATCCAACACG 26 >KY502091.1 Human papillomavirus isolate HPV18 P2 transforming protein E6 (E6) gene, partial cds Length=474 Score = 37.4 bits (40), Expect = 2.6Identities = 20/20 (100%), Gaps = 0/20 (0%) Strand=Plus/Plus Query 1 CGCTTTGAGGATCCAACACG 20 Sbjct 7 CGCTTTGAGGATCCAACACG 26 >KY502090.1 Human papillomavirus isolate HPV18 P1-50 transforming protein E6 (E6) gene, partial cds Length=474 Score = 37.4 bits (40), Expect = 2.6 Identities = 20/20 (100%), Gaps = 0/20 (0%) Strand=Plus/Plus CGCTTTGAGGATCCAACACG 20 Query 1 Sbjct 7 CGCTTTGAGGATCCAACACG 26

>KY502089.1 Human papillomavirus isolate HPV18 P1-40 transforming protein E6 (E6) gene, partial cds Length=474

Score = 37.4 bits (40), Expect = 2.6
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus

Query 1 CGCTTTGAGGATCCAACACG 20

```
BLAST INFORMATION
         Sbjct 7 CGCTTTGAGGATCCAACACG 26
>KY502088.1 Human papillomavirus isolate HPV18 P1-30 transforming protein
E6 (E6) gene, partial cds
Length=474
Score = 37.4 bits (40), Expect = 2.6
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
        CGCTTTGAGGATCCAACACG 20
Query 1
          Sbjct 7
        CGCTTTGAGGATCCAACACG 26
>KY502086.1 Human papillomavirus isolate HPV18 P1-10 transforming protein
E6 (E6) gene, partial cds
Length=474
Score = 37.4 bits (40), Expect = 2.6
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
        CGCTTTGAGGATCCAACACG 20
Query 1
          Sbjct 7 CGCTTTGAGGATCCAACACG 26
>KY502085.1 Human papillomavirus isolate HPV18 P1 transforming protein E6
(E6) gene, partial cds
Length=474
Score = 37.4 bits (40), Expect = 2.6
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
        CGCTTTGAGGATCCAACACG 20
Query 1
```

>KX545359.1 Human papillomavirus type 18 isolate NGSk274-18 E6 protein (E6) gene, complete cds; and E7 protein (E7) gene, partial cds Length=516

Sbjct 7 CGCTTTGAGGATCCAACACG 26

```
Score = 37.4 bits (40), Expect = 2.6
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
          CGCTTTGAGGATCCAACACG 20
Query 1
          Sbjct 7 CGCTTTGAGGATCCAACACG 26
>KX545354.1 Human papillomavirus type 18 isolate NGSk256-18 E6 protein (E6)
gene, complete cds; and E7 protein (E7) gene, partial cds
Length=579
Score = 37.4 bits (40), Expect = 2.6
Identities = 20/20 (100%), Gaps = 0/20 (0%)
Strand=Plus/Plus
Query 1
          CGCTTTGAGGATCCAACACG 20
          Sbjct 7 CGCTTTGAGGATCCAACACG 26
Database: Nucleotide collection (nt)
Posted date: Jul 10, 2017 6:27 PM
Number of letters in database: 150,108,365,041
Number of sequences in database: 43,336,626
Lambda
          K
                    Η
0.634
       0.408 0.912
Gapped
Lambda
           K
                    Н
       0.410
                 0.780
0.625
Matrix: blastn matrix:2 -3
Gap Penalties: Existence: 5, Extension: 2
Number of Sequences: 43336626
Number of Hits to DB: 789966
Number of extensions: 241
Number of successful extensions: 241
Number of sequences better than 10: 30
Number of HSP's better than 10 without gapping: 0
Number of HSP's gapped: 241
Number of HSP's successfully gapped: 30
Length of query: 20
Length of database: 150108365041
Length adjustment: 17
Effective length of query: 3
Effective length of database: 149371642399
```

Effective search space: 448114927197

Effective search space used: 448114927197

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)