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(54) PROTEINS AND IMMUNIZING  
COMPOSITIONS CONTAINING  
PASTEURELLA PROTEINS AND METHODS  
OF USE

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## (57) ABSTRACT

The present invention provides isolated proteins isolatable from a *Pasteurella* spp. such as *P. multocida*. Also provided by the present invention are compositions that include one or more of the proteins, and methods for making and methods for using the proteins.

Specification includes a Sequence Listing.

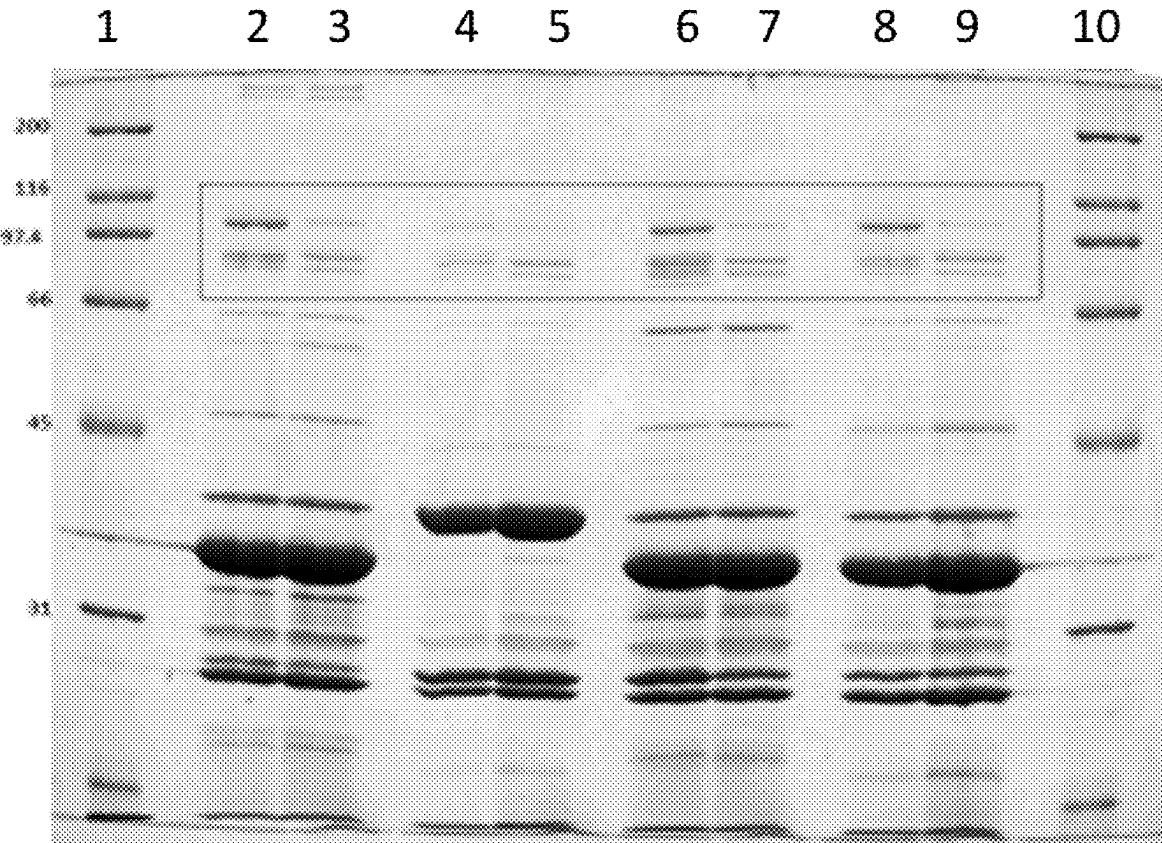


FIG. 1

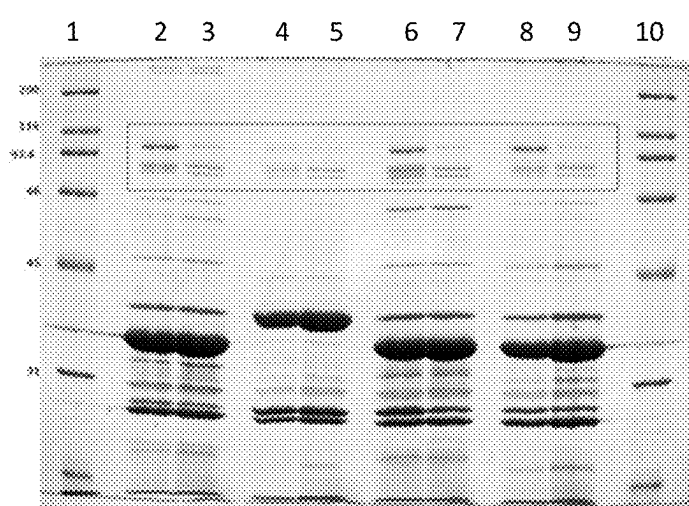


FIG. 2

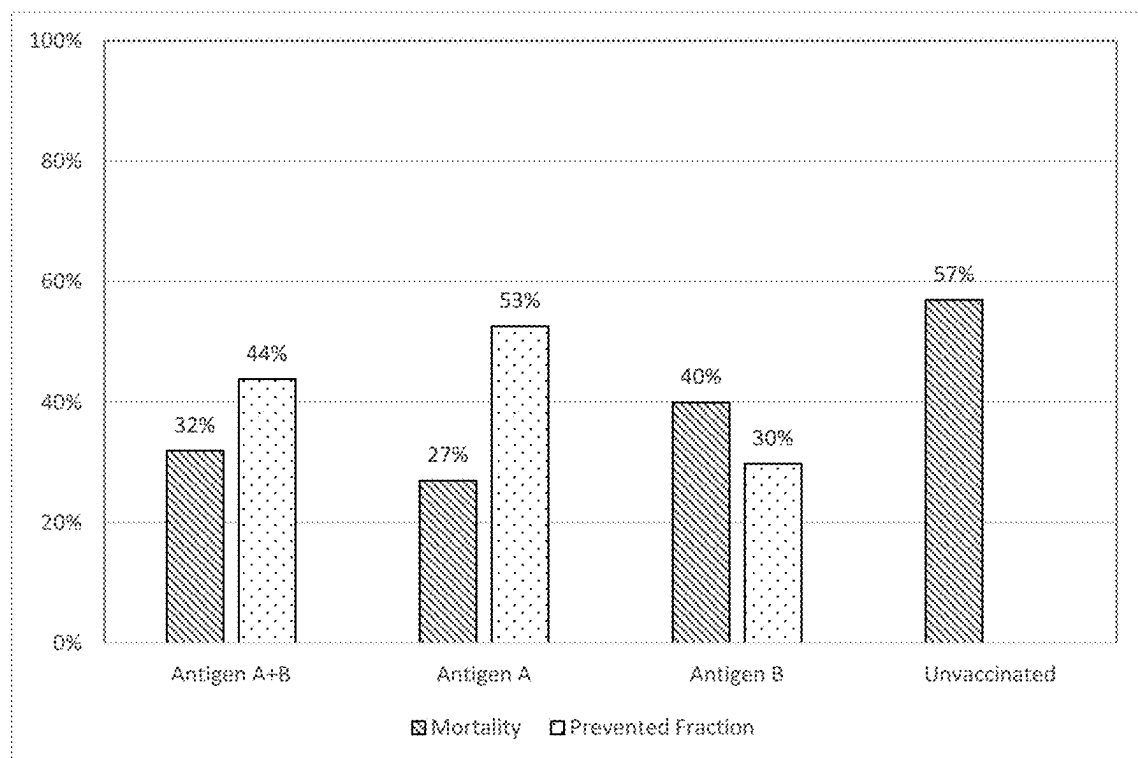


FIG. 3

Protein ID	MW	Description	Vaccine Strain MS061130 type A: 3x4	X73: type A:1	p1059 type A:3
1	128,216	Putative Ton-B dependent heme receptor	Present	Present	Present
2	114,216	Outer membrane receptor protein, mostly Fe transport	Present	Present	Absent
3	109,900	Outer membrane receptor protein, mostly Fe transport	Absent	Present	Absent
4	109,863	Outer membrane receptor protein, mostly Fe transport	Present	Absent	Present
5	104,901	Outer membrane receptor protein, mostly Fe transport	Absent	Present	Present
6	91,308	Outer membrane receptor protein, mostly Fe transport	Absent	Present	Absent
7	90,917	Outer membrane receptor protein, mostly Fe transport	Absent	Present	Present
8	90,887	Outer membrane receptor protein, mostly Fe transport	Absent	Present	Absent
9	90,601	Outer membrane receptor protein, mostly Fe transport	Present	Absent	Present
10	89,635	TonB-dependent hemoglobin receptor	Present	Present	Present
11	88,107	Outer membrane receptor protein, mostly Fe transport	Absent	Present	Absent
12	84,792	Outer membrane receptor protein, mostly Fe transport, HemR	Absent	Absent	Present
13	81,394	PfhR	Present	Present	Present
14	59,543	Heme binding protein A HbpA	Absent	Present	Present

FIG. 4

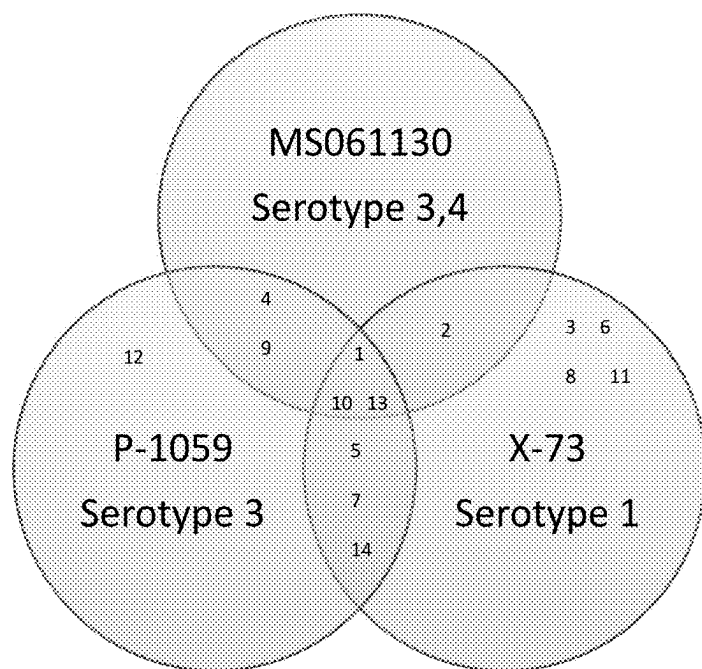


FIG. 5

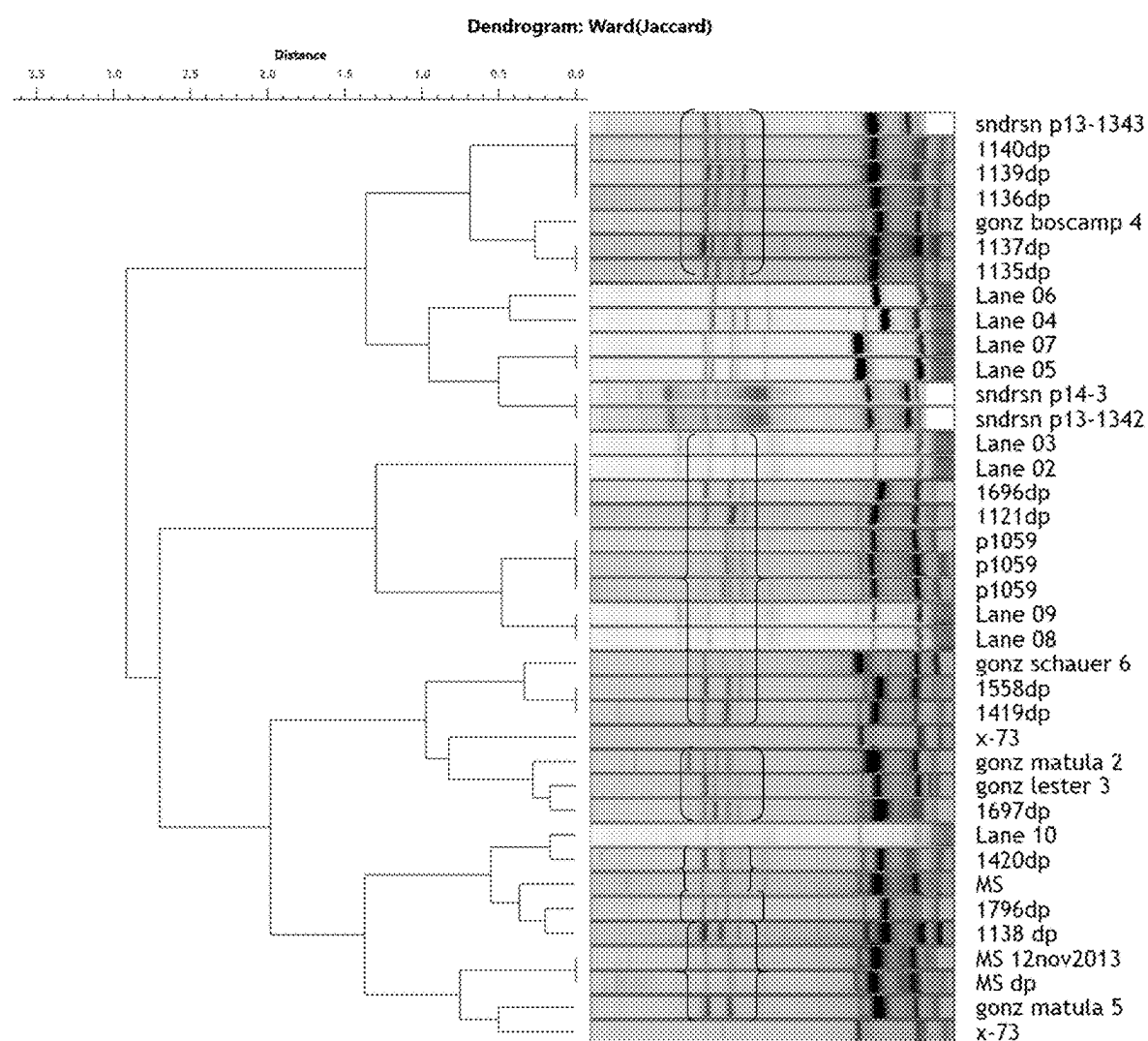


FIG. 6

MW	% identity	MS	x-73	p1059	p1662	1121	1135	1138	1558	1696	gonz2	gonz4	gonz5	gonz6
109,862		(+)	(-)	(+)	(+)	(+)	(-)	(+)	(-)	(-)	(-)	(+)	(+)	(-)
109,900	97%	(-)	(+)	(-)	(-)	(-)	(+)	(-)	(+)	(+)	(-)	(-)	(-)	(+)
92,670		(-)	(-)	(-)	(-)	(-)	(+)	(-)	(-)	(-)	(+)	(+)	(-)	(-)
91,308	>93%	(+)	(+)	(-)	(+)	(+)	(-)	(+)	(+)	(+)	(-)	(-)	(+)	(+)
89,631	99%	(+)	(-)	(-)	(-)	(-)	(+)	(-)	(-)	(-)	(+)	(+)	(-)	(+)
84,910	99%	(-)	(-)	(+)	(-)	(-)	(+)	(-)	(+)	(+)	(-)	(-)	(-)	(-)
81,332	99%	(-)	(+)	(-)	(-)	(+)	(-)	(+)	(-)	(-)	(-)	(+)	(-)	(+)

FIG. 7

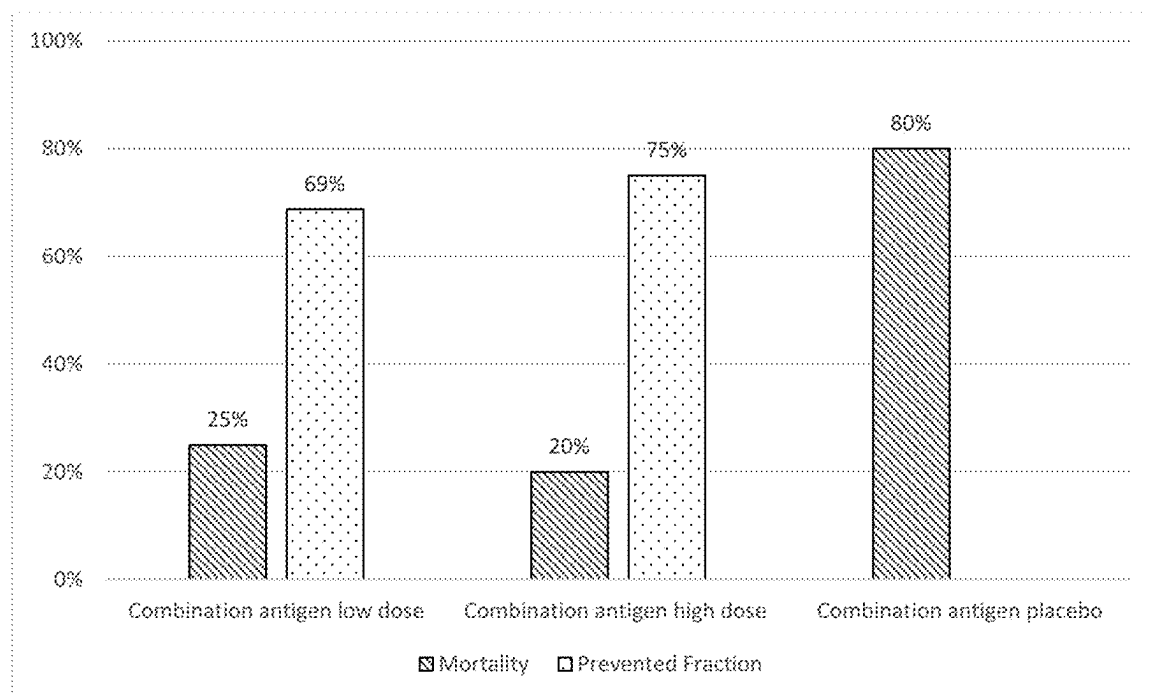


FIG. 8-1.

SEQ ID NO:1

ATGCGTACAACAACAATAAAATTCTGCAATTACATTGGCATTATTGAGTTATTGTGGGCTAT  
TTTGGCGGATAGTCATCAAGAGGCAGACTGAACCTGATACGATTACCCTTCTCAACAAGATG  
AGATGAATATTAAGAGAAAAAGTCGGTGAAACTGTGAAAACGGCAGTCATTGAAACGCCAG  
CAAGTACAGGATAGTCGTATCTGCGCTATGAAACCGGTGTGACTGTGGTAGAAGCTGGACG  
TTTGGGTCAGCGGTTATGCCATTCTGGTGGATGAGAACCGAGTGGCAATTACAGTAGATG  
GCTTACATCAAGCAGAAACCTTCTCAAGGTTAAAGAATTATTTGAAGGTTACGGCAAT  
TTAACAAATACCCGAAATAGTGTGAAATTGAGACGTTGAAAGTCGCTAAATCGCAGAAAGGTGC  
TGATTCTGAAAAGTGGGTAGTGGTTCTTGGGAGGCCTGTACTTTGAAACAAAAGATGCCA  
GAGATTCCCTGACTGAAAAAGATTGGCATATCGGCTATAAAGCAGGCTACTCAACGGCAGATAAT  
CAGGGATTAAATGCAGTGAATCTGCAGGTCGCTATCAAATGTTGATGCATTGATTATGCATT  
TAAGCGACATGGACATGAATTAGAAAATTATGACTATAAAATGGCAGAGATATTCAAGGGAAAG  
AAAGAGAGAAAGCGGATCCTTATACGATTACGAAAGAAAGTACATTAGTGAATTCTCTTTCG  
CCAACAGAAAATCATCGTTTACAGTCGCTCTGATACTTATCTCAGCATTCCGGACATGA  
TCTTCATACAATCTGTTGCAACACATATTCAAGTTAGATGAGAAAGAATCTCGTCATGCAA  
ATGATCTGACAAAACGTAAAATGTTCTTACTTATGAAAATTATACTGTTACGCCATTG  
GATACGCTCAAGTAAAGCTATTCAACAAAGAATTACAACAAGAGCAAGAACAGAAGATTACTG  
TGATGGTAATGAAAATGTGACTCTTATAAGAATCCTTAGGGCTCAATTAAAAGAGGGAAAAA  
TCGTTGATCGTAATGGCGATCCTGTTAATTGAAGCTGTTGATGGTAAACATCAAGTTG  
AAAGCTGGTAAGCCTTGTGATGTAGCCTCTGGAACTAATTATGCGGCTTCAGGTAAAGAATT  
AAGTCCTTCTTGGTTAGATTGCTCTATTGATTGCTAAGCCTATCAATACTTATA  
AATATCGCTATACTCTCGGAGCCAATTGCAAGCAAATTACTTAAATAAAACCATG  
AAATGGAAAGACATTGCTACTTATGATGGCGTGGACACTATATTACCAAAATTCTAAAGG  
TTACTTGCTTGGATTATAAGAGCGTGAATTAAACAAAGACGAAACAAATTAAATTAGATT  
TAACAAAAGCATTACTCTTTGAGATTGAAAATGAACCTTCCTATGGTGGTGTACGCGAAA  
ACGACCAAGGAAATGGTGAATAAAGCAGGATATTGGCGTAATCCTACTGGTGGCGAGAG  
AACGTTAGGGCAATCATGGGAAAATTGAGAGAGTGAAGACAAGTCTTCATATAATGGGATGC  
TATGTCCTCGTCATGAACCATTAAACCTCTTCTTAATTCCGGTAGAAGCAACAAC  
TAAGTCTTATACTGAGACAATATCAAGTTGCAAAATATGTTAAGTGTAGATTAG  
TATTAAATATCAGCCAGAATATATTCTGGTGTGACACCTAAATTGCAAGTGA  
GGTTATTATTCCATTACCTGAAGGTGAAAAGTAACGTAGGGACAATGGTATT  
CTCACTCAGCGCAAATTGTAAGAATGCGGAGGAAAATTGCTTATATTG  
CACAAGAAAAACGCTTAAAGAAACATTCTTATTCTGGTGCAACGTT  
GATTTACGATTTACCGAACCCCGTGTGAAACCAGAGGAAGCAAA  
ATCAAGAGATTGCATTAAACAGTCACGATAATTGGGATT  
TTGATTAGCGTATTAGGTTCAAGAAATTATCGAATTCCGTGG  
GAGGGCAGGCACAAGCAAG  
GATTTCAGTTATCAAATGTCAATGTCGATAATG  
CAGGTTAAAGGACTTGAAATT  
ACGTTGAATTGGGATT  
ACCAACGTGGCGTTGGATGGCGATCGT  
CCAATGAATGCGATT  
CAGCCTAAAGCTCTGTTT  
GGTTAGGCTATGAT  
CATAAAGAAATAAATT  
GGCGCTGATT  
TATATCACAC  
GTGTGAGTGA

FIG. 8-2.

AAAAAAAGCGAAAGACACTTATAATATGTTCTATAAAGAACAGGGATATAAGATAGTGCTGTTCG  
TTGGAGAAGTGATGACTATACGCTAGTTGATGCGGTTGGTTATATTAAACCGATTAAGAATTAA  
CGTTACAGTTGGCCTTATAATTGACAGACCGTAAACTTGACATGGGAATCTGCTCGTTCG  
ATTAAACCATTGGTACAAGTAATTAAATTACAGAAAACAGGCAGGAATTACGTTTTA  
CTCACCAAGGTCGTAAATTAAATTAGTGCCGAAATTACCTTCTAA

SEQ ID NO:2

MRTTIKFSAITLALLSYCGAILADSHQEATELDITVSSQQDEMNIKEKKVGETVKTASQLKRQ  
QVQDSRDLVRYETGVTVVEAGRFGSSGYAIRGVDENRAITVDGLHQAETLSSQGFKELFEGYGN  
FNNTRNSVEIETLKVAKIAKGADSVKGSGSLGGAVLFETKDARDFLTEKDWHIGYKAGYSTADN  
QGLNAVTLAGRYQMFDALIMHSKRHGHELENYDYKNGRDIQGKEREKADPYTITKESTLVKFSFS  
PTENHRFTVASDTYLQHSRGHDLSYNLVATTHIQLDEKESRHANDLTKRKNVSFTYENYTVPFW  
DTLKLSYSQRITTRARTEDYCDGNEKCDSYKNPLGLQLKEGKIVDRNGDPVNKLVDGKHQVVD  
KAGKPFVASGTYAAFSGKELSPSSFWLDCSIFDCSKPINTKYRYSSEPTLQQITLNKTMEI  
NGKTFTAYDGRGHYIILPNSKGYLPDYKERDLNTKTKQINLDLTKAFTLFEIENELSYGGVYAK  
TTKEMVNKAGYYGRNPTWWAERTLGQSWGKLRECKTSSSYNGMLCPRHEPLTSFLIPVEATTKSL  
YFADNIKLHNMLSVDLGYRYDDIKYQPEYIPGVTPKIADDMVKGFIPLPEGEKVTGTMVFTKP  
LTQAQIRKNAEENIAYIAQEKFKKHSYSLGATFDPLNFLRVQVKYSKGFRAPTSDELYFTFKHP  
DFTILPNPVLKPEEAKNQEIALTVHDNWGFVTSVFQTKYRFIDLAYLGSRNLSNVGGQAQAR  
DFQVYQNVNVDNAVKGLEINARLNLYFWHVLDFNTSYKFTYQRGRLDGDRPMNAIQPKASVF  
GLGYDHKENKGADLYITRVSEKKAKDTYNMFYKEQGYKDSAVRWRSDDYTLDAVGYIKPIKNL  
TLQFGVYNLTDRKYLTwESARSIKPFGTSNLINQKTGAGINRFYSPGRNFKFSAEITF\*

FIG. 9-1.

SEQ ID NO:3

ATGGAATCCGAAAAATCCACTCAAAAGACGACACTGCACTCCTCTGTTCTACCGCCTTCTCGCTTCCGCAAAACCGATAACGAACGCCGATAAAAATCACTTCTGACGGAAATTGTCGTGTATGCGGATCAAATAATCAATGAGTTCAACACAGAGTGTCACTCAAGATGACATGAAAAAGCCTGTACAAATGGTAATATTACTGACTATTACGTTCAAATCCGATGTGCGTTATGAGAATAGCGATCAAATGGATTGCAACGAGGTGAAATTAAACCGAAAACATTCAATAATGGTGCAGATTATCAGCAGACCACTTTTGTCGATAATGTAATATCAATAATGATATGGGATTGGTAGTGATCTTTGACGGCACCATGGAACAGTCCCCTTGCCAATCATTCTCAAGGCTACTTTTGATGCAATCTGCTCTTCATTGAGTGATGACAGTAATGTTCTGCCAGCTAGGAGGATTGCGGGGGAGCGTCGGCGAAAACCAAACAATATGATGGGAAAGATCGGTTAAAATTAGTTATCGTACAAACTGATGCCTCATGGGCAAATTAAAGTAGAACAGAACATCTGAACGTTAAAATGCGATTCCCTGAAGGATCGGTGCCGAATTCAACCTAAGTATTCTAAACACTTTCAATATCACGGCAGAAAAAGGGTTAAGTAAAATTAGGTATGGTATCGGATTAAGTCGTAGAACATCAAGTATCACAACATCACGTCAGTCAAATTAAATCCACAAGGTGATCGTGTAAACAAACACATACAAGACGTTCAGACAATGCGTTACTGAATTAACTGACACCAACGACAAACACCGTTGAACTCGTTTCAGACTCAAATTATCGTGGCGTAAATTCTTAATACTAATATAGACAGTAATGTTTGACTATCACCGTGTATGGTGTGACTTCTCTGGATAAACGCCCTCAGTCTGGCATTCTCACGACGACACTCGCCTATGATAACTTGATGACACGAGAAAATCAGCGTCAACTTATATGAAAACGACGCTCACTGAAGAGGGTGAGGAATACCGAAGGTGGAATGGTAACAGTCATTGAAACCAAAAAATTACATACGTCACCTGAATATGCGATGAATCTTTAATTAGGTTCAATAGAACACTCTGTATCATTAGCGGTATTATCAAGCAACGAAATATCGCTTACGCGACATTAGATGCAGTCAGTCGGTGAATTGTATACTCCTGATTGGTTAAATGGTAATACAGATAAAACTTATATTGACGCAACGTAACATAGCCAAAAAGGGACCGTAAAACACGTTATCAAATATCGCACTTACGTGGAAGATTAAATGACGTGGAAAAATCTCGAATTCGGCGGGACTTCGCTTCTGAAACGTGATGATTATTAAAAACACGAATTGGCGCCAAGAACCGTTTCGTTATAAACCATTTGAAGACACCGCATTAGTCAGTGTGGTTGGAACCGTTAC TACGGACGTTCTTGCCTCAATGAAATTATCTGAAGGTATCTCAAATTAGATGGTATGATGACCTTCCGTTACAAAGACCTTAGCCAATTAAACCCCTTATTAGACGAACTGAGCTTGGTGTAGAACATATGTTGCTAATCTCGCCTTCACCTGAAATACATTACGTGATAACAAACACGTATTGTTTACAAGAAGAAGAGGTGATGTTAAATGGAGAAAGGAAAAACTCCGCTATTACCAACGAGGAAAAGACTACAAACGAATGTGTTAACCTTCAAATTAAACACACAAGCGCCTTGGGAGCTTGGTCACACACGCTGGACAAGTGCAGTGGCATTGATTGGTAGATAGCAAAGCTATCGATCATGGCAGGGCTATAACGGTTCGACACCTGTGATTTAGACGGGAAATTAAATGACTTATGAGCAAATGTTGAAAAAAGTCATGCTTACAAAGAACATGGGTCTACGTTAAATCTTGACATGTTGTTCAAATTATGATCTTCTTGGGCTAATACAATCTATGTCAAACCCACCAACACCTAACCGAACGCGTTAGCAGTAATACACCTGAGGTATATCGTAGCTATGATTGGTACTTACACAATGGGATACCAACAGTCTCGCTGGCAGCCAACCTTCGCAAGAAAACACCGCTTATATTAAATTAGATGTTACTTAATGTTGTTAAATACACGTAAGGCGCGGGTCAAATGGACAAGATCTGGCATCTACACCCCCGGTCGTTAGTTCTGGCTTGAAGTTGTTACGAATTCTAA

FIG. 9-2.

SEQ ID NO:4

MESAKNPLKTTLALLCCSTAFSLSAKTDTNADKNHFLTEIVVYADQNKSMSSTQSVTQDDMKKS  
PVTNGNITDYLRSNPHVRYENSNDQNGLQRGEIKPENISINGADYQQTTFFVDNVNIINNDMFGFSD  
LFDGTMATVPFANHSQGYFFDANLSSIVHDSNVASLGGFAGGAVVAKTKQYDGKDRLKFSYR  
TTDASWAKFKVEDKDLERFKNAIPEGSVAEFQPKYSKFFFNITAEGGLSENLGMVIGLSRRTSSI  
QQSRQINPQGDRDKQTHRRSDNALLNFNLTPNDKHRFELGFRYSNYRERKFFNTNIDSNVFDYH  
RAYGVTFSWINALQSGILTTLAYDNFDDTRKSASTYMKTTLTEEEGEYTEGGMGNSQLNQKNLH  
TSLEYAMNPFLGSIEHSVSLGGIYQATKYRFTRHSDAVGELYTPDWLNGNTDKLILTQRNIACK  
GTVKTRYQNIALYVEDLMTWKNLEFRAGLRLERDDYLKNTNLAPRTVFYKPFEDTAFSVGWNRY  
YGRSFASMKLSEGIFKLDGHDTFRYKDLSQFKTPYSDELSFGVEQYVANLAFHLKYILRDNKQRI  
VLQEEEVMLNGERKKLRYYQRGKDYKTNVLTFQINTQAPWELEGPTRWTSAVAFDWLDSKAIDHGR  
GYNGSTPVILDGKLMTYEQMLKKVNAYKETWGLRLNLDMFVPIFDLSWANTIIYVKPPTLTERVS  
SNTPEVYRSYDYGTYTQWDTSLRWQPTFAEKHRPYIKLDVLNVLNKTRKGAGPNGQDLGIYTPGR  
EFWLEVGYEF\*

FIG. 10-1.

SEQ ID NO:5

ATGTCATTCAAACATAAAACACTGGCGTTTGTGACATGCTGCTGCACCTCTGCCTAGC  
AGAAAACGTGGCTACCACGTTAGAACCCATCGGGTTCTGATCTCAGTCATACCAACGCTGAACC  
TTGATCAAATAAGCTGAAAAAGAAAGTCCAAAAGATTAAAGCTATTGCCCACAACGCCA  
AATATTAAATGTTATCCATACGGGACATGCACAATTAGCGATATTGAAATTGCGGTATGGGAAG  
CAGCCGAGAAATCTTGCTACCGGGCAAACCGCGTCACAATGGAATTAGACGGTATGGACATTA  
GCCCGAGTTTATTGGACACAGCTCACGCCATGGTCGGCAATATTGATCCCAGTGTATCTA  
AAACGTGTTGAGATTCAAAGGTCCAACAGTCAGGCGTGGCAGGGCATGTCGTTCCAAAC  
GAAAGATCCTCGTGAATTCTCTGCCTAACCAACGTACAGGTGCACAACACTCGTGCTGGCTATT  
TAGGCAGATGTGATGCTTATTATGTTGGATAACTGGTGCACCTTATTGGATGAACACAGTAGT  
GCTTTAGTGAGCTATACACGACGCTGGTTAATGAATTAAATAAAGGGAGGCTTGGATGTCAC  
AGGTAGTCACGTACTAAAAGCAATCCTCCAGGGTTAGTAATGCAAGTGAACAGTAAATTAC  
GCTATTCAACAAATGACCGCCATAAATTACGCTGAATTGCAACATTATGATTAAACGCACC  
GCCTATTAGAAGATAGCTAGGAACAACGACAACACGACGTGGCACAAAACAGTTCATCATAA  
TACCAACATTCAAGAAAATCAGCGTATGCTATTGCTTCAGTCATGACATGCAACAAACCG  
CATTGATCACCTGCACTGGCAAATTGCGTTACAACAAACGAAAAGCACGAGGCCGTAATACA  
GGGGCAGTCACGAATAACATCAGCATCTCCTCCCCAAGTACGCCAAATTAGCCAAGAGCGTTC  
ACTTGATGGCTTAAAACCAAAACCATCAGCTAAAACGGAATTCAATAAAAGCATTGGCAAC  
ATGTCGTACATGAACCTCACTATGGACTAAAATTACAATATGCCAAATGCAAGCTTACGCCAA  
ACACAATCCCTAACGAACAAGGGAGTAACACTCGAACCGCCTTTCCGCACACAGCAACA  
ATGGCAAAGTAAACTCCATCTTCCGATCGGATCAGTTGGTAAATCTGGTTAAGCTTGACAC  
CATCCATACATCTCACACAGATTAGAATCAAACCGAAAACAGAAAATGTATCGAAGAAAACCGT  
GAACAATTATTACTTACAAGGATACGCCATTGGTTACGGTCTGCGTGTGATTATGCACTCAA  
TGAAGCGAATTACTGAGTCTGAACATCAGCACGCCACTCGCTACCCGGCTATGGTAAAACA  
ATGCGCAAAGCTATGGACACTGGCAGCAAAACCGAACCCCATCTACAGCCAGAAACCTCAGAT  
GGTATTGAATTAAGTTGGCGTAGTGCAGGGGGCGATTGGTCAACAAACACGACCTGTTCTATAA  
CCGTTACAATGACTTAATTATCTCGATACCACGGCATGTTATGCTGACCGAACAGGTCAAGTGC  
CTTGTGATTTAGCAAATGAAAAGGACGTAGTTATAGCTATGGAATAGAATTGACGGTAAACTC  
AATCTTGATACGATCGCTCGCTCAAGGAACATATTAAATGCTGGCTCGCTTACAGCAAAGG  
GAAGACCGCGAACAGCAACCACAAGGACGTCTGATCCCTAACAGGCTTGTGGCTTGGCT  
ACCAACAGCCAATGGATTTGGGCATTGAAGGTTAAACTGAAATTGCCCGAAGAAAAAAACT  
AAAGACTTACCCGCCAATCAAGGTTTGAAGGCTTACCGGGCTATGCTGTAGTTGATCTTACCGC  
CTATTATAATGTGACGAAACAGCTTATCTTGGCATCGGCATCTATAATGTCGCTAGATAAAAAT  
ATGCTCGCTGGCAATGGCAAGAGGGACATTAAACATGGTAACTATGACAAGCACACTGAAGCA  
GGTCGTCAATTGGTGCCTATTGCTTACCACTTTAA

SEQ ID NO:6

MSFKHKTIALFVAHACCTSALAENVATTLEPIVVSDSLHTTLNLDQNKLEKESPKDLKAIFATT  
NINVIHTGHAQLGDIEIRGMGSSREIFATGANRVTMELDGMDISPSFYFGHSSRHGRQYFDPSDL  
KRVEIHKGPNSQGVAGHVRFQTKDPDYLLPNQRTGAQLRAGYLGDSDAYVGITGATLLDEHSS  
ALVSYTRRWNFENNKGGLDVTGSQRTKSNPSSGYSNAVNSKLRYSPNDRHKFTLNLQHYDLKRT

FIG. 10-2.

AYLEDSLGGTTTRRGKTVHHNTNIQKNQRHAIASFHDHQQTAAFFDHLHWQIALQQTKSTS RNT  
GAVTN TSASPPPSTPKFSQERSLDGFKTKTISLKTEFNKSIGQHVVHELHYGLKLQYSQM QALRQ  
TQSLNEQGSNRTSAFFPTQQQWQSKLHLSDRISFGKSGLSLTPSIHLTQIRIKPKTENVSKKNR  
EQLFTYKDTAIGYGLRVDYALNEANLLS LNYQHATRLPGYGENNAQS YGHWPAKPNPHLQPETSD  
GIELSWRSAGAIGQQTTLFYNRYNDLIYLDTTACYADRTGQVPCDLANEKGRSY SYGIEFDGKL  
NLDTIGFAQGTYL NAGFAYSKGKTANKQPQGRLDPLTGFVGLGYQQPMDVWGIEGKLKFAAKKKT  
KDLPANQGFEGLPGYAVVDLTAYYNTKQLYLGIGIYNVLDKKYARWAMARGDIKHGNYDKHTEA  
GRHFGANIRYHF\*

FIG. 11-1.

SEQ ID NO:7

ATGCGTACAACAACAATAAAATTCTGCAATTACATTGCATTATTGAGTTATTGTGGGTCA  
TTGGCGGATAGTCATCAAGAGGGCAGTGAACCTGATACGATTACCGTTCTCAACAAGATG  
AGATGAATATTAAAGAGAAAAAGTCGGTGAAACTGTGAAAACGGCAGTCATTGAAACGCCAG  
CAAGTACAGGATAGTCGTATCTTGCGCTATGAAACCGGTGTGACTGTGGTAGAAGCTGGACG  
TTTGGGTCAGCGGTTATGCCATTCTGGTGTGGATGAGAACCGAGTGGCAATTACAGTAGATG  
GCTTACATCAAGCAGAAACCCCTTCTCAAGGTTAAAGAATTATCGAAGGTTACGGCAAT  
TTTAACAATACCGAAATAGTGTGAAATTGAGACGTTGAAAGTCGCTAAATCGCAAAGGTGC  
TGATTCTGAAAAGTGGTAGTGGTCTTGAGGCGCTGTACTTTGAAACAAAAGATGCCA  
GAGATTCCCTGACTGAAAAAGATTGCATATCGGCTATAAAGCGGGCTACTCAACGGCAGATAAT  
CAGGGATTAAATGCAGTGAATCTTGAGGCGCTATCAAATGTTGATGCATTGATTATGCATTC  
TAAGCGACATGGACATGAATTAGAAAATTATGACTATAAAATGGCAGAGATAATTCAAGGGAAAG  
AAAGAGAGAAAGCGGATCCTTACGATTACGAAAGAAAGTACATTAGTGAATTCTCTTTCG  
CCAACAGAAAATCATCGTTTACAGCCTCAATCTAATTATTCACATATGACGAGAAAGAATCGCTCATG  
TCGCTCTTATTCTTACAGCCTCAATCTAATTATTCACATATGACGAGAAAGAATCGCTCATG  
CAAATGATTGACAAAACGTTAAAGTGTGATTCTTACAGCCTCAATCTAATTATTCACATATGACGAGAAAGAATCGCTCATG  
TGGGATACGCTAAATGAGTTATTCAACAAAAGATCAGAACAGAGCAAGAACAGAAGATTA  
TTGTGATGGTAATGAAAATGTGATTCTTACAGCCTCAATCTAATTATTCACATATGACGAGAAAGAATGGCA  
AAATTGTTGATCCAGAGGGAAATCAGATTACTTTAAAGGAACAGGATTTAACAGAAATAGTT  
GATAAAAATGGTAACCCATTCCCTACGACATCTGGTACTAATAATGCAGCATTAGTAATAATAT  
TCAGTTAGGACCTAAAGAATTGGTTAGATTGTTCTCTTGTGATTCTCAGCCATTACTG  
TTTATAACTATCAAATGGCAATATACGCCAAAACAAGTTGAGTTATCTGAAGAAATCACTGTC  
AATGGTAAATTATATAACAGCTAAAGAAGAAAGAGGTGTTAGGAACATTAAATTACCTAA  
TTCAAAAGGTTATTACCATATGATTACAAAGAAAGAGATCTGATTCAAATACAAACAAATCA  
ATTGGATTTAACAAAACATTTCGACTTTAATATAGAAAATGAATTATTACGGTGCATT  
TATTACGTACAGAGAAAGAAAATGGTAATAAAGCAGGTTACGATGGGAGAAATCCTACATGGTGG  
GGCTGATAGAATTAGGGAAAGAGTACGAATTGTAACGAAATGGACTGAAATGTCCTCGTCATG  
AACCTTAACTCTTCTTAATTCCAGTAGAACGACAACCAAGTCTCTATATTTCAGATAAT  
ATCAAATTACACAACATGTTGAGTGTAGATTAGGTTATCGTTATGACGATATTAAATATCAACC  
AGAATATATTCCCTGGTGTAAACACCTAAATTGCAGATGATATGGTGAAGGTTATTTCAT  
TACCTAAGGGAGAAAAGTAACAACACCTGGGGGGCCGAATATAACAAACCACTCACACAGGAA  
CAAATCGTAAGAATGCGGAGGAAAATATTGCTTATATTGACAAAGAAAACGCTTAAAAACAA  
TTCTTATTCTCTGGTGCACGTTCGATCCTCTGAATTTCAGAGTACAAGTAAATATTCAA  
AAGGGTTAGAGCCCCGACTTCGGATGAACATTATTACCTTAAGCATCAGATTTCAGATT  
TTACCGAACCCCGTGTGAAACCAGAGGAAGCAAAAATCAAGAGATTGCATTAACAGTGCACGA  
TAATTGGGATTGTTAGCACAAGTGTGTTCAAACAAAGTATCGTCATTATTGATTAGCGT  
ATTAGGTTCAAGAAATTATCGAATTCCGTGGGAGGGCAGGCACAAGCAAGAGATTCCAAGTT  
TATCAAATGTCAATGTCGATAATGCCAAGGTTAAAGGACTTGAAATTAAATGCACGTTGAATT  
GGGATATTCTGGCATGTGTTGGATGGATTAAACGAGCTATAAATTCACTTACCAATGTGGTC  
GTTGGATGGCGATCGTCAAATGAATGCGATTCTGAGCTAAAGCTTCTGTTGGTTAGGCTAT  
GATCATAAAAGAAAATAATTGGCGCTGATTATATACACACGTGTGAGTGAAGAAAAAGCGAA  
AGACACCTATAATATGTTCTATAAAGAACAGGGTATAAAGATAGTGCATTGTTGGAGAAGTG  
ATGACTATACGCTAGTTGATGCGGTTGGTTATTAAACCGATTAAGAATTAAACGTTACAGTT

FIG. 11-2.

GGCGTTATAATTCACAGACCGTAAACTTGACATGGAATCTGCTCGATTAAACCATT  
TGGTACAAGTAATTAAATTAAATCAGAAAACAGGCAGGAATTACGTTTACTCACCAGGTC  
GTAATTAAATTAGTGCCGAAATTACCTTCTAA

SEQ ID NO:8

MRTTIKFSAITLALLSYCGVILADSHQEATELDITVSSQDEMNIKEKKVGETVKTASQLKRQ  
QVQDSRDLVRYETGTVVEAGRFGSSGYAIRGVDENRAITVDGLHQAETLSSQGFKELFEGYGN  
FNNTRNSVEIETLKVAKIAKGADSVKGSGSLGGAVLFETKDARDFLTEKDWHIGYKAGYSTADN  
QGLNAVTLAGRYQMFDALIMHSKRHGHELENODYKNGRDIQGKEREKADPYTITKESTLVKFSFS  
PTENHRFTVASDTYIQRSRGHDRSYSLQPQSNYFTYDEKECSRHANDLTKRNVSFTYENYSVTPF  
WDTLKLSYSQQKIRTRARTEDYCDGNEKCDSYKNPLGLQLKDGIVDPEGNQITLKGTFNTEIV  
DKNGNPFPPTSGTNNAAFSNNIQLGPKEFWLDCSLFDCTQPFTVNYQNGQYTPKQVELSEEITV  
NGKLYKTAKEERGVRNYLILPNSKGYLPYDYKERDLDNSTKQINLDLTKTGSTFNIENELLYGAI  
YSRTEKKMVNKAGYDGRNPTWWADRILGKSTNCNYNGLKPCRHEPLTSFLIPVEATTKSLYFSDN  
IKLHNMLSVDLGYRYDDIKYQPEYIPGVTPKIADDMVKGIFIPLPKGEKVTPWGAEYTKPLTQE  
QIRKNAEENIAYIAQEKFHKHSYSLGATFDPLNFLRVQVKYSKGFRAPTSDELYFTFKHPDFTI  
LPNPVLKPEEAKNQEIALTVHDNWGFVSTSVEQTKYRHFIIDLAYLGSRNLSNSVGGQAQARDFQV  
YQNVNVDNAKVKGLEINARLNLYFWHVLDFNTSYKFTYQCGRLDGDRPMNAIQPKASVFGLGY  
DHKENKFGADLYITRVSEKKAKDTYNMFYKEQGYKDSAIRWRSDDYTLDAVGYIKPIKNLTLQF  
GVYNLTDRKYLWESARSIKPFGTSNLINQKTGAGINRFYSPGRNFKFSAEITF\*

FIG. 12-1.

SEQ ID NO:9

ATGCAGAAACAGCAACCTATCCCATTACACCTGGGATTTTGATGTTGGGTTACCAACATGGCGTTCAGTCAGCTAATTAGAGAAAATCAACAATCAATAATTGGAAACGATTTGGTCAATGAGAGTGAAGAGAAAATAATTGATGAGAATTGATCAAAACTTATCTGTCTCGGGTTCTTATCTCTTATTATCGCAATCGGATATCAGTACGTTCAGAGGTAGCTCGGTAGGGATTCCCTCTCTGGTGTACCAGGAGTTATTGTGGGAAATAAGCGTAATAGCGCGCTTATCCGTTAATATTGAGGAAATGCGAATGAAAATCGTGTGCCTGTTGGATAGATAAAAGGTCTACAATCGGTACCCCTTACCAAAGGTTATGCAGGTTCTCAACTCGAACCTATTAGATCCGATTGATCAGCCAAGTCGAGATTGAAAAGGTCCCTTTGCAAATGGACGCAACAGGCGCACGGGAGGGTAGTGGAGTAGAGACTTTACGTTGGCAAGATATTATCCTCAAGGGAAAATTGGGGCGTGCCTTGAACACTAGGGACGATGACTAACACCGTATCACCTCCCCTTATTATACAAGAGGAGGATATCAAACAACTAAGTATATTAGTAAATGTCTTCTAATGATACTGGTTATGTCAAACACAAACATATGCACCAAATGCACGCTATTCTTCTCATGGCTTGAATTGATGCATAAACTATAGCCTGGCTTGTCTAATAATGGCAAAATGCTGATCTTGTACTTGCATGCAAAACGTAACAGGCAACTATTGGTGGGCGTATGGACAAACCCCAGTGAATCCATTGAATTGAGGAAGATTCACTAGTAGAAGTCAAAGAGCCTCGCGTTCATGAAGAGGTTGAAATTGGTTCTAACATTAAAGAAAATCGCAGCACCTTATATCGACCGGGTGAAAGGCCCTGAATACCTCACAAGATAAACCTCTTATCTCGCTAAAATAATGTCTACAATGATGTTCATCGTTAGGGTTAGCGTATGCCATTATCATAGCGTTTGGTGAGATTAGAGCTCAATTITGAATTTCAGAGCGTATGGCGCATTGCAAGGTGAAGGGACAGAAGTCAAAGTCGATAGCTATCATGCAAATTATAGCTATAACCCAACGACACCTTATGTGAATTGATGCTATGCAATTTCAGTGAAGCAGTCTAATTTCACCCATTATTGAAGAATATGGTTACTCTTATCCAGTCGTATGCCATTTCAGGTTCTAACAGCAGAAAGGGTAAGTATTGAAAATACTAGCATTCCAGCTAACGACAACCGTTAGTTAAAATATGGCTTGCCTAGTTATGAAACGGATTATCAACCACGTTAGCTCAAGCAGTGTGAGAGCTAACGGGTATCCAGAAGATGCGATTGGTCCACTTATATCGAGATGGTAAGCGTAAAGAATGGAGCGCTTGTGCTGCAACTATCCAATCATTGCTGGTTAAAAGCCGACATCGGGCTACGTTATCTCAATCTACTATTGATTATTGTGAGAACGGAAAGAGTGAATATTGGAGGGCACTTGTGCTTAATCCAATGGATCCGGTAATATTGGGTGGAAAATATAAAGATGTTGTGCATAAACAGGCGCCAGTGAAGGAAATAAGGCATGTCGCAATTGTGCTCACAATTGAACTTAAACGGAGTACAAATTATACGAAATATGCAAGAACGATTGCGTTGCCAAGTTATTCCAAGCAACTAAAGGCTGGTCCATGAGTGCAGCAGGAGATAATCTAGAACAAATTGAGACCTGAAACGCCAAAATTGGGAGGCAGGTATTAACCTGGTTATTTAATAATAGGATAAAAGATTATTGACGCGGAGTTATTCGCCTAAAGATAAGGTGACCGAGACAATTAAACAAAGTCGACAATTAAAGGAATTGAGTTACAGCGTATTGATATGGAAATTGAGTTACAGCGTCTCGCTAATGCGCTAATTGAGTAAACACATCAAGTATTGAGTGGGCGCCTTATTCTTAGTGTAGATTGACGTTAATCGCTATTAAATTAAATAATGCCGTTCTCCGCTTAAATTACATGCGACTTACAGGTTATGTATATCGTAGAACAAAAACTCGATATTGGTGCAGCTATAGTTACTACAGTAAGCGATTAGTACCAAGTCGCTTCTGCAGAACGTTTGTAAACACATCAAGTATTGAGTGGGCGCCTTATTCTTAGTGTAGATTGACGTTAATCGCTATTAAATTAGATATCAATAATATGGGATTAACCGCACCGGGTAGAACATTGCAATTAGGATTAGAAATTACGGTTTAG

Fig. 12-2.

SEQ ID NO:10

MQKQQPYPIHLGIFLMLGLPTWAFSQANLEKSTINKLETILVNESEEKNKFDENLIKTYLSSGSY  
SYLSQSDISTFRGSSVGDFLSGVPGVIVGNKRNSGALSVNIRGIANENRVPVWIDKGLQSVP SYQ  
GYAGSSTRTYLDPDLISQEIEKGPSLQMDATGATGGVVRVETLRWQDIIPQGKNWGVRLKLGTM  
TNTVSPPPYYTRGGYQTKYISKCLSNDTGLCQTQTYAPNARYSSHGFDLNAYNSLAFAWKQNA  
DLVLAYAKRKQGNYFVGRHGQTPVIESIEFEEDSVEVKEPRVHEEEVEIGSLTFKENRSTLYRPGE  
EALNTSQDNTSYLAKINVYNDVHRLGLAYRHYSRGEIMSSIlnFRAYGALQGEGTEVKVDSYH  
ANYSYNPTTPYVNLNVNAYFTDSDSSNFTPFIIEYGYSLSSRAHFLVSKQKGLSIENTSIFQLN  
DKPFSLKYGLAHSYERIYQPRNAQARVRAKGYPEDAIGPLYIRDGKRKEWSAFVAANYPITSWLK  
ADIGLRYLQSTIYDYIVRTERVNIGGALVPNPNGSGNIWVEKYKDVVHKQAPVKNKGSPIVMLT  
FEPINGVQIYT KYAEALRSPSLFQATKGWSMSATADNLEQLRPERAQNWEAGINLFYENLGGKDN  
ILGFKLAYFNNRIKDYLT RSYSPKDKVTQTINIQS AQFKGIELSAYDMGKFYAKLAGTYYT KTK  
FCLTAEQAGKGEQCNSGYVYRSNLNAVPPRLNLHATLGTRLFEQKLDIGARYSYYSKRLVPVLS  
AERFVNNTSSIEWAPYSLVDLYANYNVSNNLKLTMTMDNVFNYYLDINNMGLNTAPGRTLHLGLE  
YRF\*

FIG. 13-1.

SEQ ID NO:11

ATGAAATATCCCTTAAGCTATAAAATAGCAAGGTCCATTCTTCTCATTGCGTT  
TCCTCTGTATGCGCAAGAACGACTGAATTAGAACAAATTACAGTACAAGAAAGCGCAACCGCTG  
AAGTGAACAAAACCTCACCAACAGTGTACAGCAAAAGGCCACGACCATTCAAACGAAATGATT  
CGAGACACCAGAGATTAGTCGCTACACAACGGATGTAGGGATTAGCGATAATGGCGTTTT  
GAAAGGCTTGCGATGCGAGGCCTGAAGATAACCGCCTGGTATCAGTATTGATGGTGTTC  
TGCCTGATTCAAGAAGAAAACTCAGTGTATGCGCTTATGGTAACTTAATAATTCCGCCTAACG  
ATTGATCCTGAATTAAATTCAAACCATGATATTGTACCGGCTCAGATTCTTAATGCAGGCAG  
CGGTTCATAGTGGCGGTGAATTATAACACTTTAGATCCACACATATTGTTAAAACAGGTA  
ATTCCGTTGGTGCTTACTACGAGGCAGTTATGCCAGTAAAATCGTGAATGGTTCGTACTTTA  
GGGATAGGCTATGTTGGCAGAAATTGATGCCTTATTGATGTATTACAACGCACAGGGCACGA  
GTTTAAAAGTCGTGGTTCAGGTCTGAATTCCGGTATTCCAGTAGCCAGCATCCGATCCTGTGA  
CACAAACGCTTCCACAATTATCTCGCTAAATGAATTATCAAATTAAATGACAAACAAACGTATTGGT  
TTAACGCTGAATGGCAAACAGGGGGCGTTACATTGATGAGCCTTATACGTTAATGGCTC  
ACAATGGCGTGAAGCCGATGATCAAACGAACGGTGAATGCAAACCTTATATTATATTATGCAC  
CAAGCACGGGATGGTTAGCCTACAGTAAATTGATTTAGATTACAAAGACCGATCTAGCAGCA  
GTAACTATAAAGGCGGACGCCATTCACGACAGATGCTAAAGAGCTGAACGAAATTATGATCG  
CCGTATGAAAACCGTGTTCAGCGCGGAGTGTAGAACCTCAATGCACAACCTGTACATTCTATG  
GTGAACATACTTAACCATAAGGCTATGTTAGTCAGCGTGTGATTCAAAATATCAATCAAGAT  
CGTATCGGTATTGGCACAAACTACGACACACAATACCACTATACGATTCAATATCCTATTAGAAC  
CAAGCAGTACGGCTTCTTGAAAGATCATGTGCGCTGGATGATACTCTCAAGCCATTAG  
GTTTACGTTATGACCATAACAAACTAAAACCAAGGAACCTCAATGCACCTGTAGCAAAGCCTGT  
TTAGAAGAAGGAAAACCTAAACCGACCCGTTCTACCGTTAGCACATTGCGGGATTGAGGC  
ACAACCTCAGCCCCATGGATGTTAGGCTACAAATTAGCACCGGTTATCGTGTACCAACGGCTT  
CAGAAATGTTCTCAGCTTACCAATGCGTATGGCACGTTGAAATCTAACCCGAGCTGAAACCC  
GAAAAAAAGTATTAAACCATACTCTCTTGAAGGCAATAGTGAAGGCTTGCTGATCTCAC  
CCTCTATCAAACAAACTATCGTCTTATTGAAACAAGAAAGTTAATTCAACGTACCGAAA  
TGCCTATGGACGCCCTTACCAAGCCAAGAACAAACAAATGGTAACCTAGATAAAGCA  
AAAATTATGGTGTGAAATTGAAAACCCATGTCAATTAGATCAGATGATCGCTGTGATACCACA  
AGGCTTAAAGTCTACGCCGCTCGGTTAGCAAAGGTAAACTCTCGAATAACGCCAGCCTAC  
TTTCCATTCAACCGCTTAAATTATTCTGGGTTGGATTATGAAGCAACAAACGGCAAATGGCT  
ATTTTCAACCGCTAACCTATTGGGTGAAAAAGAGCTAGTGTGCGAAAGTGTATGAAATTAA  
ACGTCGCTGTACTGAATTGTCAGAACAGATCCTGGACTGGTCAACAAATTACTCGCTGTA  
AAAAACGGGAATTGTATCCAGATTCTACTTATAAACACTTAAATAATCTGCTTGTGTT  
GATACTTTGGTTATTACAAGATCACGGACGATACAGCTTCCGAGCTGGCATTATAATCTGTT  
TAATAAAAAATACCACACTGGGATGCCTTACGTGGTATTAAATGCCAATAGTACGTTAAATTCA  
TTGACCGTGAAGGAAAGGGTTACAACGCTTCTATGCGCCGGACGTAACATGCGGCTTCCCTT  
GAAATCCGTTCTAA

FIG. 13-2.

SEQ ID NO:12

MKYPLSYKNIARSIPFLSFIAPPLYAQETTELEQITVQESATAEVNKTSPTVISKSATTIQNEMI  
RDTDLVRYTTDVGISDNGRFLKGFMARGVEDNRVGISIDGVSLPSEENSLYARYGNFNNNSRLS  
IDPELIQTIDIVRGSDSFNAGSGSLGGGVNYNTLDPQHIVKTGNSVGALLRGSYASKNREWVRTI  
GIGYVGEKFDALLMYSQRTGHEFKSRGSGPEFRYSSSQHPDPVTQRFHNYLAKMNYQINDKQRIG  
LTLngQtGGRYIDERSYTLMSQWRADDQNERLNANLYIYAPSTGWLAYSKFDLDYQKTDLAA  
VNYKGRHFTTDAKELENEIYDRRMKTVFTRGSVELNAQPVFYGETLTIKGYVSQRDFKNINQD  
RIGIGTNYDTQYHYTIQYPIRTQYGLSLKDHRWNTFSSHGLRYDHTKLKPTELNAPCSKAC  
LEEGKPKPTRFSTVSTFAGFEAQLSPSWMLGYNISTGYRVPTASEMFFSFTNAYGTWKSNPSLKP  
EKSINHTLSLKGNSEKGLLDLTLYQTNYRHFLFEQESLIQRTEMRYGRPYTYQSQQMVNLDA  
KIYGVELKTHVNLDQMIAVIPQGFKFYAAALGYSKGKLSNNASLLSIQPLKIIILGLDYEATNGKWA  
IFNRLTYLGEKRASDAKVYEIKRRCTEFVTETDPWTGQQITRCKKRELYPDLSITYKHLNKS AFVF  
DTFGYYKITDDITFRAGIYNLFNKKYHTWDALRGINANSTLNSVDREGKGLQRFYAPGRNYAASL  
EIRF\*

FIG. 14-1.

SEQ ID NO:13

ATGGATAAAAATTAAATGAAGGGATGTATTCTTATCAATAGTCGGTTGCGGTATCAAATAGG  
GCTAGCATCAAATCCAAATCCTCCAGATGTGGATGAGTTATTACCTATTATGTGAATGCTGATG  
AAGATAATAAATTACCAAGGTCGTTCTGTATTAAAACAGAAAAATATCGATCAACAACAAGCAGAT  
AATGCTGCTGACTTAATAAATATTTGCCTGGTGTAAATATGGCGGGAGGATTCGCCCTAGTGG  
TCAAACATTAAATATTAATGGAATGGGTATGCTGAAGATGTTAGAGTTCAACTAGACGGCGCAA  
CAAAAGTTCGAAAAATATCAACAAGGCTCTATTTATTGAACCTGAGTTATTAAGAAAGGTG  
ACAGTAGACAAAGGAAATTATTCTCCTCAATATGGCAATGGTGGCTTGCTGGTACTGTAAGAATT  
TGAAACAAAAGATGCAACTGATTTTGAAAGAAAATCAGAAAATAGGTGGATTATTAATATG  
GAAATAATAGCAATAATAACCAAAAAACTTATAGTACAGCCCTAGTTACAGAATGAACAAAAA  
AATATTGATTTGTTATTATTGGTTCTGTAAGAAATGCAAGCAATTATAACAAGACCTGATAAAAG  
TAAAATTCTTTCAAAAACAATCAAAAAGTGGATTAAATAAAAGTAAATTGGCAAATTACTC  
CTGAACATTTATTAAACTTATCCAGTGTATGGCATTCAAAAGGGTGGGACCTGGCAGCA  
AAAAGAGATGTGATGTCGAGACCAACAGAAACAGAAATAAACACTATGGGATTGATGTTGCGTG  
GAAACGTAACCTGTTATCGAGATCAAAAAGATGAAAGTTATTGATTGAAATATCGCTATTAC  
CTGAAAATAATAAGTGGATTAAATTATCTGTCAGCTGAGTTATAGTAAAACAGAACAGAACATGAT  
ACTCGCCATGAGAAAGTCACCTCTTCAATTCTAGGTACATTAGGAAATAAAAGTTGGATAACTTA  
TTCAGATCTTACTTTGATATAAGTAACACAAGTACTCTAAATATTGGCGTGCTGAGCATGAAC  
TACTATTGGTTACAGTGGTAAAAAATAAAAGAAATACCCTATGTATCATAAAGGGGAGTC  
AAGAAGGCAGACTATAATTATGGCTATTTCAGCCTTATTATATGCCTCTGGACGCCAGTATAAC  
ACAAGCATTATTACAAGATCAAATAAAATGGCAGAATTTCCTCTTACAGGAGGGATAAGAT  
ATGACCATATCAATAATATAGGGCAGAAAATTTAGCGCCACGATATAATGATATCTGCAGGA  
CATGATTATGCCAGAAAATTATAATGGTTGGTCTTATTATTAGGTCTTAAGTATGATGTAAA  
TCATTATTAAAGTTATTACGAATTTCAGTAAACTGGCAGGCCCTGTTATTGATGAACAGT  
ATGAGACACAATATAGTCAGCTCTGTATCTGCGACTCTTAAATTAGAAAAGAAATGATT  
AATCAAACCAAGAGTGGTGBAATTATTACTCTCAATCTATTTCAGGAAATGATGCTTTCA  
ATTAGAAACTACTATTTCAGCAGGAAATTCAGGAGGGATAAGT  
GCCTAGAAAATGCTTATGTTAGATGTTGATAATAGTGTGCTTAAATTAGTAATTACGTAAT  
TTACCTGGTTATGTTATTCAAGGAGGGATTAGAAGCTTATTATCAATCATCGTATTATTGG  
TGGACTGACATATTCTTATGTAAGGAAAACCGCATACTCACCAAGAAATCCATGGAGTAAA  
CATCTACATGGATCGCAGAAAACATCACCTAGAAAAGCAATCGCTACTTTAGGTTAATATTCCG  
GAATATTATTACGGCAGGTTGGCGTGTGAGTTGAGAAAGCAAGATAGATCACCACATC  
TAATGATTCTAAAGCATCATATTGGCATTACCTCTTCAAAAGGATATAGCCTACATAGTGTAT  
TCTTCTTGGAGTCCTACAAAGATTAAGGAATGAATTCAAAGTTACTGTTGATAATTATT  
AACCGACCCTATTATCCTTACTTAGGAGAATTAGCTTCAGGAACAGGTAGGAATGTCAAATTAG  
CCTGACTCAGCAATTAA

FIG. 14-2.

SEQ ID NO:14

MDKNLMKGCVFLSIVGCGIQIGLASNPNNPPDVDELLPIIVNADEDNKLPGRSVLKQKNIDQQQAD  
NAADLINILPGVNMAGGFRPSGQTLNINMGDAEDVRVQLDGATKSFEKYQQGSIFIEPELLRKV  
TVDKGNYSPOYGNGGFAGTVKFETKDATDFLKENQKIGGLFKYGNNSNNNQKTYSTALVLQNEQK  
NIDLLLFGSVRNASNYTRPDKSKILFSKNNQKSGLIKVNWQITPEHLLTLSVYGIHKGWEPWAA  
KRDVMSRPTETEIKHYGIDVAWKRKLVYRDQKDESYSKYRYLPENNWKWINLSVQLSYSKTEQND  
TRHEKVTSFLGTLGNKSWITYSDLTFDISNTSTLNIGRAEHELLFGLQWLKNKRNTLMYHKGGV  
KKADYNYGYFQPYMPSGRQYTQAFYLQDQIKWQNFLTGGIRYDHINNIQQKNLAPRYNDISAG  
HDYSQKNYNGWSYYLGLKYDVNHYLSLFTNFSKTWRAPVIDEQYETQYSQASVSATSILNLEKEMI  
NQTRVGGIITLNHLFQENDAFQFRRTYFYNRGKNEIFKTRGVNCVENALDVDNSVCPKIISNYRN  
LPGYVIQGAELEAAYYQSSYLFGGLTYSYVKGKRDTSPRNPWSKTSTWIAETSPRKAIATLGFNIP  
EYYFTAGWRAEFVRKQDRSPLSNDSKASYWALPSSKGYSLHSVFFSWSPTKIKGMNFKVTVVDNL  
NRPYYPYLGEASGTGRNVKFSLTQQF\*

FIG. 15-1.

FIG. 15-2.

WP\_053521090.1 ILPNSKGYLPLDYKERDLNTKTKQINLDDLTKAFTLFEIENELSYGGVYAKTTKEMVNKAG  
1121\_Hgba\_968aa ILPNSKGYLPLDYKERDLNTKTKQINLDDLTKAFTLFEIENELSYGGVYAKTTKEMVNKAG  
WP\_005756141.1 ILPNSKGYLPLDYKERDLNTKTKQINLDDLTKAFTLFEIENELSYGGVYAKTTKEMVNKAG  
AAQ14873.1 ILPNSQGYLPLDYKERDLNTKTKQINLDDLTKAFTLFEIENELSYGGVYAKTTNGVNVKAG  
WP\_061405928.1 ILPNSQGYLPLDYKERDLNTKTKQINLDDLTKAFTLFEIENELSYGGVYAKTTKEMVNKAG  
\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:  
  
WP\_053521090.1 YYGPNPTWWAERTLQGDLSGNQHNCNTNSSYNGMLCPRHEPLTSFLIPVEATTKSLYFSD  
1121\_Hgba\_968aa YYGRNPTWWAERTLQGSW-GKLRECKTSSSYNGMLCPRHEPLTSFLIPVEATTKSLYFAD  
WP\_005756141.1 YYGRNPTWWAERTLQGSW-GKLRECKTSSSYNGMLCPRHEPLTSFLIPVEATTKSLYFAD  
AAQ14873.1 YYGRNPTWWAERTLQGSWNGTLRECKTSSSYNGMLCPRHEPLTSFLIPVEATTKSLYFAD  
WP\_061405928.1 YYGRNPTWWAERTLQGSWNGTLRECKTSSSYNGMLCPRHEPLTSFLIPVEATTKSLYFAD  
\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:  
  
WP\_053521090.1 NIKLNHNMLSVDLGYRYDDIKYQPEYIIPGVTPKIADDMVKGJIFIPLPKGEKIKIGNYETTK  
1121\_Hgba\_968aa NIKLNHNMLSVDLGYRYDDIKYQPEYIIPGVTPKIADDMVKGJIFIPLPKGEKIKIGNYETTK  
WP\_005756141.1 NIKLNHNMLSVDLGYRYDDIKYQPEYIIPGVTPKIADDMVKGJIFIPLPKGEKIKIGNYETTK  
AAQ14873.1 NIKLNHNMLSVDLGYRYDDIKYQPEYIIPGVTPKIADDMVKGJIFIPLPKGEKIKIGNYETTK  
WP\_061405928.1 NIKLNHNMLSVDLGYRYDDIKYQPEYIIPGVTPKIADDMVKGJIFIPLPKGEKIKIGNYETTK  
\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:  
  
WP\_053521090.1 PLTPEQIRKNAEENIAYIAQEKRFKKHSYSLGATFDPLNFLRVQVKYSKGFRAPTSDELY  
1121\_Hgba\_968aa PLTQAOQIRKNAEENIAYIAQEKRFKKHSYSLGATFDPLNFLRVQVKYSKGFRAPTSDELY  
WP\_005756141.1 PLTQAOQIRKNAEENIAYIAQEKRFKKHSYSLGATFDPLNFLRVQVKYSKGFRAPTSDELY  
AAQ14873.1 PLTPEQIRKNAEENIAYIAQGKRFKKHSYSLGATFDPLNFLRVQVKYSKGFRAPTSDELY  
WP\_061405928.1 PLTPEQIRKNAEENIAYIAQEKRFKKHSYSLGATFDPLNFLRVQVKYSKGFRAPTSDELY  
\*\*\* \*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:  
  
WP\_053521090.1 FTFKHPDPDTILPNPVLKPEEAKNQEIALTVHDNWGFVSTSVFQTQYRHFIDLAYLGSRNLL  
1121\_Hgba\_968aa FTFKHPDPDTILPNPVLKPEEAKNQEIALTVHDNWGFVSTSVFQTQYRHFIDLAYLGSRNLL  
WP\_005756141.1 FTFKHPDPDTILENPVLKPEEAKNQEIALTVHDNWGFVSTSVFQTQYRHFIDLAYLGSRNLL  
AAQ14873.1 FTFKHPDPDTILENPVLKPEEAKNQEIALTVHDNWGFVSTSVFQTQYRHFIDLAYLGSRNLL  
WP\_061405928.1 FTFKHPDPDTILENPVLKPEEAKNQEIALTVHDNWGFVSTSVFQTQYRHFIDLAYLGSRNLL  
\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:  
  
WP\_053521090.1 SNSVGGQAQARDFQVYQNWNVDNAVKVGLEINARLNLYFWHVLDFGNTSYKFTYQRGRLL  
1121\_Hgba\_968aa SNSVGGQAQARDFQVYQNWNVDNAVKVGLEINARLNLYFWHVLDFGNTSYKFTYQRGRLL  
WP\_005756141.1 SNSVGGQAQARDFQVYQNWNVDNAVKVGLEINARLNLYFWHVLDFGNTSYKFTYQRGRLL  
AAQ14873.1 SNSVGGQAQARDFQVYQNWNVDNAVKVGLEINARLNLYFWHVLDFGNTSYKFTYQRGRLL  
WP\_061405928.1 SNSVGGQAQARDFQVYQNWNVDNAVKVGLEINARLNLYFWHVLDFGNTSYKFTYQRGRLL  
\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:  
  
WP\_053521090.1 DGDRPMNAIQPKASVFGLYDHKENKFGADLYITRVSEKKAKDTONMFYKEEKKKDSAIH  
1121\_Hgba\_968aa DGDRPMNAIQPKASVFGLYDHKENKFGADLYITRVSEKKAKDTONMFYKEQGYKDSAVR  
WP\_005756141.1 DGDRPMNAIQPKASVFGLYDHKENKFGADLYITRVSEKKAKDTONMFYKEQGYKDSAVR  
AAQ14873.1 DGDRPMNAIQPKASVFGLYDHKENKFGADLYITRVSEKKAKDTONMFYKEQGYKDSAIH  
WP\_061405928.1 DGDRPMNAIQPKASVFGLYDHKENKFGADLYITRVSEKKAKDTONMFYKEQGYKDSAVR  
\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:  
  
WP\_053521090.1 WRSDDYTLDVFVTYIKPVKNVTLQFGIYNLTDKRKYLTWESARS1KPGTSNLINQQTGAG  
1121\_Hgba\_968aa WRSDDYTLDVDAVGYIKPIK1NLTQFGVYNLTDKRKYLTWESARS1KPGTSNLINQQTGAG  
WP\_005756141.1 WRSDDYTLDVDAVGYIKPIK1NLTQFGVYNLTDKRKYLTWESARS1KPGTSNLINQQTGAG  
AAQ14873.1 WRSDDYTLDVDAVGYIKPIK1NLTQFGVYNLTDKRKYLTWESARS1KPGTSNLINQQTGAG  
WP\_061405928.1 WRSDDYTLDVDAVGYIKPIK1NLTQFGVYNLTDKRKYLTWESARS1KPGTSNLINQQTGAG  
\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:  
  
WP\_053521090.1 INRFYSPGRNYKLSAEITF-  
1121\_Hgba\_968aa INRFYSPGRNFKFSAEITF\*  
WP\_005756141.1 INRFYSPGRNFKFSAEITF-  
AAQ14873.1 INRFYSPGRNFKFSAEITF-  
WP\_061405928.1 INRFYSPGRNFKFSAEITF-  
\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*

FIG. 16-1.

SEQ71136.1 WP_014391205.1 WP_016534044.1 1121_FepA_790aa WP_005756883.1	MEKIDMESAKNPLKKTTLALLCCSTAFSLSAKTDTNADKNHFLTEIVVYADQNKSMSSTO -----MESAKNPLKKTTLALLCCSTAFSLSAKTDTNADKNHFLTEIVVYADQNKSMSSTO -----MEYAKNPLKKTTLALLCCSTAFSLSAKTDTNADKNHFLTEIVVYADQNKSMSSTO -----MESAKNPLKKTTLALLCCSTAFSLSAKTDTNADKNHFLTEIVVYADQNKSMSSTO -----MESAKNPLKKTTLALLCCSTAFSLSAKTDTNADKNHFLTEIVVYADQNKSMSSTO *** ****; ****;
SEQ71136.1 WP_014391205.1 WP_016534044.1 1121_FepA_790aa WP_005756883.1	SVTQDEDMKKSPVTNGNITDYLRSNPHVRYENSQNGLQRGEIKEPKENISINGADYQQTFFF SVTQDEDMKKSPVTNGNITDYLRSNPHVRYENSQNGLQRGEIKEPKENISINGADYQQTFFF SVTQDEDMKKSPVTNGNITDYLRSNPHVRYENSQNGLQRGEIKEPKENISINGADYQQTFFF SVTQDEDMKKSPVTNGNITDYLRSNPHVRYENSQNGLQRGEIKEPKENISINGADYQQTFFF SVTQDEDMKKSPVTNGNITDYLRSNPHVRYENSQNGLQRGEIKEPKENISINGADYQQTFFF ***; ****;
SEQ71136.1 WP_014391205.1 WP_016534044.1 1121_FepA_790aa WP_005756883.1	VDNVNINNDMGMGSDFDGTMATVPFANHSQGYFFDANLSSIVVHDNSVSASLGGFAGG VDNVNINNDMGMGSDFDGTMATVPFANHSQGYFFDANLSSIVVHDNSVSASLGGFAGG VDNVNINNDMGMGSDFDGTMATVPFANHSQGYFFDANLSSIVVHDNSVSASLGGFAGG VDNVNINNDMGMGSDFDGTMATVPFANHSQGYFFDANLSSIVVHDNSVSASLGGFAGG VDNVNINNDMGMGSDFDGTMATVPFANHSQGYFFDANLSSIVVHDNSVSASLGGFAGG ****;
SEQ71136.1 WP_014391205.1 WP_016534044.1 1121_FepA_790aa WP_005756883.1	AVVAKTKQYDGKDQLKFYSRITDASWAKFKVEDKDLERFKNAIPEGSVAEFQPKYSKHFF AVVAKTKQYDGKDQLKFYSRITDASWAKFKVEDKDLERFKNAIPEGSVAEFQPKYSKHFF AVVAKTKQYDGKDQLKFYSRITDASWAKFKVEDKDLERFKNAIPEGSVAEFQPKYSKRFF AVVAKTKQYDGKDQLKFYSRITDASWAKFKVEDKDLERFKNAIPEGSVAEFQPKYSKHFF AVVAKTKQYDGKDQLKFYSRITDASWAKFKVEDKDLERFKNAIPEGSVAEFQPKYSKHFF ****;
SEQ71136.1 WP_014391205.1 WP_016534044.1 1121_FepA_790aa WP_005756883.1	NTAEKGLSENLCMVGILSRRTSSIQQSRQINPQGDRDKQTHTRRSNDALLNFNLTPNDK NTAEKGLSENLCMVGILSRRTSSIQQSRQINPQGDRDKQTHTRRSNDALLNFNLTPNDK NTAEKGLSENLCMVGILSRRTSSIQQSRQINPQGDRDKQTHTRRSNDALLNFNLTPNDK NTAEKGLSENLCMVGILSRRTSSIQQSRQINPQGDRDKQTHTRRSNDALLNFNLTPNDK ****;
SEQ71136.1 WP_014391205.1 WP_016534044.1 1121_FepA_790aa WP_005756883.1	HRFELGFRYSNYRERKFFNTNIDSNVFDYHRAYGVTFSWINALQSGILTTLAYDNFDDT HRFELGFRYSNYRERKFFNTNIDSNVFDYHRAYGVTFSWINALQSGILTTLAYDNFDDT HRFELGFRYSNYRERKFFNTNIDSNVFDYHRAYGVTFSWINALQSGILTTLAYDNFDDT HRFELGFRYSNYRERKFFNTNIDSNVFDYHRAYGVTFSWINALQSGILTTLAYDNFDDT ****;
SEQ71136.1 WP_014391205.1 WP_016534044.1 1121_FepA_790aa WP_005756883.1	RKSASTSMKTI -EDENDYTLGGMGNSQLNQKNSHFSLEYAMNSFDLSHINHSISLGSVF RKSASTYMKTTLDDGEEYTEGGMGNSQLNQKNLHTSLEYAMNPFNLSIEHSVSLGGIY RKSASTYMKTTLTEEGEEYTEGGMGNSQLNQKNLHTSLEYAMNPFNLSIEHSVSLGGIY RKSASTYMKTTLTEEGEEYTEGGMGNSQLNQKNLHTSLEYAMNPFNLSIEHSVSLGGIY ****;
SEQ71136.1 WP_014391205.1 WP_016534044.1 1121_FepA_790aa WP_005756883.1	QATKYRFTRHSDAEAEIINRIDLEN-----EKIEIKSSNLAKKGTVKTRYQNIALYVED QATKYRFTRHSDAVGELYTPNWLDDNNNSNDIYDELTIAQRNIAKKGTVKTRYQNIALYVED QATKYRFTRHSDAVGELYTPDWLNNGNT---DKLILTORNIAKKGTVKTRYQNIALYVED QATKYRFTRHSDAVGELYTPDWLNNGNT---DKLILTORNIAKKGTVKTRYQNIALYVED QATKYRFTRHSDAVGELYTPDWLNNGNT---DKLILTORNIAKKGTVKTRYQNIALYVED *;*** .** . *; . *; . *; ****;
SEQ71136.1 WP_014391205.1 WP_016534044.1 1121_FepA_790aa WP_005756883.1	LMTWKNLEFRAGLRLERDDYLKNTNLAPRTVPRYKPFEDTAFSFGWNRYYGRSFASMKLS LMTWKNLEFRAGLRLERDDYLKNTNLAPRTVPRYKPFEDTAFSFGWNRYYGRSFASMKLS LMTWKNLEFRAGLRLERDDYLKNTNLAPRTVPRYKPFEDTAFSFGWNRYYGRSFASMKLS LMTWKNLEFRAGLRLERDDYLKNTNLAPRTVPRYKPFEDTAFSFGWNRYYGRSFASMKLS ****;

FIG. 16-2.

ESQ71136.1  
WP\_014391205.1  
WP\_016534044.1  
1121\_FepA\_790aa  
WP\_005756883.1

EGIFKLDGHDTFRYKDLSQFKTPYSDELSFGVEQYVANLAFHLKYILRDNKQRIVLQEED  
EGIFKLDGHDTFRYKDLSQFKTPYSDELSFGVEQYVANLAFHLKYILRDNKQRIVLQEED  
EGIFKLDGHDTFRYKDLSQFKTPYSDELSFGVEQYVANLAFHLKYILRDNKQRIVLQEED  
EGIFKLDGHDTFRYKDLSQFKTPYSDELSFGVEQYVANLAFHLKYILRDNKQRIVLQEED  
EGIFKLDGHDTFRYKDLSQFKTPYSDELSFGVEQYVANLAFHLKYILRDNKQRIVLQEED  
\*\*\*\*\*;

ESQ71136.1  
WP\_014391205.1  
WP\_016534044.1  
1121\_FepA\_790aa  
WP\_005756883.1

VMLNGEKKKLRYYQRGKDYKTNVLTFQINTQAPWELGPTRWTSAVAFDWLDSSKAIDHGRG  
VMLNGEKKKLRYYQRGKDYKTNVLTFQISTQAPWELGPTRWTSALAFDWLDSSKAIDHGRG  
VMLNGERKKKLRYYQRGKDYKTNVLTFQINTQAPWELGPTRWTSAVAFDWLDSSKAIDHGRG  
VMLNGERKKKLRYYQRGKDYKTNVLTFQINTQAPWELGPTRWTSAVAFDWLDSSKAIDHGRG  
VMLNGEKKKLRYYQRGKDYKTNVLTFQINTQAPWELGPTRWTSAVAFDWLDSSKAIDHGRG  
\*\*\*\*\*;

ESQ71136.1  
WP\_014391205.1  
WP\_016534044.1  
1121\_FepA\_790aa  
WP\_005756883.1

YNGSTPVILDGKLMTYEQMLRKVNAYKETWGLRLNLDMFVPRFDLSWANTIVVKPPTTLT  
YNGSTPVILDGKLMTYEQMLRKVNAYKETWGLRLNLDMFVPKFDSLWANTIVVKPPTTLT  
YNGSTPVILDGKLMTYEQMLRKVNAYKETWGLRLNLDMFVPRFDLSWANTIVVKPPTTLT  
YNGSTPVILDGKLMTYEQMLRKVNAYKETWGLRLNLDMFVPIFDSLWANTIVVKPPTTLT  
YNGSTPVILDGKLMTYEQMLRKVNAYKETWGLRLNLDMFVPRFDLSWANTIVVKPPTTLT  
\*\*\*\*\*;

ESQ71136.1  
WP\_014391205.1  
WP\_016534044.1  
1121\_FepA\_790aa  
WP\_005756883.1

ERVSSNTPEVYRSYDYGTYTQWDTSILRWQPTFAEKHRPYIKLDVLNVLNKTFRKGAGPNGQ  
ERVSSNTPEVYRSYDYGTYTQWDTSILRWQPTFAEKHRPYIKLDVLNVLNKTFRKGAGPNGQ  
ERVSSNTPEVYRSYDYGTYTQWDTSILRWQPTFAEKHRPYIKLDVLNVLNKTFRKGAGPNGQ  
ERVSSNTPEVYRSYDYGTYTQWDTSILRWQPTFAEKHRPYIKLDVLNVLNKTFRKGAGPNGQ  
EHVSSSTPEVYRSYDYGTYTQWDTSILRWQPTFAEKHRPYIKLDVLNVLNKTFRKGAGPNGQ  
\*\*\*\*\*;

ESQ71136.1  
WP\_014391205.1  
WP\_016534044.1  
1121\_FepA\_790aa  
WP\_005756883.1

DLGIYTPGREFWLEVGYEF  
DLGIYTPGREFWLEVGYEF  
DLGIYTPGREFWLEVGYEF  
DLGIYTPGREFWLEVGYEF  
DLGIYTPGREFWLEVGYEF  
\*\*\*\*\*

FIG. 17-1.

WP\_071522773.1 MSFKHKTALFVVAHACCTSVAENAATTLEPIVVSDSHTTLNLQDNKLEKESPKDLKAI  
WP\_016534444.1 MSFKHKTALFVVARACCTSALAENAATTLEPIVVSELSTSHTTLNLQDNKLEKESPKDLKAI  
WP\_005755819.1 MSFKHKTALFVVAHACCTSALAENVATTLEPIVVSDSHTTLNLQDNKLEKESPKDLKAI  
1121\_PfhR\_727aa MSFKHKTALFVVAHACCTSALAENAATTLEPIVVSDSHTTLNLQDNKLEKESPKDLKAI  
EGP05580.1 \*\*\*\*\*;\*\*\*\*\*;\*\*\*\*\*;\*\*\*\*\*;\*\*\*\*\*;\*\*\*\*\*;\*\*\*\*\*;\*\*\*\*\*;  
  
WP\_071522773.1 FATTPNINVINTGHAQLGDIIEIRGMGSSREIFATGANRVTMELDGMMDISPSFYFGHSSRM  
WP\_016534444.1 FTTTPNINVINTGHAQLGDIIEIRGMGSSREIFATGANRVTMELDGMMDISPSFYFGHSSRM  
WP\_005755819.1 FATTPNINVINTGHAQLGDIIEIRGMGSSREIFATGANRVTMELDGMMDISPSFYFGHSSRM  
1121\_PfhR\_727aa FATTPNINVINTGHAQLGDIIEIRGMGSSREIFATGANRVTMELDGMMDISPSFYFGHSSRM  
EGP05580.1 \*;\*\*\*\*\*;\*\*\*\*\*;\*\*\*\*\*;\*\*\*\*\*;\*\*\*\*\*;\*\*\*\*\*;\*\*\*\*\*;  
  
WP\_071522773.1 GROYFDPSDLKRVEVKGPNSQGVAGHVRFQTKDPHDYLLPNQRTGAQLRAGYLGDSDAY  
WP\_016534444.1 GROYFDPSDLKRVEIKGPNSQGVAGHVRFQTKDPHDYLLPNQRTGAQLRAGYLGDSDAY  
WP\_005755819.1 GROYFDPSDLKRVEIKGPNSQGVAGHVRFQTKDPHDYLLPNQRTGAQLRAGYLGDSDAY  
1121\_PfhR\_727aa GROYFDPSDLKRVEIKGPNSQGVAGHVRFQTKDPHDYLLPNQRTGAQLRAGYLGDSDAY  
EGP05580.1 \*\*\*\*\*;\*\*\*\*\*;\*\*\*\*\*;\*\*\*\*\*;\*\*\*\*\*;\*\*\*\*\*;\*\*\*\*\*;  
  
WP\_071522773.1 YVGITGAALLDEHSSALVSYTRRWNFNEFFNNKGGLDVGSQRTKSNPSSGYSNAVNSKLRY  
WP\_016534444.1 YVGITGATLLDEHSSALVSYTRRWNFNEFFNNKGGLDVGSQRTKSNPSSGYSNAVNSKLRY  
WP\_005755819.1 YVGITGATLLDEHSSALVSYTRRWNFNEFFNNKGGLDVGSQRTKSNPSSGYSNAVNSKLRY  
1121\_PfhR\_727aa YVGITGATLLDEHSSALMSYTRRWNFNEFFNNKGGLDVGSQRTKSNPSSGYSNAVNSKLRY  
EGP05580.1 \*\*\*\*\*;\*\*\*\*\*;\*\*\*\*\*;\*\*\*\*\*;\*\*\*\*\*;\*\*\*\*\*;\*\*\*\*\*;  
  
WP\_071522773.1 SPNDRHKFTLNLQHYDLKRATAYLEDLSLGTTFTRRGTKTVHHNTNIQKNQRHIAFSHDMQ  
WP\_016534444.1 SPNDRHKFTLNLQHYDLKRATAYLEDLSLGTTFTRRGTKTVHHNTNIQKNQRHIAFSHDMQ  
WP\_005755819.1 SPNDRHKFTLNLQHYDLKRATAYLEDLSLGTTFTRRGTKTVHHNTNIQKNQRHIAFSHDMQ  
1121\_PfhR\_727aa SPNDRHKFTLNLQHYDLKRATAYLEDLSLGTTFTRRGTKTVHHNTNIQKNQRHIAFSHDMQ  
EGP05580.1 \*\*\*\*\*;\*\*\*\*\*;\*\*\*\*\*;\*\*\*\*\*;\*\*\*\*\*;\*\*\*\*\*;\*\*\*\*\*;  
  
WP\_071522773.1 QTNHSIFDHLHWQIALQQTSTSRTNTGAVTSTSCTSGSEPPSTPKFSQERSLGFKTKTISLK  
WP\_016534444.1 QTT-AFFDHLHWQIALQQTSTSRTNTGAVTSTSCTSAPPPSTPKFSQERSLGFKTKTISLK  
WP\_005755819.1 QTT-AFFDHLHWQIALQQTSTSRTNTGAVTSTSCTSAPPPSTPKFSQERSLGFKTKTISLK  
1121\_PfhR\_727aa QTT-AFFDHLHWQIALQQTSTSRTNTGAVTNTSCTSAPPPSTPKFSQERSLGFKTKTISLK  
EGP05580.1 QTT-AFFDHLHWQIALQQTSTSRTNTGAVTNTSCTSAPPPSTPKFSQERSLGFKTKTISLK  
\*\*\*\*\*;\*\*\*\*\*;\*\*\*\*\*;\*\*\*\*\*;\*\*\*\*\*;\*\*\*\*\*;  
  
WP\_071522773.1 TEFNKS LGQHVHRELHYGLKLQYSQMQLRQTQSLNEQGSNTRTSAFFPTQQWQSKLHL  
WP\_016534444.1 TEFNKS LGQHVHRELHYGLKLQYSQMQLRQTQSLNEQGSNTRTSAFFPTQQWQSKLHL  
WP\_005755819.1 TEFNKS LGQHVHRELHYGLKLQYSQMQLRQTQSLNEQGSNTRTSAFFPTQQWQSKLHL  
1121\_PfhR\_727aa TEFNKS SIGQHVHRELHYGLKLQYSQMQLRQTQSLNEQGSNTRTSAFFPTQQWQSKLHL  
EGP05580.1 TEFNKS SIGQHVHRELHYGLKLQYSQMQLRQTQSLNEQGSNTRTSAFFPTQQWQSKLHL  
\*\*\*\*\*;\*\*\*\*\*;\*\*\*\*\*;\*\*\*\*\*;\*\*\*\*\*;  
  
WP\_071522773.1 SDRISFGKSGSLSLTPSIHLTQIRIKPKTENVSKKNREQLFTYKDTAIGYGLRVDYALNEA  
WP\_016534444.1 SDRISFGKSGSLSLTPSIHLTQIRIKPKTENVSKKNREQLFTYKDTAIGYGLRVDYALNEA  
WP\_005755819.1 SDRISFGKSGSLSLTPSIHLTQIRIKPKTENVSKKNREQLFTYKDTAIGYGLRVDYALNEA  
1121\_PfhR\_727aa SDRISFGKSGSLSLTPSIHLTQIRIKPKTENVSKKNREQLFTYKDTAIGYGLRVDYALNEA  
EGP05580.1 SDRISFGKSGSLSLTPSIHLTQIRIKPKTENVSKKNREQLFTYKDTAIGYGLRVDYALNEA  
\*\*\*\*\*;\*\*\*\*\*;\*\*\*\*\*;\*\*\*\*\*;

FIG. 17-2.

WP\_071522773.1  
WP\_016534444.1  
WP\_005755819.1  
1121\_PfhR\_727aa  
EGP05580.1

NILSINYQHATRLPGYGENNAQSYGHWPAKPNPHLQPETSDGIELSWRSAGAIGQQTTI  
NILSINYQHATRLPGYGENNAQSYGHWPAKPNPHLQPETSDGVIELSWRSAGAIGQQTTI  
NILSINYQHATRLPGYGENNAQSYGHWPAKPNPHLQPETSDGIELSWRSAGAIGQQTTI  
NILSINYQHATRLPGYGENNAQSYGHWPAKPNPHLQPETSDGIELSWRSAGAIGQQTTI  
NILSINYQHATRLPGYGENNAQSYGHWPAKPNPHLQPETSDGIELSWRSAGAIGQQTTI  
\*\*\*\*\*;

WP\_071522773.1  
WP\_016534444.1  
WP\_005755819.1  
1121\_PfhR\_727aa  
EGP05580.1

FYNRYNDLIYLDTTACYADRTGQVCPDLANEKGRSYSYGYIEFDGKLNLDTIGFAQGTY  
FYNRYNDLIYLDTTACYADRTGQ--VPCDLANEKGRSYSYGYIEFDGKLNLDTIGFAQGTY  
FYNRYNDLIYLDTTACYADRTGQ--VPCDLANEKGRSYSYGYIEFDGKLNLDTIGFAQGTY  
FYNRYNDLIYLDTTACYADRTGQ--VPCDLANEKGRSYSYGYIEFDGKLNLDTIGFAQGTY  
FYNRYNDLIYLDTTACYADRTGQ--VPCDLANEKGRSYSYGYIEFDGKLNLDTIGFAQGTY  
\*\*\*\*\*;

WP\_071522773.1  
WP\_016534444.1  
WP\_005755819.1  
1121\_PfhR\_727aa  
EGP05580.1

LNAGFAYSKGKTANKQPQGRLDPLTFVGVLGYQQPMWDVWVGIEGKLKFAAKKKTKDLPANQ  
LNAGFAYSKGKTANKQPQGRLDPLTFVGVLGYQQPMWDVWVGIEGKLKFAAKKKTKDLPANQ  
LNAGFAYSKGKTANKQPQGRLEPLTGFVGLGYQQPMWDVWVGIEGKLKFAAKKKTKDLPANQ  
LNAGFAYSKGKTANKQPQGRLEPLTGFVGLGYQQPMWDVWVGIEGKLKFAAKKKTKDLPANQ  
LNAGFAYSKGKTANKQPQGRLEPLTGFVGLGYQQPMWDVWVGIEGKLKFAAKKKTKDLPANQ  
\*\*\*\*\*;

WP\_071522773.1  
WP\_016534444.1  
WP\_005755819.1  
1121\_PfhR\_727aa  
EGP05580.1

GFEGLPGYAVVDLTAYYNTKQLYLGIGIYNVLDKKYARWAMARGDIKHGNYDKHTEAGR  
GFEGLPGYAVVDLTAYYNTKQLYLGIGIYNVLDKKYARWAMARGDIKHGNYDKHTEAGR  
GFEGLPGYAVVDLTAYYNTKQLYLGIGIYNVLDKKYARWAMARGDIKHGNYDKHTEAGR  
GFEGLPGYAVVDLTAYYNTKQLYLGIGIYNVLDKKYARWAMARGDIKHGNYDKHTEAGR  
GFEGLPGYAVVDLTAYYNTKQLYLGIGIYNVLDKKYARWAMARGDIKHGNYDKHTEAGR  
\*\*\*\*\*;

WP\_071522773.1  
WP\_016534444.1  
WP\_005755819.1  
1121\_PfhR\_727aa  
EGP05580.1

HFGANIRYHF  
HFGANIRYHF  
HFGANIRYHF  
HFGANIRYHF  
HFGANIRYHF  
\*\*\*\*\*

FIG. 18-1.

1135_PM0300_964aa WP_005753642.1 WP_016534557.1 WP_010906573.1 AAQ14873.1 1121_HgbA_968aa	MRTTTIKFSAITLALLSYCGVILADSHQEATELDITVSSQQDEMNIKEKKVGETVKTAS MRTTTIKFSAITLALLSYCGVILADSHQEATELDITVSSQQDEMNIKEKKVGETVKTAS MRTTTIKFSAITLALLSYCGAILADSHQEATELDITVSSQQDEMNIKEKKVGETVKTAS MRTTTIKFSAITLALLSYCGVILADSHQEATELDITVSSQQDEMNIKEKKVGETVKTAS MRTTTIKFSAITLALLSYCGAILADSHQEATELDITVSSQQDEMNIKEKKVGETVKTAS *****
1135_PM0300_964aa WP_005753642.1 WP_016534557.1 WP_010906573.1 AAQ14873.1 1121_HgbA_968aa	QLKRQQVQDSRDLVRYETGVTVVEAGRFGSSGYAIRGVDENRAITVDGLHQAEETLSSQG QLKRQQVQDSRDLVRYETGVTVVEAGRFGSSGYAIRGVDENRAITVDGLHQAEETLSSQG QLKRQQVQDSRDLVRYETGVTVVEAGRFGSSGYAIRGVDENRAITVDGLHQAEETLSSQG QLKRQQVQDSRDLVRYETGVTVVEAGRFGSSGYAIRGVDENRAITVDGLHQAEETLSSQG QLKRQQVQDSRDLVRYETGVTVVEAGRFGSSGYAIRGVDENRAITVDGLHQAEETLSSQG *****
1135_PM0300_964aa WP_005753642.1 WP_016534557.1 WP_010906573.1 AAQ14873.1 1121_HgbA_968aa	FKELFEGYGNFNTRNSVEIETLKVAKIAKGADSVKVGSGLGGAVLFETKDARDFLTEK FKELFEGYGNFNTRNSVEIETLKVAKIAKGADSVKVGSGLGGAVLFETKDARDFLTEK FKELFEGYGNFNTRNSVEIETLKVAKIAKGADSVKVGSGLGGAVLFETKDARDFLTEK FKELFEGYGNFNTRNSVEIETLKVAKIAKGADSVKVGSGLGGAVLFETKDARDFLTEK FKELFEGYGNFNTRNSVEIETLKVAKIAKGADSVKVGSGLGGAVLFETKDARDFLTEK *****
1135_PM0300_964aa WP_005753642.1 WP_016534557.1 WP_010906573.1 AAQ14873.1 1121_HgbA_968aa	DWHIGYKAGYSTADNQGINAVTLAGRYQMFDALIMHSKRHGHELENIDYRNGRDIQGKER DWHIGYKAGYSTADNQGINAVTLAGRYQMFDALIMHSKRHGHELENIDYRNGRDIQGKER DWHIGYKAGYSTADNQGINAVTLAGRYQMFDALIMHSKRHGHELENIDYRNGRDIQGKER DWHIGYKAGYSTADNQGINAVTLAGRYQMFDALIMHSKRHGHELENIDYRNGRDIQGKER DWHIGYKAGYSTADNQGINAVTLAGRYQMFDALIMHSKRHGHELENIDYRNGRDIQGKER *****
1135_PM0300_964aa WP_005753642.1 WP_016534557.1 WP_010906573.1 AAQ14873.1 1121_HgbA_968aa	EKADPYTITKESTLVKFSSPTENHRTVASDTYIQRSGRHDYSLSQPQSNYFTYDEKE EKADPYTITKESTLVKFSSPTENHRTVASDTYIQRSGRHDYSLSQPQSNYFTYDEKE EKADPYTITKESTLVKFSSPTENHRTVASDTYIQRSGRHDYSLSQPQSNYFTYDEKE EKADPYTITKESTLVKFSSPTENHRTVASDTYIQRSGRHDYSLSQPQSNYFTYDEKE EKADPYTITKESTLVKFSSPTENHRTVASDTYIQRSGRHDYSLSQPQSNYFTYDEKE EKADPYTITKESTLVKFSSPTENHRTVASDTYIQRSGRHDYSLSQPQSNYFTYDEKE *****
1135_PM0300_964aa WP_005753642.1 WP_016534557.1 WP_010906573.1 AAQ14873.1 1121_HgbA_968aa	SRHANDLTKRKNVSFTYENYSVTFWDTLKLSSYQQKIRTRARTEDYCDGNEKCDSYKNP SRHANDLTKRKNVSFTYENYSVTFWDTLKLSSYQQKIRTRARTEDYCDGNEKCDSYKNP SRHANDLTKRKNVSFTYENYSVTFWDTLKLSSYQQKIRTRARTEDYCDGNEKCDSYKNP SRHANDLTKRKNVSFTYENYSVTFWDTLKLSSYQQKIRTRARTEDYCDGNEKCDSYKNP SRHANDLTKRKNVSFTYENYSVTFWDTLKLSSYQQKIRTRARTEDYCDGNEKCDSYKNP SRHANDLTKRKNVSFTYENYSVTFWDTLKLSSYQQKIRTRARTEDYCDGNEKCDSYKNP *****
1135_PM0300_964aa WP_005753642.1 WP_016534557.1 WP_010906573.1 AAQ14873.1 1121_HgbA_968aa	LGLQLKDGKIVDPFGNQITLKGTGFNTIEVDKNGNPFPTSGTNNAAFSNNIQLGPKEFW LGLQLKDGKIVDPFGNQITLKGTGFNTIEVDKNGNPFPTSGTNNAAFSNNIQLGPKEFW LGLQLKDGKIVDPFGNQITLKGTGFNTIEVDKDGKPFPTSGTNNAAFSNNLQLSPTGFW LGLQLKDGKIVDPFGNQITLKGTGFNTIEVDKDGKPFPTSGTNNAAFSNNLQLSPTGFW LGLQLKEGKIVDRNGDPVNLLQVLGKHQVVDKAGKPFDTSGTNYAAFSG-KQLGPSPYFW LGLQLKEGKIVDRNGDPVNLLQVLGKHQVVDKAGKPFDTSGTNYAAFSG-KELSPSSFW *****

FIG. 18-2.

1135\_PM0300\_964aa      LDCLSLFDCTQPETVNYQNGQYTFKQ--VELSEEITVNGKLYKTAKEERGVNLYLILPNS  
 WP\_005753642.1      LDCLSLFDCTQPETVNYQNGQYTFKQ--VELSEEITVNGKLYKTAKEERGVNLYLILPNS  
 WP\_016534557.1      LBCTIFDCTKPETVNYQKQNKYEPR--VMLSEEITIDGKLYKTAKEESGVNLYLILPNS  
 WP\_010906573.1      LDCTIFDCTKPETVNYQKQNKYEPR--VMLSEEITIDGKLYKTAKEESGVNLYLILPNS  
 AAQ14873.1      LECTVFDCSKPVTYYKYRYSTETPVKEDIQLNKTMEVNGKTFATYDM-GRERRYIILPNS  
 1121\_HgbA\_968aa      LDCSIFDCSKPINTYKYRYSSEPTLQIQLNKTMEVNGKTFATYDG---RGHYIILPNS  
 \*;\*:\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:  
  
 1135\_PM0300\_964aa      KGYLPLYDYKERDLDSNTKQINLDLTKTFSFNIENELLYGAIYSRTEKKMVNKAGYDGRN  
 WP\_005753642.1      KGYLPLYDYKERDLDSNTKQINLDLTKTFSFNIENELLYGAIYSRTEKKMVNKAGYDGRN  
 WP\_016534557.1      KGYLPLYDYKERDLDSNTKQINLDLTKTFSFNIENELLYGAIYSRTEKKMVNKAGYDGRN  
 WP\_010906573.1      KGYLPLYDYKERDLDSNTKQINLDLTKTFSFNIENELLYGAIYSRTEKKMVNKAGYDGRN  
 AAQ14873.1      QGYLPLDYKERDLNTKQINLDLTCAFTLFEIENELSYGGVIAKTTNGVNVKAGYYGRN  
 1121\_HgbA\_968aa      KGYLPLDYKERDLNTKQINLDLTCAFTLFEIENELSYGGVIAKTTNGVNVKAGYYGRN  
 \*;\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:  
  
 1135\_PM0300\_964aa      PTWWADRLGKST-----NCNYNGLKCPHRHEPLTSFLIPVEATTKSILYFSDNIKLH  
 WP\_005753642.1      PTWWADRLGKST-----NCNYNGLKCPHRHEPLTSFLIPVEATTKSILYFSDNIKLH  
 WP\_016534557.1      PTWWADRLGKST-----NCNYNGLKCPHRHEPLTSFLIPVEATTKSILYFSDNIKLH  
 WP\_010906573.1      PTWWADRLGKST-----NCNYNGLKCPHRHEPLTSFLIPVEATTKSILYFSDNIKLH  
 AAQ14873.1      PTWWAERTLGQSWNGTIRECKTSSSYNGMLCPRHEPLTSFLIPVEATTKSILYFADNIKLH  
 1121\_HgbA\_968aa      PTWWAERTLGQSW-GKLRECKTSSSYNGMLCPRHEPLTSFLIPVEATTKSILYFADNIKLH  
 \*;\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:  
  
 1135\_PM0300\_964aa      NMLSDLGYRYDDIKYQPEYIPGVTPKIADDVKGIFIFIPLPKGEKVTPWGAETYTKPLTQ  
 WP\_005753642.1      NMLSDLGYRYDDIKYQPEYIPGVTPKIADDVKGIFIFIPLPKGEKVTPWGAETYTKPLTQ  
 WP\_016534557.1      NMLSDLGYRYDDIKYQPEYIPGVTPKIADDVKGIFIFIPLPKGEKIKIE-NYETTKPLTP  
 WP\_010906573.1      NMLSDLGYRYDDIKYQPEYIPGVTPKIADDVKGIFIFIPLPKGEKIKIG-NYETTKPLTP  
 AAQ14873.1      NMLSDLGYRYDDIKYQPEYIPGVTPKIADDVKGIFIFIPLPKGEKVTVG-TVVFTKPLTP  
 1121\_HgbA\_968aa      NMLSDLGYRYDDIKYQPEYIPGVTPKIADDVKGIFIFIPLPKGEKVTVG-TMVFTKPLTQ  
 \*;\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:  
  
 1135\_PM0300\_964aa      EQIRKNAEENIAYIAQEKRFKHHSYSLGATFDPLNFLRVQVKYSKGFRAPTSDELYFTFK  
 WP\_005753642.1      EQIRKNAEENIAYIAQEKRFKHHSYSLGATFDPLNFLRVQVKYSKGFRAPTSDELYFTFK  
 WP\_016534557.1      EQIRKNAEENIAYIAQEKRFKHHSYSLGATFDPLNFLRVQVKYSKGFRAPTSDELYFTFK  
 WP\_010906573.1      EQIRKNAEENIAYIAQEKRFKHHSYSLGATFDPLNFLRVQVKYSKGFRAPTSDELYFTFK  
 AAQ14873.1      EQIRKNAEENIAYIAQEKRFKHHSYSLGATFDPLNFLRVQVKYSKGFRAPTSDELYFTFK  
 1121\_HgbA\_968aa      AQIRKNAEENIAYIAQEKRFKHHSYSLGATFDPLNFLRVQVKYSKGFRAPTSDELYFTFK  
 \*;\*\*\*\*\*:  
  
 1135\_PM0300\_964aa      HPDFTILPNPVLPKPEEAKNQEIALTVHDNWGFVSTSVDQKTYRHFIDLAYLGSRNLSNSV  
 WP\_005753642.1      HPDFTILPNPVLPKPEEAKNQEIALTVHDNWGFVSTSVDQKTYRHFIDLAYLGSRNLSNSV  
 WP\_016534557.1      HPDFTILPNPVLPKPEEAKNQEIALTVHDNWGFVSTSVDQKTYRHFIDLAYLGSRNLSNSV  
 WP\_010906573.1      HPDFTILPNPVLPKPEEAKNQEIALTVHDNWGFVSTSVDQKTYRHFIDLAYLGSRNLSNSV  
 AAQ14873.1      HPDFTILPNPVLPKPEEAKNQEIALTVHDNWGFVSTSVDQKTYRHFIDLAYLGSRNLSNSV  
 1121\_HgbA\_968aa      \*;\*\*\*\*\*:  
  
 1135\_PM0300\_964aa      GGQAQARDFQVYQNVDNAVKGLEINARLNLYFWHLDFGNTSYKFTYQCGRLDGDR  
 WP\_005753642.1      GGQAQARDFQVYQNVDNAVKGLEINARLNLYFWHLDFGNTSYKFTYQCGRLDGDR  
 WP\_016534557.1      GGQAQARDFQVYQNVDNAVKGLEINARLNLYFWHLDFGNTSYKFTYQCGRLDGDR  
 WP\_010906573.1      GGQAQARDFQVYQNVDNAVKGLEINARLNLYFWHLDFGNTSYKFTYQCGRLDGDR  
 AAQ14873.1      GGQAQARDFQVYQNVDNAVKGLEINARLNLYFWHLDFGNTSYKFTYQCGRLDGDR  
 1121\_HgbA\_968aa      \*;\*\*\*\*\*:

FIG. 18-3.

1135\_PM0300\_964aa  
WP\_005753642.1  
WP\_016534557.1  
WP\_010906573.1  
AAQ14873.1  
1121\_HgbA\_968aa

PMNAIQPKASVFGLGYDHKENKFGALLYITRVSEKKAKDTYNMFYKEQGYKDSAIRWRSD  
PMNAIQPKASVFGLGYDHKENKFGALLYITRVSEKKAKDTYNMFYKEQGYKDSAIRWRSD  
PMNAIQPKASVFGLGYDHKENKFGALLYITRVSEKKAKDTYNMFYKEQGYKDSAIRWRSD  
PMNAIQPKASVFGLGYDHKENKFGALLYITRVSEKKAKDTYNMFYKEQGYKDSAIRWRSD  
PMNAIQPKASVFGLGYDHKENKFGALLYITRVSEKKAKDTYNMFYKEQGYKDSAIRWRSD  
PMNAIQPKASVFGLGYDHKENKFGALLYITRVSEKKAKDTYNMFYKEQGYKDSAVPWRSD  
\*\*\*\*\*;

1135\_PMC300\_964aa  
WP\_005753642.1  
WP\_016534557.1  
WP\_010906573.1  
AAQ14873.1  
1121\_HgbA\_968aa

DYTLVDAVGYIKPIKNLTLOFGVYNLTDRKYLTWESARSIKPFGTSNLINQKTGAGINRF  
DYTLVDAVGYIKPIKNLTLOFGVYNLTDRKYLTWESARSIKPFGTSNLINQKTGAGINRF  
DYTLVDAVGYIKPIKNLTLOFGVYNLTDRKYLTWESARSIKPFGTSNLINQKTGAGINRF  
DYTLVDAVGYIKPIKNLTLOFGVYNLTDRKYLTWESARSIKPFGTSNLINQKTGAGINRF  
DYTLVDAVGYIKPIKNLTLOFGVYNLTDRKYLTWESARSIKPFGTSNLINQKTGAGINRF  
\*\*\*\*\*;

1135\_PM0300\_964aa  
WP\_005753642.1  
WP\_016534557.1  
WP\_010906573.1  
AAQ14873.1  
1121\_HgbA\_968aa

YSPGRNFKFSAEITF  
YSPGRNFKFSAEITF  
YSPGRNFKFSAEITF  
YSPGRNFKLSAEITF  
YSPGRNFKFSAEITF  
YSPGRNFKFSAEITF  
\*\*\*\*\*;

FIG. 19-1.

WP_017861186.1 AAU29202.1 EGP04511.1 1135_HasR_848aa WP_005752163.1	MQKSQPYPIHLGIFIMLGLPTWAFAQANLEKSTINKLETILVNESEEKNKFDENIKTYL MQQQPYPIHLGIFIMLGLPTWAFAQANLEKSTINKLETILVNESEEKNKFDENIKTYL MQQQPYPIHLGIFIMLGLPTWAFAQANLEKSTINKLETILVNESEEKNKFDENIKTYL MQQQPYPIHLGIFIMLGLPTWAFAQANLEKSTINKLETILVNESEEKNKFDENIKTYL MQQQPYPIHLGIFIMLGLPTWAFAQANLEKSTINKLETILVNESEEKNKFDENIKTYL *****
WP_017861186.1 AAU29202.1 EGP04511.1 1135_HasR_848aa WP_005752163.1	SSGSYSYLSQS DISTFRGSSVGDFLSGVPGVIVGNKRNSGALSVNIRGIANENRVPVWID SSGSYSYLSQS DISTFRGSSVGDFLSGVPGVIVGNKRNSGALSVNIRGIANENRVPVWID SSGSYSYLSQS DISTFRGSSVGDFLSGVPGVIVGNKRNSGALSVNIRGIANENRVPVWID SSGSYSYLSQS DISTFRGSSVGDFLSGVPGVIVGNKRNSGALSVNIRGIANENRVPVWID SSGSYSYLSQS DISTFRGSSVGDFLSGVPGVIVGNKRNSGALSVNIRGIANENRVPVWID *****
WP_017861186.1 AAU29202.1 EGP04511.1 1135_HasR_848aa WP_005752163.1	RGLQSVP SYQGYAGSSTRYLD PDLI SQVE IERGPSLQMDATGATGGVVR VETLRWQDII RGLQSVP SYQGYAGSSTRYLD PDLI SQVE IERGPSLQMDATGATGGVVR VETLRWQDII *****
WP_017861186.1 AAU29202.1 EGP04511.1 1135_HasR_848aa WP_005752163.1	PQGNKLGVR LKLGTMTNTVSPPEYYTRGGYQT KYISKCLSNHTGLCQTO TYAPNARYSSH PQGNWGVRLKLGTMTNTVSPPEYYTRGGYQT KYISKCLSNHTGLCQTO TYAPNARYSSH PQGNWGVRLKLGTMTNTVSPPEYYTRGGYQT KYISKCLSNHTGLCQTO TYAPNARYSSH PQGNWGVRLKLGTMTNTVSPPEYYTRGGYQT KYISKCLSNHTGLCQTO TYAPNARYSSH PQGNWGVRLKLGTMTNTVSPPEYYTRGGYQT KYISKCLSNHTGLCQTO TYAPNARYSSH *****
WP_017861186.1 AAU29202.1 EGP04511.1 1135_HasR_848aa WP_005752163.1	GFDLNAYN YSLAFANKWSNADLVLAYAKRKQGNYFVGRHGT PVI ESEFEEDSVEVKEP GFDLNAYN YSLAFANKWQNADLVLAYAKRKQGNYFVGRHGT PVI ESEFEEDSVEVKEP GFDLNAYN YSLAFANKWQNADLVLAYAKRKQGNYFVGRHGT PVI ESEFEEDSVEVKEP GFDLNAYN YSLAFANKWQNADLVLAYAKRKQGNYFVGRHGT PVI ESEFEEDSVEVKEP GFDLNAYN YSLAFANKWQNADLVLAYAKRKQGNYFVGRHGT PVI ESEFEEDSVEVKEP *****
WP_017861186.1 AAU29202.1 EGP04511.1 1135_HasR_848aa WP_005752163.1	RVHEDVEIGSLTFKENRSTLYRPGEAEALNTSQDNTSYLAKINVYNDVHRLGLAYRHYSR RVHEDVEIGSLTFKENRSTLYRPGEAEALNTSQDNTSYLAKINVYNDVHRLGLAYRHYSR RVHEDVEIGSLTFKENRSTLYRPGEAEALNTSQDNTSYLAKINVYNDVHRLGLAYRHYSR RVHEEVEIGSLTFKENRSTLYRPGEAEALNTSQDNTSYLAKINVYNDVHRLGLAYRHYSR RVHEEVEIGSLTFKENRSTLYRPGEAEALNTSQDNTSYLAKINVYNDVHRLGLAYRHYSR *****
WP_017861186.1 AAU29202.1 EGP04511.1 1135_HasR_848aa WP_005752163.1	FGEIMSSILNFRAY GALQGEGTEVKVD SYHANYSYNPTTPVNVLSVNAYFTDSDSSNFTP FGEIMSSILNFRAY GALQGEGTEVKVD SYHANYSYNPTTPVNVLSVNAYFTDSDSSNFTP FGEIMSSILNFRAY GALQGEGTEVKVD SYHANYSYNPTTPVNVLSVNAYFTDSDSSNFTP FGEIMSSILNFRAY GALQGEGTEVKVD SYHANYSYNPTTPVNVLSVNAYFTDSDSSNFTP FGEIMSSILNFRAY GALQGEGTEVKVD SYHANYSYNPTTPVNVLSVNAYFTDSDSSNFTP *****
WP_017861186.1 AAU29202.1 EGP04511.1 1135_HasR_848aa WP_005752163.1	FIEEYGYSLSSRHAHFLVSKQKGLSIENTSIFQLNDKPFTLKYGGLAHSYERIYQPRNAQA FIEEYGYSLSSRHAHFLVSKQKGLSIENTSIFQLNDKPFTLKYGGLAHSYERIYQPRNAQA FIEEYGYSLSSRHAHFLVSKQKGLSIENTSIFQLNDKPFTLKYGGLAHSYERIYQPRNAQA FIEEYGYSLSSRHAHFLVSKQKGLSIENTSIFQLNDKPFTLKYGGLAHSYERIYQPRNAQA FIEEYGYSLSSRHAHFLVSKQKGLSIENTSIFQLNDKPFTLKYGGLAHSYERIYQPRNAQA *****

FIG. 19-2.

WP\_017861186.1  
AAU29202.1  
EGP04511.1  
1135\_HasR\_848aa  
WP\_005752163.1

RVRAKGYPEDAIGPLYIRDGKRKEWSAFVAANYPITSWLKADIGLRYLQSTIYDYIVRTE  
RVRAKGYPEDAIGPLYIRDGKRKEWSAFVAANYPITSWLKADIGLRYLQSTIYDYIVRTE  
RVRAKGYPEDAIGPLYIRDGKRKEWSAFVAANYPINSWLKADIGLRYLQSTIYDYIVRTE  
RVRAKGYPEDAIGPLYIRDGKRKEWSAFVAANYPITSWLKADIGLRYLQSTIYDYIVRTE  
RVRAKGYPEDAIGPLYIRDGKRKEWSAFVAANYPITSWLKADIGLRYLQSTIYDYIVRTE  
\*\*\*\*\*

WP\_017861186.1  
AAU29202.1  
EGP04511.1  
1135\_HasR\_848aa  
WP\_005752163.1

RVNIGGALVPNPNGSGNIWVEKYKDVVKQAPVKNKGMSPIVMFTFEPINGVQIYTKEYAE  
RVNIGGALVPNPNGSGNIWVEKYKDVVKQAPVKNKGMSPIVMFTFEPINGVQIYTKEYAE  
RVNIGGALVPNPNGPGNIIWVEKYKDVVKQAPVKNKGMSPIVMLTFEPINGVQIYTKEYAE  
RVNIGGALVPNPNGSGNIWVEKYKDVVKQAPVKNKGMSPIVMLTFEPINGVQIYTKEYAE  
RVNIGGALVPNPNGSGNIWVEKYKDVVKQAPVKNKGMSPIVMLTFEPINGVQIYTKEYAE  
\*\*\*\*\*

WP\_017861186.1  
AAU29202.1  
EGP04511.1  
1135\_HasR\_848aa  
WP\_005752163.1

ALRSPSLFQATKGWSMSATEDNLLEQLRPERAKNWEAGINLFYENLGGKDNILGFKLAYFN  
ALRSPSLFQATKGWSMSATADNLLEQLRPERAKNWEAGINLFYENLGGKDNILGFKLAYFN  
ALRSPSLFQATKGWSMSATADNLLEQLRPERAKNWEAGINLFYENLGGKDNILGFKLAYFN  
ALRSPSLFQATKGWSMSATADNLLEQLRPERAKNWEAGINLFYENLGGKDNILGFKLAYFN  
ALRSPSLFQATKGWSMSATADNLLEQLRPERAKNWEAGINLFYENLGGKDNILGFKLAYFN  
\*\*\*\*\*

WP\_017861186.1  
AAU29202.1  
EGP04511.1  
1135\_HasR\_848aa  
WP\_005752163.1

NRIKDYLTRSYSPKDKVTQTINIQSQAQFKGIELSAYYDMGKFYAKLAGTYYTKTFCCLTA  
NRIKDYLTRSYSPKDKVTQTINIQSQAQFKGIELSAYYDMGKFYAKLAGTYYTKTFCCLTA  
NRIKDYLTRSYSPKDKVTQTINIQSQAQFKGIELSAYYDMGKFYAKLAGTYYTKTFCCLTA  
NRIKDYLTRSYSPKDKVTQTINIQSQAQFKGIELSAYYDMGKFYAKLAGTYYTKTFCCLTA  
NRIKDYLTRSYSPKDKVTQTINIQSQAQFKGIELSAYYDMGKFYAKLAGTYYTKTFCCLTA  
\*\*\*\*\*

WP\_017861186.1  
AAU29202.1  
EGP04511.1  
1135\_HasR\_848aa  
WP\_005752163.1

EQAGKGEQCNSGYVYRSNLNNNAVPPRNLHATLGTRLFEQKLDIGARYSYYSKRLVPVLS  
EQAGKGEQCNSGYVYRSNLNNNAVPPRNLHATLGTRLFEQKLDIGARYSYYSKRLVPVLS  
EQAGKGEQCNSGYVYRSNLNNNAVPPRNLHATLGTRLFEQKLDIGARYSYYSKRLVPVLS  
EQAGKGEQCNSGYVYRSNLNNNAVPPRNLHATLGTRLFEQKLDIGARYSYYSKRLVPVLS  
EQAGKGEQCNSGYVYRSNLNNNAVPPRNLHATLGTRLFEQKLDIGARYSYYSKRLVPVLS  
\*\*\*\*\*

WP\_017861186.1  
AAU29202.1  
EGP04511.1  
1135\_HasR\_848aa  
WP\_005752163.1

AERFVNTSSIEWAPYSLVDLYANYNVSNNLKLTMDNMFNRYYLDINNMGLNTAFGRTL  
AERFVNTSSIEWAPYSLVDLYANYNVSNNLKLTMDNMFNRYYLDINNMGLNTAFGRTL  
AERFVNTSSIEWAPYSLVDLYANYNVSNNLKLTMDNMFNRYYLDINNMGLNTAFGRTL  
AERFVNTSSIEWAPYSLVDLYANYNVSNNLKLTMDNMFNRYYLDINNMGLNTAFGRTL  
AERFVNTSSIEWAPYSLVDLYANYNVSNNLKLTMDNMFNRYYLDINNMGLNTAFGRTL  
\*\*\*\*\*

WP\_017861186.1  
AAU29202.1  
EGP04511.1  
1135\_HasR\_848aa  
WP\_005752163.1

BLGLEYRF  
BLGLEYRF  
BLGLEYRF  
BLGLEYRF  
BLGLEYRF  
\*\*\*\*\*

FIG. 20-1.

WP_005751557.1 1135_PM0741_784aa	MKYPLSYKNIARSIPFLSFIAPPLYAQETTELEQITVQESATAEVNKTSPTVISKSATTI MKYPLSYKNIARSIPFLSFIAPPLYAQETTELEQITVQESATAEVNKTSPTVISKSATTI MKYPLSYKNIARSIPFLSFIAPPLYAQETTELEQITVQESATAEVNKTSPTVISKSATTI MKYPLSYKNIARSIPFLSFIAPPLYAQETTELEQITVQESATAEVNKTSPTVISKSATTI MKYPLSYKNIARSIPFLSFIAPPLYAQETTELEQITVQESATAEVNKTSPTVISKSATTI *****
WP_005751557.1 1135_PM0741_784aa	QNEMIRDTRDLVRYTTEVGISDNGRFLKGFMARGVEDNRVGISIDGVSLPDSEENSLYAR QNEMIRDTRDLVRYTTEVGISDNGRFLKGFMARGVEDNRVGISIDGVSLPDSEENSLYAR QNEMIRDTRDLVRYTTEVGISDNGRFLKGFMARGVEDNRVGISIDGVSLPDSEENSLYAR QNEMIRDTRDLVRYTTEVGISDNGRFLKGFMARGVEDNRVGISIDGVSLPDSEENSLYAR QNEMIRDTRDLVRYTTEVGISDNGRFLKGFMARGVEDNRVGISIDGVSLPDSEENSLYAR ***
WP_005751557.1 1135_PM0741_784aa	YGNFNNRSLSIDPPELIQTIDIVRGSDSFNAGSGSLGGGVNYNTLDPQHIVKAGNSVGALL YGNFNNRSLSIDPPELIQTIDIVRGSDSFNAGSGSLGGGVNYNTLDPQHIVKAGNSVGALL YGNFNNRSLSIDPPELIQTIDIVRGSDSFNAGSGSLGGGVNYNTLDPQHIVKAGNSVGALL YGNFNNRSLSIDPPELIQTIDIVRGSDSFNAGSGSLGGGVNYNTLDPQHIVKAGNSVGALL YGNFNNRSLSIDPPELIQTIDIVRGSDSFNAGSGSLGGGVNYNTLDPQHIVKAGNSVGALL *****
WP_005751557.1 1135_PM0741_784aa	RGSYASKNREWRVTLGIGYVGEKFALLMYSQRTGHEFKSRGSGPEFRYSSSQHPDPVTQ RGSYASKNREWRVTLGIGYVGEKFALLMYSQRTGHEFKSRGSGPEFRYSSSQHPDPVTQ RGSYASKNREWRVTLGIGYVGEKFALLMYSQRTGHEFKSRGSGPEFRYSSSQHPDPVTQ RGSYASKNREWRVTLGIGYVGEKFALLMYSQRTGHEFKSRGSGPEFRYSSSQHPDPVTQ RGSYASKNREWRVTLGIGYVGEKFALLMYSQRTGHEFKSRGSGPEFRYSSSQHPDPVTQ *****
WP_005751557.1 1135_PM0741_784aa	RFHNYLAKMNYQINDNQRIGLTENGQTCGRYIDERSYTLMGSQWREADDQNERLNANLYY RFHNYLAKMNYQINDKQRIGLTENGQTCGRYIDERSYTLMGSQWREADDQNERLNANLYY RFHNYLAKMNYQINDNQRIGLTENGQTCGRYIDERSYTLMGSQWREADDQNERLNANLYY RFHNYLAKMNYQINDKQRIGLTENGQTCGRYIDERSYTLMGSQWREADDQNERLNANLYY RFHNYLAKMNYQINDNQRIGLTENGQTCGRYIDERSYTLMGSQWREADDQNERLNANLYY *****
WP_005751557.1 1135_PM0741_784aa	YYAPSTGWLAYSFKFDLQYKTDLLAAVNYKGGRHFTTDAKELENEIYDRRMRTVFTRGSVEL YYAPSTGWLAYSFKFDLQYKTDLLAAVNYKGGRHFTTDAKELENEIYDRRMRTVFTRGSVEL YYAPSTGWLAYSFKFDLQYKTDLLAAVNYKGGRHFTTDAKELENEIYDRRMRTVFTRGSVEL YYAPSTGWLAYSFKFDLQYKTDLLAAVNYKGGRHFTTDAKELENEIYDRRMRTVFTRGSVEL YYAPSTGWLAYSFKFDLQYKTDLLAAVNYKGGRHFTTDAKELENEIYDRRMRTVFTRGSVEL *****
WP_005751557.1 1135_PM0741_784aa	NAQPVHFYGEHTLTICKYVSQRDFKNINQDRIGIGTNYDTQYHYTIQYPIRTKQYGLSLK NAQPVHFYGEHTLTICKYVSQRDFKNINQDRIGIGTNYDTQYHYTIQYPIRTKQYGLSLK NAQPVHFYGEHTLTICKYVSQRDFKNINQDRIGIGTNYDTQYHYTIQYPIRTKQYGLSLK NAQPVHFYGEHTLTICKYVSQRDFKNINQDRIGIGTNYDTQYHYTIQYPIRTKQYGLSLK NAQPVHFYGEHTLTICKYVSQRDFKNINQDRIGIGTNYDTQYHYTIQYPIRTKQYGLSLK *****
WP_005751557.1 1135_PM0741_784aa	DHVRWNDTFSSHLLGLRYDHTKLKPKELNAPCSKACLEEGKPCKPTRFSTVSTFAGLEAQLS DHVRWNDTFSSHLLGLRYDHTKLKPKELNAPCSKACLEEGKPCKPTRFSTVSTFAGLEAQLS DHVRWNDTFSSHLLGLRYDHTKLKPKELNAPCSNACLEEGKPCKPTRFSTVSTFAGLEAQLS DHVRWNDTFSSHLLGLRYDHTKLKPKELNAPCSKACLEEGKPCKPTRFSTVSTFAGLEAQLS DHVRWNDTFSSHLLGLRYDHTKLKPKELNAPCSKACLEEGKPCKPTRFSTVSTFAGLEAQLS *****
WP_005751557.1 1135_PM0741_784aa	PSWMLGYNISTGYRVPTASEMFSTNAYGTWKSNPSLKPKEKSINHTLSQLKGNSEKGLLD PSWMLGYNISTGYRVPTASEMFSTNAYGTWKSNPSLKPKEKSINHTLSQLKGNSEKGLLD PSWMLGYNISTGYRVPTASEMFSTNAYGTWKSNPSLKPKEKSINHTLSQLKGNSEKGLLD PSWMLGYNISTGYRVPTASEMFSTNAYGTWKSNPSLKPKEKSINHTLSQLKGNSEKGLLD PSWMLGYNISTGYRVPTASEMFSTNAYGTWKSNPSLKPKEKSINHTLSQLKGNSEKGLLD *****

FIG. 20-2.

WP\_005751557.1  
1135\_PM0741\_784aa  
WP\_016534554.1  
WP\_064972816.1  
WP\_074865020.1

LTLYQTNYRHFLFEQESLIQRTIEIRYGPYTYQSQEQQMVNLKAKIYGVELKTHVNLDQ  
LTLYQTNYRHFLFEQESLIQRTIEMRYGRPYTYQSQEQQMVNLKAKIYGVELKTHVNLDQ  
LTLYQTNYRHFLFEQESLIQRTIEMRYGRPYTYQSQEQQMVNLKAKIYGVELKTHVNLDQ  
LTLYQTNYRHFLFEQESLIQRTIEMRYGRPYTYQSQEQQMVNLKAKIYGVELKTHVNLDQ  
LTLYQTNYRHFLFEQESLIQRTIEMRYGRPYTYQSQEQQMVNLKAKIYGVELKTHVNLDQ  
\*\*\*\*\*;\*\*\*\*\*;\*\*\*\*\*;\*\*\*\*\*;\*\*\*\*\*;

WP\_005751557.1  
1135\_PM0741\_784aa  
WP\_016534554.1  
WP\_064972816.1  
WP\_074865020.1

MIAVLPQGFKFYAALGYSKGKLSNNASLLSIQPLKILLGLDYATNGKWAIFNRLITYLGE  
MIAVLPQGFKFYAALGYSKGKLSNNASLLSIQPLKILLGLDYATNGKWAIFNRLITYLGE  
MIAVLPQGFKFYAALGYSKGKLSNNASLLSIQPLKILLGLDYATNGKWAIFNRLITYLGE  
MIAVLPQGFKFYAALGYSKGKLSNNASLLSIQPLKILLGLDYATNGKWAIFNRLITYLGE  
MIAVLPQGFKFYAALGYSKGKLSNNASLLSIQPLKILLGLDYATNGKWAIFNRLITYLGE  
\*\*\*\*\*;\*\*\*\*\*;\*\*\*\*\*;\*\*\*\*\*;

WP\_005751557.1  
1135\_PM0741\_784aa  
WP\_016534554.1  
WP\_064972816.1  
WP\_074865020.1

KRASDAKVEIKRRCTEFVTETDPWSGEQITHCKKRELYPDLSYSTYKHLNKSASFVFDTEGY  
KRASDAKVEIKRRCTEFVTETDPWTGQITRCKKRELYPDLSYSTYKHLNKSASFVFDTEGY  
KRASDAKVEIKRRCTEFVTETDPWSGEQITRCKKRELYPDLSYSTYKHLNKSASFVFDTEGY  
KRASDAKVEIKRRCTEFVTETDPWSGEQITRCKKRELYPDLSYSTYKHLNKSASFVFDTEGY  
KRASDAKVEIKRRCTEFVTETDPWSGEQITRCKKRELYPDLSYSTYKHLNKSASFVFDTEGY  
\*\*\*\*\*;\*\*\*\*\*;\*\*\*\*\*;\*\*\*\*\*;

WP\_005751557.1  
1135\_PM0741\_784aa  
WP\_016534554.1  
WP\_064972816.1  
WP\_074865020.1

YKITDDITFRAGIYNLFNKKYHTWDALRGINANSTLNSVDREGKGLQRFYAPGRNYAASL  
YKITDDITFRAGIYNLFNKKYHTWDALRGINANSTLNSVDREGKGLQRFYAPGRNYAASL  
YKITDDITFRAGIYNLFNKKYHTWDALRGINANSTLNSVDREGKGLQRFYAPGRNYAASL  
YKITDDITFRAGIYNLFNKKYHTWDALRGINANSTLNSVDREGKGLQRFYAPGRNYAASL  
YKITDDITFRAGIYNLFNKKYHTWDALRGINANSTLNSVDREGKGLQRFYAPGRNYAASL  
\*\*\*\*\*;\*\*\*\*\*;\*\*\*\*\*;\*\*\*\*\*;

WP\_005751557.1  
1135\_PM0741\_784aa  
WP\_016534554.1  
WP\_064972816.1  
WP\_074865020.1

EIRF  
EIRF  
EIRF  
EIRF  
EIRF  
\*\*\*\*\*

FIG. 21-1.

WP\_050948957.1  
WP\_014391043.1  
WP\_016533738.1  
1135\_P1062\_0207600\_742aa  
WP\_025248456.1

WP\_050948957.1  
WP\_014391043.1  
WP\_016533738.1  
1135\_P1062\_0207600\_742aa  
WP\_025248456.1

NIEQQQADNAANLNLN1LPGVNMAGGFRPGQTLN1NGMGAEDVRVQLDGATKSFEKYQQ  
NIIQQQADNAADLNLN1LPGVNMAGGFRPGQTLN1NGMGAELRVVQLDGATKSFEKYQQ  
NIDQQQADNAADLNLN1LPGVNMAGGFRPGQTLN1NGMGAEDVRVQLDGATKSFEKYQQ  
NIDQQQADNAADLNLN1LPGVNMAGGFRPGQTLN1NGMGAEDVRVQLDGATKSFEKYQQ  
NIDQQQADNAADLNLN1LPGVNMAGGFRPGQTLN1NGMGAEDVRVQLDGATKSFEKYQQ  
\*\*\*\*\*

WP\_050948957.1  
WP\_014391043.1  
WP\_016533738.1  
1135\_P1062\_0207600\_742aa  
WP\_025248456.1

WP\_050948957.1  
WP\_014391043.1  
WP\_016533738.1  
1135\_P1062\_0207600\_742aa  
WP\_025248456.1

NNNQKTYSTALVLQNEQKNIDLLLLFGSVPNAGDYKRPDKSKILFGSKNNQKTGLIKVNWQI  
MNNQKTYSTALVLQNEQKNIDLLLLFGSVPNRNASNYTRPDKSKILFGSKNNQKSGSLIKVNWQI  
NNNQKTYSTALVLQNEQKNIDLLLLFGSVPNRNASNYTRPDKSKILFGSKNNQKSGSLIKVNWQI  
NNNQKTYSTALVLQNEQKNIDLLLLFGSVPNRNASNYTRPDKSKILFGSKNNQKSGSLIKVNWQI  
NNNQKTYSTALVLQNEQKNIDLLLLFGSVPNRNASNYTRPDKSKILFGSKNNQKSGSLIKVNWQI

WP\_050948957.1  
WP\_014391043.1  
WP\_016533736.1  
1135\_P1062\_0207600\_742aa  
WP\_025248456.1

TPEHLLTLLSSVYGIHKGWPWAAKRDUVSRPTETEIKRYGIDVAWKRKLVYRDQKDESYS  
TPEHLLTLLSSVYGIHKGWPWAAKRDUVSRPTETEIKRYGIDVAWKRKLVYRDQKDESYS  
TPEHLLTLLSSVYGIHKGWPWAAKRDUVSRPTETEIKRYGIDVAWKRKLVYRDQKDESYS  
TPEHLLTLLSSVYGIHKGWPWAAKRDUVSRPTETEIKRYGIDVAWKRKLVYRDQKDESYS  
TPEHLLTLLSSVYGIHKGWPWAAKRDUVSRPTETEIKHYGIDVAWKRKLVYRDQKDESYS  
TPEHLLTLLSSVYGIHKGWPWAAKRDUVSRPTETEIKHYGIDVAWKRKLVYRDQKDESYS

WP\_050948957.1  
WP\_014391043.1  
WP\_016533738.1  
1135\_P1062\_0207600\_742aa  
WP\_025248456.1

LKYRYLPENNKKWINLSVQLSYSKTEQNDTRHEKVTSFLGTLGNKSWITYSDLTFDISNT  
LKYRYLPENNKKWINLSVQLSYSKTEQNDTRHEKVTSFLGTLGNKSWITYSDLTFDISNT  
LKYRYLPENNKKWINLSVQLSYSKTEQNDTRHEKVTSFLGTLGNKSWITYSDLTFDISNT  
LKYRYLPENNKKWINLSVQLSYSKTEQNDTRHEKVTSFLGTLGNKSWITYSDLTFDISNT  
LKYRYLPENNKKWINLSVQLSYSKTEQNDTRHEKVTSFLGTLGNKSWITYSDLTFDISNT

WP\_050948957.1  
WP\_014391043.1  
WP\_016533738.1  
1135\_P1062\_0207600\_742aa  
WP\_025242456.1

STLNIGRAEHELLFGQLWLNTRNLMYHKGKMNDKTYNYGYFQPYYMPSGRQYTQAFYL  
STLNIGRAEHELLFGQLWLNKRNTLMYHKEGVKADYNNGYFQPYYMPSGRQYTQAFYL  
STLNIGRAEHELLFGQLWLNKRNTLMYHKEGVKADYNNGYFQPYYMPSGRQYTQAFYL  
STLNIGRAEHELLFGQLWLNKRNTLMYHKGGVKADYNNGYFQPYYMPSGRQYTQAFYL  
STLNIGRAEHELLFGQLWLNKPNTLMYHKGGVKADYNNGYFQPYYMPSGRQYTQAFYL

WP\_050948957.1  
WP\_014391043.1  
WP\_016533736.1  
1135\_P1062\_0207600\_742aa  
WP\_0265242456.1

QDQIKWKWNIIIPSTGARYDHINNIGQKLNAPQYNDISAGHNYNSQKNYNGWSYYLGLKYDVN  
QDQIKWKWNFLFTGGIRYDHINNIGQKLNAPRYNDISAGHDYSQKNYNGWSYYLGLKYDVN  
QDQIKWKWNFLFTGGIRYDHINNIGQKLNAPRYNDISAGHDYSQKNYNGWSYYLGLKYDVN  
QDQIKWKWNFLFTGGIRYDHINNIGQKLNAPRYNDISAGHDYSQKNYNGWSYYLGLKYDVN  
QDQIKWKWNFLFTGGIRYDHINNIGQKLNAPRYNDISAGHDYSQKNYNGWSYYLGLKYDVN

WP\_050948957.1  
WP\_014391043.1  
WP\_016533738.1  
1135\_P1062\_0207600\_742aa  
WP\_050948957.1

HYLSLFTNFSRTWRAPVIDEQYETQYSKASVSPATSLNLEKEMISQTRGGGIVTLLNLHFQE  
HYLSLFTNFSKTKWRAPVIDEQYETQYSQASVSATSLNLEKEMINQTRVGGIITLNHLFQE  
HYLSLFTNFSKTKWRAPVIDEQYETQYSQASVSATSLNLEKEMINQTRVGGIITLNHLFQE  
HYLSLFTNFSKTKWRAPVIDEQYETQYSQASVSATSLNLEKEMINQTRVGGIITLNHLFQE  
HYLSLFTNFSKTKWRAPVIDEQYETQYSQASVSATSLNLEKEMINQTRVGGIITLNHLFQE

FIG. 21-2.

WP\_050948957.i  
WP\_014391043.i  
WP\_016533738.i  
1135\_P1062\_0207600\_742aa  
WP\_025248456.i

DDTFQFRATYFYHRGKNEIFKTRGVNCVGNALDVNKICPKIISNYRNLPGYVIQGAELE  
NDAFQFPRTTYFYNRGKNEIFKTRGVNCVENEALDVNSVCVKIISNYRNLPGYVIQGAELE  
NDAFQFPRTTYFYNRGKNEIFKTRGVNCVENEALDVNSVCVKIISNYRNLPGYVIQGAELE  
NDAFQFRRTTYFYNRGKNEIFKTRGVNCVENEALDVNSVCVKIISNYRNLPGYVIQGAELE  
NDAFQFRRTTYFYNRGKNEIFKTRGVNCVENEALDVNSVCVKIISNYRNLPGYVIQGAELE  
\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:  
\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:

WP\_050948957.1  
WP\_014391043.1  
WP\_016533738.1  
1135\_P1062\_0207600\_742aa  
WP\_025248456.1

AYYQSTYLFGELETYSVKGKRDTSPRNPWGKTSTWIAEIPFRKATATLGFnPKYYLTVG  
AYYQSSSYLFGGLTYSVKGKRDTSPRNPWSKTSTWIAETPPRKATATLGFnIPEYYFTAG  
AYYQSSSYLFGGLTYSVKGKRDTSPRNPWSKTSTWIAETPPRKATATLGFnIPEYYFTAG  
AYYQSSSYLFGGLTYSVKGKRDTSPRNPWSKTSTWIAETSPPRKATATLGFnIPEYYFTAG  
AYYQSSSYLFGGLTYSVKGKRDTSPRNPWSKTSTWIAETPPRKATATLGFnIPEYYFTAG  
\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*:\*\*\*\*\*

WP\_050948957.1  
WP\_014391043.1  
WP\_016533738.1  
1135\_P1062\_0207600\_742aa  
WP\_025248456.1

WRAEFVRKQDRSPSSRPKASYYLSQLFASPFGYSLHNLFLSWSTPEKIKGMNIIKTVNDLFN  
WRAEFVRKQDRSPLSNDSKASYW-ALPSSKGYSLSHVSFFWSNSTKIKGMNFKVTVNDLFN  
WRAEFVRKQDRSPLSNDSKASYW-ALPSSKGYSLSHVSFFWSNSTKIKGMNFKVTVNDLFN  
WRAEFVRKQDRSPLSNDSKASYW-ALPSSKGYSLSHVSFFWSNSTKIKGMNFKVTVNDLFN  
WRAEFVRKQDRSPLSNDSKASYW-ALPSSKGYSLSHVSFFWSNSTKIKGMNFKVTVNDLFN  
\*\*\*\*\*

WP\_050948957.1  
WP\_014391043.1  
WP\_016533738.1  
1135\_P1062\_0207600\_742aa  
WP\_025249456.1

RAYNPYLGELASGTGPNIKFSLSQKF  
RPYYPYLGELASGTGRNVKFSLTQQF  
RPYYPYLGELASGTGRNVKFSLTQQF  
RPYYPYLGELASGTGRNVKFSLTQQF  
RPYYPYLGELASGTGRNVKFSLTQQF

FIG. 22-1.

SEQ ID NO:43

ATGATTCAAGAGGTTGAAAGTAAATAAATTGGCTGTTGATGATGTGTTGCATTCCGCA  
AGTAGTTGGCAAATACAGAAAAGAAGCAAATTGTTTTAGATGAAATTAGTGTGGAGTCGA  
AAGGTGCTGCTTCTCGTAGCGATCCCTTCCGGCTTACAAAACAAAATGACATTAGTCAGT  
AAGCAAAATTAACAGGATCCAGTACATTAGGTAATGCCTAGCAGGTGAACTCAGTGTACA  
TAGTAATCAATTGGTGGTGGATCCAGTGCACCTGTCGTTGAGGACAGGAAGGCGTGCCTTAA  
AAATCTACAAATGGTCTGATGTCATTGATATGTCTCAACTTCTCCTGACCATGCGATTGGT  
GTCGATACGTTATTGGCAGAGCAAGTCGAAATTGTACGTGGTGCCTAACATTGTTATGCCAA  
TGCCTCACCTGCTGGTGTGATTAATGTAGTAGATAAACGCATTCCAACACAGCTCCTCAAAAGG  
GCTATGAAGTGGATTCAACACGCGTTATAACACGAATAGTCATGAAAATTGGTAACGGCTGCA  
CTGACTTCGGGTTAGGTAAGCACATTGCTTACGGTAGAAGAACTGTTACGTGGCTCAAATAA  
TTACCATGTGCCAGCATTAGCTAGACAAAACATTAAATTATGCCCCGATACTCAAAATAAAA  
CGAAGTCAGGTAACTATGGTGTGGCTTTATCGGTGAGCGAGGTTATGTCGGTTTGCGCTACAAAT  
CTTCGTCGTGAGAAATATGGGTTACCGGGACATAACCATAAGCTAGATAGTTGTGCTGCGCATAT  
TTGGGGTGGCAATGTACGTAATGACTATTATTAGGGCTTATCCTCATTAATGCATGATAACGG  
ATCTAGTAAATACGCATTCCATTGTGGATCAAATCATGACATGGATGGAAAACACAGTCATGAT  
CATCCGTACGGTCACGATCATGACCATTCTATTGCCGGCATTGATTGATTCTTATGCTAAACG  
TTATGACATACGTGCTGAAGTCAAACAGCCGATGAAAGCGATTGAGAAAATCAAACAGTTATT  
CGGAAACCCGTTATAAACACGATGAAAAGATGGCAATATGCCGTGAATTATTAAAAATAAC  
GGATATAACCTGCGCGTAGAAATTCCATACGCCATAGCGGGTTGAGTGGCGTTATAGGGC  
GCAGTATCAAACACAAACCAGCAGTGCAGTATTCCGCGTATTGCACCATGCTCAAATAATGCAA  
GCGATCCTGTACATAAGAAAAGCAACCGATCCGTCAAATCACTAAGGGTGATCGTAAGTC  
TGGGCATTGATTGAAAATACTCAATCACAAATGAGTTTTGCTATAGAACAAATTGGCGTTGGCA  
AGATTTTATTGAAATTGGTGTACGTACGGAAAACACGCAATTGACTATTGAATATGATCGCG  
CTTGGCTTTAAAGTAAAGCGAAAGCTAGAAGGCTGTGATCCGAATTGCTCTTTATAGCCC  
TCAGGATGTCGTCAAGGCAGTTATCCAGCACCTGATTGCTTATCATGATCGCGACCTC  
TTATTCTGGTGTACGTTGAAATATGACGCCACTACACCCTTCTTGACTTACTCACATA  
ACGAACGTCACCCAAACACCGATGGAATTGTATTACACGGCAAACATTAGCGACGGTGTAC  
GAACATGGCAATCGTAATCTGAAAAAGAAGTTCTGATAACTGGGAAGTGGTCTTGTATCT  
TGGTGACAAGCTAAGTTAAAGTGAATGTATATTACAATGATTAAAATCGGATTTAATC

FIG. 22-2.

AAACATTGAACAAATCCGTAATTTATCTTGAATCGTTATAATCAATCCAAAGCGAAATATTAT  
GGTGTGGAAGGGCGTATTGACTATGCCTGACACCTGAACACTGCACATGGGACTTTGGTGATTA  
TGTGCGTGGAAAATGTATGATTACCGCCAACGTACCGTGTGATCATGTGGCGAATAGCTTAG  
AACCCGTTCTCAACCTGATCAAGACGCACCACGTGTCCGCCAATGCCTTACGGCTTCGTGTG  
AATATGGAGATGACTGAGAGTTAACAGTTCACTCGAATACACTTACGTTATCAACAAAAGAA  
AGTAGGCCGTTAGAAAATCAAACGGCTGCATATAGTTATTAAATATCGGAGTGGATTATTCAC  
GCCAAATAGCTGGGTAAATTATCAATTATTGTTCAAGCAAATAATGTGTTAACCGTAAAGTT  
TATTCTCATACTTCTTTTACCCCTTGTACCGCAGATGGGCGTAATGTGACTTACGGATTAAA  
CATCCATTCTAA

SEQ ID NO: 44

MISRGCKVNKFFAVLMMCCI PQVVWANTEKKQIVFLDEISVESKGAAFRSDPLSLPKQNDILVS  
KQKLKTGSSTLGNALAGELSVHSNQFGGSSAPVVRGQEGVRLKILQNGSDVIDMSQLSPDHAIG  
VDTLLAEQVEIVRGASTLLYANASPAGVINVVDKRIPTQLPQKGYEVDFNTRYNTNSHEKLVTA  
LTFLGLKHIALRVEELLRGSNNYHVPFKLDKTLNYVPDTQNKTSGNYGVAFIGERGYVGFAYN  
LRREKYGLPGHNHKLDSCAAHIWGGNVRNDYLYLGLYPHLMHDTDLVNTHFHCGSNHDMDGKHSHD  
HPYGHHDHSIAGPLIDSYAKRYDIRAEVKQPMKAIEKIKLSYSETRYKDEKDGNIAVNLFKNN  
GYNLRVEIFHTPIAGLSGVIGAQYQTQTSSANIPRIAPCSNNASDPCHKKQRDPSKITKGDRKS  
WALIENTQSQMSFFAIEQLRWQDFLFEIGVRTEKQRIDIEYDRAWLFVKRKLEGCDPNSFFYSP  
SGCRQGSYPAPDFASYHDRATSYSGAISWNMTPDYTLSLTYSHNERHPTPMELYHGKHLATVSF  
EHGNRNLKKEVSDNWEVGLAYLGDKLSYKVNVYYNDFKNRIFNQTLNKSGNLSLNRYNQSKAKYY  
GVEGRIDYALTPELHMGLFGDYVRGKLYDLPPTYRVDHVANSLEPVPQPDQDAPRVPPMRLGFRV  
NMEMTESLTSSLEYTYVYQQKKVAPLENQTAAYSLLNIGVDYSRQIAGVNYQLFVQANNVLRKV  
YSHTSFLPFVPQMGRNVTLGLNIHF

FIG. 23.

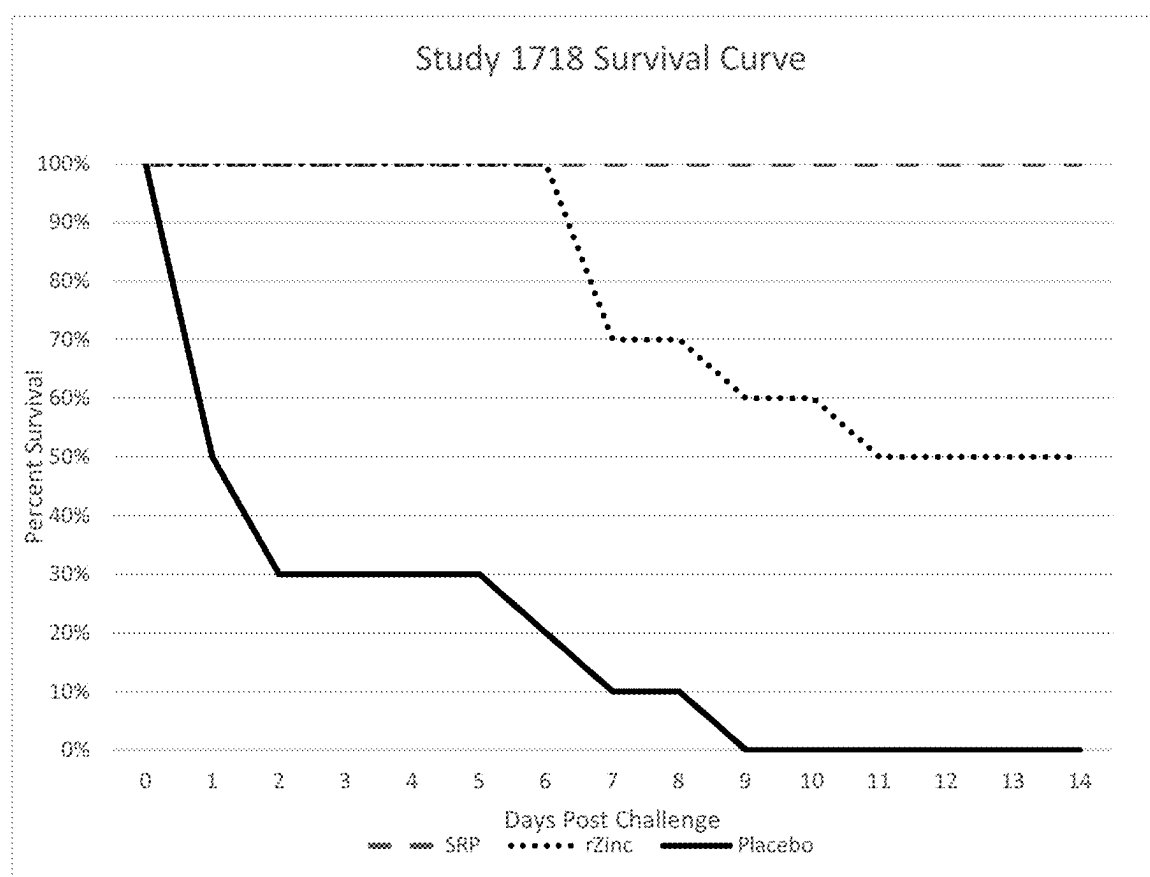


FIG. 24

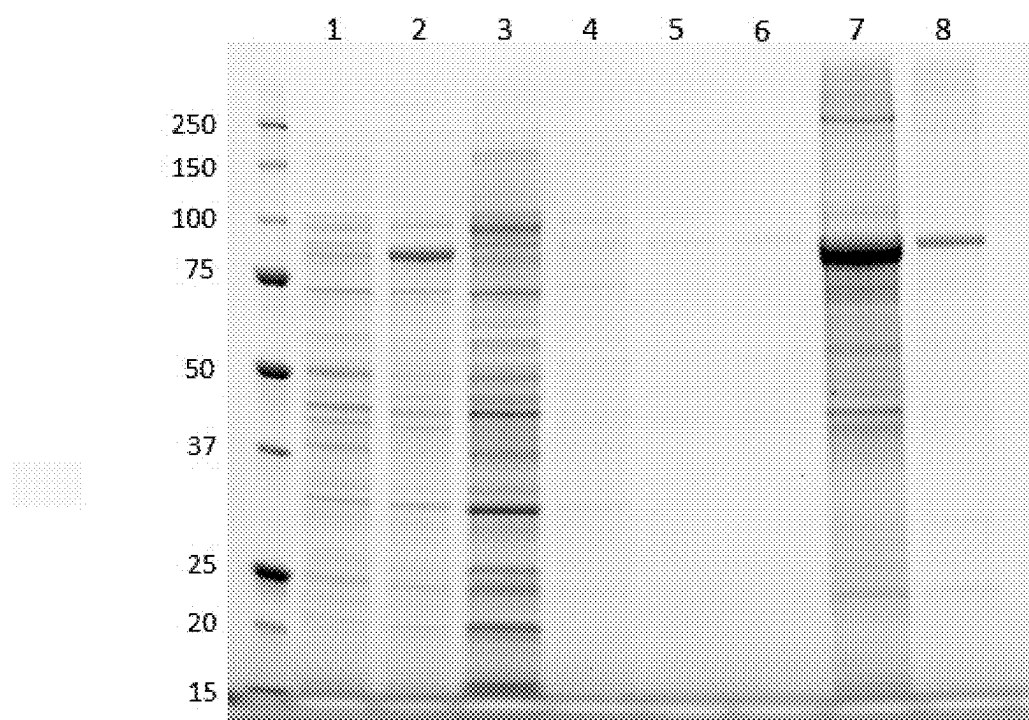


FIG. 25.

ZAP	1	-----SKR8DNYGLPGRHMKFDFTGHIYGNK-RD---KYAYTYLYPHLIG	42
AHG81836.1	239	LSPVGEQGYIGASYSKK8DNYGLPGRHMKFDFTGHIYGNK-RD---KYAYTYLYPHLIG	234
WP_005612269.1	231	LSPVGEQGYIGASYSKK8DNYGLPGRHMKFDFTGHIYGNK-QG---KYAYTYLYPHLIG	286
WP_027074597.1	242	ISPIGERGYIGAAYSHRKDTYGLPGRHMKFDFTGHIYGV-D-RD---KHAHTYLYPHLLT	297
AHG73391.1	242	LSPVGERGYIGAAAYMERTOTYGLPGRHMKFDECIGRHYNEV-RD---KYAYTYLYPHLLD	297
WP_021114857.1	252	LSPIGEKGY1GASYNQRKDRYGLPGRHMKFDTCIAHIIYDMRLQG---KHSYTNLYPHLMS	308
WP_026212957.1	232	LSYIIGNQGHIGLAYSERDOKYVOLVSHMKFDNCERGAFTNTSRGLWOPERRYLIPYPHMS	291
WP_028858792.1	274	LSYVGERGHIGVAYSEBEOKYGLVSHMKFLDGCGYGRVYYPQKINY--KKNKPYLAAYPHUNG	331
WP_016534590.1	246	VAFIGERGYVGFAVILBREKYGGLPGRHMKLDSCAAHINGGNVRN---OY-YLELYPHUTH	301
KDN24548.1	246	VSYIGOKGHIQVSYSRRQDKYQIPGRHMKAYDNCIAHVLTEAS---ISRYYLKAYPHLIQ	302
WP_027021676.1	249	VSWVGQNGHLGVSYSHRDYGLPGRHMKLDNCSGHVFDVTIAS-AVKRMYLLPYPHLIG	307
ZAP	43	EENIGSNPMPFHCGTNAEDGTHSHDNPPFGMHDHMTHKGPVWOLE-----	87
AHG81836.1	295	EENIGSNPMPFHCGTNAEDGTHSHDNPPFGMHDHMTHPGPVWOLESKRFOVKAELRQPFKG	354
WP_005612269.1	287	EENIGSNPMPFHCGTNAEDGTHSHDNPPFGMHDHMTHPGPVWOLESKRFOVKAELRQPFKG	346
WP_027074597.1	298	NELISENPMPFHCGSDPGLDHMHSHDNPPFGMHDHMTHKGPVWOLRSKRDLKMELKQPFSG	357
AHG73391.1	298	EDLISHGPMPFHCGTNAEDGTHSHDNPPFGMHDHMTHKGPVWOLSKSKRDLVKGELLKPFRG	357
WP_021114857.1	309	DEMVTENPHFHCGTNAEDGTHSHDNPPFGMHDHMTHKGPVWOLHSKRDLIKGEIKQPLPH	368
WP_026212957.1	292	DEDNITSLMPFHCGTNAEDGTHSHDNPPFGMHDHMTHKGPVWOLHSKRDLIKGEIKQPLPH	351
WP_028858792.1	332	DEDLAESFHMPFHCGTNAEDGTHSHDNPPFGMHDHMTHKGPVWOLHSKRDLIKGEIKQPLPH	391
WP_016534590.1	302	DTDL-VNTHMPFHCGTNAEDGTHSHDNPPFGMHDHMTHKGPVWOLHSKRDLIKGEIKQPLPH	360
KDN24548.1	303	NWDSSSANFHCGTNAEDGTHSHDNPPFGMHDHMTHKGPVWOLHSKRDLIKGEIKQPLPH	362
WP_027021676.1	308	DEDVNLSQHFHCGTNAEDGTHSHDNPPFGMHDHMTHKGPVWOLHSKRDLIKGEIKQPLPH	367

**PROTEINS AND IMMUNIZING  
COMPOSITIONS CONTAINING  
PASTEURELLA PROTEINS AND METHODS  
OF USE**

**CROSS-REFERENCE TO RELATED  
APPLICATIONS**

[0001] This application is a continuation of U.S. application Ser. No. 18/203,917, filed May 31, 2023, which is a continuation of U.S. application Ser. No. 17/191,913, filed Mar. 4, 2021, now U.S. Pat. No. 11,696,945, which is a continuation of U.S. application Ser. No. 16/484,332, filed Aug. 7, 2019, now U.S. Pat. No. 10,967,056, and which is a § 371 U.S. National Stage of International Application No. PCT/US2018/017682, filed 9 Feb. 2018, which claims the benefit of U.S. Provisional Application Ser. No. 62/457,599, filed Feb. 10, 2017, the disclosures of which are incorporated by reference herein in their entireties.

**SEQUENCE LISTING**

[0002] This application contains a Sequence Listing electronically submitted via Patent Center to the United States Patent and Trademark Office in XML format entitled “0293000056US03”, having a size of 106,580 bytes and created Sep. 18, 2023. This information contained in the Sequence Listing is incorporated by reference herein.

**BACKGROUND**

[0003] *Pasteurella* spp. are gram negative, facultatively anaerobic and pleomorphically rod shaped bacteria able to cause disease in a variety of animals as well as humans. Species of *Pasteurella* in domestic and wild animals have evolved a particular niche for mucosal membranes of the upper respiratory, mucosa of the oral cavity and lower genital tract. In farm animals, diseases are based on clinical symptoms and on the particular *Pasteurella* species associated with disease. Today *Pasteurella multocida* and *Mannheimia haemolytica* are the two most widely recognized species of the Pasteurellaceae family associated with clinical disease in animal husbandry with particular reference to poultry, cattle, pigs and rabbits. The family Pasteurellaceae currently comprises five genera; *Pasteurella*, *Mannheimia*, *Actinobacillus*, *Haemophilus*, and *Lonepinella* and have been the subject of extensive reclassification (Angen et al. Int. J. Syst. Bacteriol. 49:67-86 (1999). Recently *Pasteurella haemolytica* has been taxonomically reclassified as *Mannheimia haemolytica* based on ribosomal sequencing and DNA-DNA hybridization (Oystein et al. Int. J. Syst. Bacteriol. 49:6-86 (1999).

[0004] *Pasteurella multocida* is the etiological agent of avian pasteurellosis, commonly referred to as fowl cholera, a widely distributed and economically important disease of poultry, having a high incidence in chickens, turkeys, geese and ducks (Rhoades et al. Fowl Cholera. In: Adlam, C. F. and Rutter, J. M. (Eds.), *Pasteurella* and Pasteurellosis. Academic Press, London, pp. 95-113. 1989). The organism is also responsible for inducing clinical disease in wild birds, commercially raised game birds and Psittacines (Rhoades and Rimler, 1989, Fowl cholera. In *Pasteurella* and Pasteurellosis, pp. 95-113. Edited by C. F. Adlam & J. M. Rutter. London: Academic Press). All species of birds are susceptible to varying degrees. Heavy breeds appear more susceptible than light breeds, and adult birds and those in late

growing phase appear more susceptible than younger birds (Jordan et al. Poultry Diseases. Ballierre Tindall, London, pp 42-50. 1990.). Four capsular serogroups (A, B, D and F) are recognized among avian strains of *Pasteurella multocida* (Rhoades et al. Avian Dis. 31:895-898 (1987). In domestic poultry, strains belonging to capsular type A are recognized as the primary agent of pasteurellosis (Rhoades and Rimler, 1987, Avian Dis 31, 895-898; Rhoades et al. Fowl Cholera. In: Adlam, C. F. and Rutter, J. M. (Eds.), *Pasteurella* and Pasteurellosis. Academic Press, London, pp. 95-113. 1989; Wilson et al. J. Clin. Microbiol. 31:255-259 1993)). It has been shown that avian strains of *Pasteurella multocida* are genetically diverse as demonstrated by multilocus enzyme electrophoresis and DNA-DNA hybridization. Based on these findings *P. multocida* has now been subdivided into three subspecies, *multocida*, *septica* and *gallicida*.

[0005] The pathogenesis of *P. multocida* is poorly understood but is dependent on both host and pathogen specific factors. To date no single virulence factor has been associated with disease among strains. Virulence factors that have been studied to date include; capsule, endotoxin, outer membrane proteins, serum resistance, iron-acquisition systems, heat shock proteins, neuraminidase production, adhesion factors, antibody cleaving enzymes and the potential existence of cytopathic toxins.

[0006] Clinical manifestation of disease may range from peracute/acute to chronic infections, and may initially arise by the colonization of the upper respiratory tract, followed by invasion and septicemia. During acute infections, few clinical signs are observed before death which is generally dominated by septicemic lesions. In chronic forms of *P. multocida* infections, suppurative lesions may be widely distributed involving the mucous membranes of the respiratory tract, the pharynx, nasal passages, conjunctiva and adjacent tissues of the head.

[0007] Because of the ubiquitous nature of *P. multocida*, and the potential for great economic loss, commercial vaccines have been developed to protect against outbreaks of Fowl cholera in layer and breeder broiler operations throughout the US and abroad. Inactivated commercial Fowl cholera vaccines currently available rely on somatic serotypes to offer protection. Most inactivated vaccines contain, at a minimum, *Pasteurella multocida* strains of serotype 1 (also referred to as A: 1), serotype 3 (also referred to as A: 3), and serotype 4 (also referred to as A: 4). Fowl cholera vaccines made with one serotype do not cross protect against other serotypes, e.g., a vaccine made using a serotype 1 strain does not protect against serotypes 3 or 4.

[0008] The ability of *Pasteurella multocida* to evade the natural defense mechanisms of the vertebrate host depends in part on its ability to obtain host iron, which in turn, directly influences the host-pathogen interaction. Because of iron's essential nature, vertebrate hosts have developed elaborate mechanisms to bind iron in body fluids (e.g., transferrin in blood and lymph fluids and lactoferrin in external secretions). These high affinity iron binding proteins create an iron restricted environment within the host, reducing the level of iron to approximately  $10^{-18}$  molar, a concentration too low to support the growth of nearly all bacteria. These iron sequestering mechanisms of the host act as a natural defense mechanism to combat bacterial invasion. To circumvent these iron-restrictive conditions many bacterial species have evolved mechanisms for obtaining iron. The most common mechanisms include the diffusion of

soluble iron through porins and specialized transport systems that mediate the uptake of iron by siderophores. This latter system is one of the most common and well-studied mechanisms for iron acquisition and involves the specific chelation of ferric iron by siderophores and the synthesis of their cognate transport systems, which permits the bacteria to continue to replicate and overcome the non-specific defense mechanisms of the host. Continued replication, and thus each step in the infectious process, is ultimately dependent on the ability of the organism to obtain iron from its host.

[0009] Divalent metal ions such as iron, cobalt, copper, magnesium, manganese, molybdenum, nickel, selenium, and zinc are trace elements often required for the survival of bacteria infecting both animal and human hosts. These trace metal elements are used by bacteria as cofactors for enzymes that catalyze biochemical reactions for various metabolic pathways required by the organism. The impact of iron on the pathogenesis of bacteria has been studied extensively. Iron is essential for nearly all life and is required for enzymatic and metabolic pathways of organisms at all phylogenetic levels. It has been well-documented that during bacterial sepsis there is an alteration in the concentration of a number of metal ions in serum such as, iron, copper, and zinc. For instance, serum levels of zinc decrease from 10 percent to 60 percent with the onset of infection. Following the onset of infection, zinc is then redistributed from plasma to liver where it is bound to metallothionein. Decreases in serum iron of up to 50 percent have been described during infectious illness, whereas serum copper has been shown to increase in response to inflammatory stimuli. The alteration of these trace metal ions in serum may directly affect the severity or progression of any bacterial infection.

[0010] With so many basic functions relying on the availability of iron, bacteria have evolved a complex regulatory network for acquiring iron under varying physiological conditions. Under anaerobic conditions, iron is present in the soluble ferrous form (Fe II) and can freely diffuse through outer membrane porins into the periplasm. For instance, in *E. coli* the FeoAB transport system present in the cytoplasmic membrane will transport the ferrous iron molecules into the cell cytoplasm. Under aerobic conditions and neutral pH, iron is primarily present in the insoluble ferric form (Fe III) and cannot pass through the outer membrane porins by passive diffusion. Instead, molecules called siderophores are secreted by bacteria, which have a high affinity for ferric iron. The ferric-siderophore complexes are recognized by receptors in the outer membrane collectively referred to as the TonB-dependent receptors. These receptors, once bound to loaded siderophores, are believed to interact with TonB and its associated proteins localized in the periplasm and cytoplasmic membrane. These protein-protein interactions, though poorly understood, serve to provide the energy necessary to transport the ferric-siderophore complexes across the outer membrane and through the periplasmic space. ABC transport systems present in the cytoplasmic membrane serve to transport the iron-siderophore complexes across the cytoplasmic membrane. Reductase enzymes reduce the ferric iron to its ferrous form, which dissociates it from the siderophore and releases iron into the cell.

[0011] Several species of pathogenic bacteria use additional mechanisms to obtain iron from mammalian hosts, including the direct binding of transferrin, heme, and other

heme-containing compounds. The receptor proteins that bind these iron-containing molecules most likely rely on the TonB complex for the energy required to transport heme across the outer membrane, similar to the iron-siderophore complexes. Specialized ABC transporters are then used to transport the heme across the cytoplasmic membrane. In addition, some bacteria secrete hemophores, small molecules that can bind heme and present it to receptors on the bacterial cell surface. Several pathogenic species also produce hemolysins, which are toxins that lyse red blood cells, releasing heme and hemoglobin for uptake by the bacteria.

[0012] The outer membrane proteins of gram-negative bacteria control the selective permeability of many essential nutrients critical to the survival of bacteria, including all pathogenic bacteria that cause disease in animals and man.

This selective permeability of nutrients is controlled by a class of membrane proteins called porins.

It now appears that the majority of the outer membrane proteins on the surface of gram-negative bacteria are porins, identified as

the general porins (e.g., OmpF), monomeric porins (e.g., OmpA), the specific porins (e.g., the maltose-specific porin LamB) and the TonB-dependent, gated porins (e.g., the siderophore receptor FepA).

The porin class of proteins generally share structural features, including the presence of beta-barrels that span the outer membrane.

[0013] Beyond the role of iron as an essential nutrient for microbial survival, there are now many other well-defined transitional metals that play critical roles in bacterial survival, homeostasis, and pathogenesis such as iron, manganese, copper, zinc, magnesium, cobalt, and nickel (Waldron and Robinson and 2009; Porcheron; 2013). Iron, zinc and copper are the three most abundant divalent metal ions in mammals in descending order of concentration. The ability of a bacterium to utilize these transitional metals by finely regulated uptake or acquisition systems significantly contributes to the virulence of pathogenic bacteria. It is well known that bacteria within the same genus/species do not have the same uptake systems for the acquisition of transitional metals owing to the difference in pathogenicity from one strain of bacteria to another. These differences in the ability of bacteria to use different transitional metals based on expressed uptake systems may specifically direct what organ or tissue an organism can invade.

[0014] Little is known regarding the iron-acquisition by *Pasteurella multocida*; it has not been studied nearly to the extent of *E. coli* iron transport systems. The iron regulated proteins of *Pasteurella* have been investigated as potential immunogens as target antigens for different vaccine strategies in multiple animal species. It has been shown that these proteins provide protective efficacy against homologous challenge yet a combination of these proteins to provide heterologous protection has eluded researchers.

[0015] Glisson et al. (Avian Diseases 37:1074-1079, 1993) attempted to induce cross-protection across serotypes by preparing bacterins from *P. multocida* strains X-73 (serotype 1) and P-1059 (serotype 3) under iron-restricted culture conditions. They found that bacterins produced in medium low in iron did not consistently induce significant protection against heterologous challenge.

## SUMMARY OF THE APPLICATION

[0016] We observed results similar to Glisson when growing two heterologous strains, one of serotype 3×4 and one of serotype 1, in iron-restricted medium and attempting a

vaccination and challenge model with one strain as vaccine strain and the other as challenge strain (see Examples 6-9), i.e., some protection was observed when the vaccine and the challenge were different serotypes. However, when we compared field strains of pathogenic *Pasteurella multocida*, and particularly when we compared the patterns of metal regulated proteins present in these strains, we found two major patterns of metal regulated proteins predominate in these pathogenic strains. One pattern contains 2 or 3 major metal regulated proteins, and the other pattern contains 4 distinct metal regulated proteins. When the metal regulated proteins of these two pattern types were obtained from two *P. multocida* strains, one of serotype 2,5 and one of serotype 3,4, and combined and used to vaccinate animals, we observed heterologous protection across serotypes of *P. multocida*, including protection against a *P. multocida* of another serotype not represented in the two strains used to make the vaccine. This result was unexpected and surprising. We are not aware of any reports of this type of heterologous protection being observed in vaccines that protect against *P. multocida* infection.

[0017] Provided herein are compositions. In one embodiment, a composition includes an isolated protein having at least 80% similarity to amino acids 25-968 of SEQ ID NO:2, an isolated protein having at least 80% similarity to amino acids 27-790 of SEQ ID NO:4, an isolated protein having at least 80% similarity to amino acids 23-727 of SEQ ID NO:6, an isolated protein having at least 80% similarity to amino acids 25-964 of SEQ ID NO:8, an isolated protein having at least 80% similarity to amino acids 26-848 of SEQ ID NO:10, an isolated protein having at least 80% similarity to amino acids 27-784 of SEQ ID NO: 12, an isolated protein having at least 80% similarity to amino acids 25-742 SEQ ID NO: 14, an isolated protein having at least 80% similarity to amino acids 26-805 of SEQ ID NO: 44, or a combination thereof. The composition protects an animal, such as a chicken, against challenge with *Pasteurella multocida*.

[0018] In another embodiment, a composition includes isolated proteins having molecular weights of 99 kDa, 81 kDa, and 80 kDa where the proteins are isolatable from a *Pasteurella multocida* when incubated in media that includes an iron chelator and not isolatable when grown in the media without the iron chelator, isolated proteins having molecular weights of 109 kDa, 89 kDa, and 87 kDa wherein the proteins are isolatable from a *P. multocida* when incubated in media comprising an iron chelator, where the proteins are expressed by the *P. multocida* when incubated in media without the iron chelator and expressed at an enhanced level during growth in media that includes an iron chelator. The composition protects an animal, such as a chicken, against challenge with *P. multocida*.

[0019] In one embodiment, a includes an isolated protein having at least 80% similarity to amino acids 26-805 of SEQ ID NO:44, wherein the composition protects a chicken against challenge with *P. multocida*.

[0020] A composition described herein can optionally include additional proteins, such as at least one isolated protein having a molecular weight of 249 kDa, 60 kDa, 42 kDa, 38 kDa, 27 kDa, 26 kDa, or 22 kDa, wherein the proteins are isolatable from the *P. multocida*.

[0021] In one embodiment, a composition includes an isolated whole cell that includes at least one of the proteins of a composition described herein. The composition protects an animal, such as a chicken, against challenge with *Pas-*

*turella multocida*. In one embodiment, the whole cell is a cell engineered to express one or more of the proteins. In one embodiment, the cell is *E. coli*. Further provided is a composition that includes isolated antibody that specifically binds to a protein of a composition described herein. In one embodiment, the antibody is polyclonal antibody.

[0022] A composition described herein can further include a pharmaceutically acceptable carrier, an adjuvant, or a combination thereof.

[0023] Also provided are methods. In one embodiment, a method includes administering to a subject an amount of a composition described herein effective to induce the subject to produce antibody that specifically binds to at least one protein of the composition. In one embodiment, a method is for treating an infection in a subject, and includes administering an effective amount of the composition described herein to a subject having or at risk of having an infection caused by a *P. multocida*. In one embodiment, a method is for treating a symptom in a subject, and includes administering an effective amount of a composition described herein to a subject having or at risk of having an infection caused by a *P. multocida*. In one embodiment, a method is for decreasing colonization in a subject, and includes administering an effective amount of a composition described herein to a subject colonized by a *P. multocida*. In one embodiment, a method is for treating an infection in a subject, and includes administering an effective amount of a composition to a subject having or at risk of having an infection caused by a *P. multocida* wherein the composition includes antibody that specifically binds to a protein of a composition described herein. In one embodiment, a method is for treating a symptom in a subject, and includes administering an effective amount of a composition to a subject having or at risk of having an infection caused by a *P. multocida*, wherein the composition that includes antibody that specifically binds to a protein of a composition described herein. In one embodiment, a method is for decreasing colonization in a subject, and includes administering an effective amount of a composition to a subject colonized by a *P. multocida*, wherein the composition that includes antibody that specifically binds to a protein of a composition described herein.

[0024] In one embodiment, the subject can be a mammal, such as a bovine, or an avian, such as a chicken or a turkey. In one embodiment, at least 700 micrograms ( $\mu$ g) to no greater than 1,200  $\mu$ g of protein is administered.

[0025] Also provided are kits. In one embodiment, a kit is for detecting antibody that specifically binds a protein, and includes in separate containers (i) an isolated protein of the composition of claim 1 or 2, and (ii) a reagent that detects an antibody that specifically binds the protein. In one embodiment, a kit is for detecting a protein, and includes in separate containers (i) an antibody that specifically binds an isolated protein of a composition described herein, and a second reagent that specifically binds the protein.

[0026] The term “and/or” means one or all of the listed elements or a combination of any two or more of the listed elements.

[0027] The words “preferred” and “preferably” refer to embodiments of the invention that may afford certain benefits, under certain circumstances. However, other embodiments may also be preferred, under the same or other circumstances. Furthermore, the recitation of one or more preferred embodiments does not imply that other embodi-

ments are not useful, and is not intended to exclude other embodiments from the scope of the invention.

[0028] The terms “comprises” and variations thereof do not have a limiting meaning where these terms appear in the description and claims. It is understood that wherever embodiments are described herein with the language “include,” “includes,” or “including,” and the like, otherwise analogous embodiments described in terms of “consisting of” and/or “consisting essentially of” are also provided.

[0029] Unless otherwise specified, “a,” “an,” “the,” and “at least one” are used interchangeably and mean one or more than one.

[0030] Also herein, the recitations of numerical ranges by endpoints include all numbers subsumed within that range (e.g., 1 to 5 includes 1, 1.5, 2, 2.75, 3, 3.80, 4, 5, etc.).

[0031] For any method disclosed herein that includes discrete steps, the steps may be conducted in any feasible order. And, as appropriate, any combination of two or more steps may be conducted simultaneously.

[0032] The above summary of the present invention is not intended to describe each disclosed embodiment or every implementation of the present invention. The description that follows more particularly exemplifies illustrative embodiments. In several places throughout the application, guidance is provided through lists of examples, which examples can be used in various combinations. In each instance, the recited list serves only as a representative group and should not be interpreted as an exclusive list.

#### BRIEF DESCRIPTION OF THE FIGURES

[0033] FIG. 1. SDS-PAGE gel of extract of bacteria grown under iron limiting and iron replete conditions. Lanes 1 and 10: Broad range molecular weight markers; Lane 2: Vaccine candidate MS061130 under iron restriction; Lane 3: Vaccine candidate strain under iron replete conditions. Lane 4: serotype 1 reference strain X-73 under iron limited conditions. Lane 5: X-73 under iron replete conditions. Lane 6: Serotype 3 reference strain P1059 under iron limiting conditions. Lane 7: P1059 under iron replete conditions. Lane 8: reference strain P-1662 under iron limiting conditions; lane 9: P-1662 under iron replete conditions. Note the similar molecular weights of the iron-regulated proteins as depicted within the rectangle.

[0034] FIG. 2. Efficacy of three *Pasteurella multocida* vaccine serials produced from a single strain against challenge with heterologous *Pasteurella* serotype 1. Three vaccine serials consisting of the same antigen were prepared from the Master Seed MS061130. Note: two antigen preparation representing two separate fermentation processes were used to prepare two antigen lots designated as antigen A and antigen B. The first vaccine serial was prepared using a mixture of antigen A and antigen B, the second vaccine serial was prepared only using antigen A, and the third vaccine serial was prepared using only antigen B.

[0035] FIG. 3. Table of proteins known to be in the reference challenge strains for serotype A: 1 and serotype A: 3 compared to the vaccine strain. Note the 8 proteins present in at least one of the reference strain genomes, but absent from the vaccine strain.

[0036] FIG. 4. Venn diagram illustrating outer membrane proteins missing from the vaccine strain (MS061130). Numbers correlate to the protein identification numbers of FIG. 3.

[0037] FIG. 5. Dendrogram of metal regulated protein banding patterns of 30+ clinical field isolates and reference strains of *Pasteurella multocida*. Note the two major banding patterns as depicted as a four band pattern in brackets [ ], and a three band pattern in braces { }. Most other strains appear to be variations on these two patterns.

[0038] FIG. 6. Comparison of 38 expressed iron regulated proteins of 13 different isolates representing pattern types of FIG. 5. (+) indicates the protein is expressed, and (-) indicates the protein is not expressed at detectable levels. The ellipse identifies two potential vaccine targets that cover the majority of proteins.

[0039] FIG. 7. Vaccine efficacy using a vaccine composed of two Master seed strains challenged with *Pasteurella* serotype 1 in chickens showing heterologous protection.

[0040] FIGS. 8-1, 8-2, 9-1, 9-2, 10-1, 10-2, 11-1, 11-2, 12-1, 12-2, 13-1, 13-2, 14-1 and 14-2. The amino acid sequences of SEQ ID NOs: 2, 4, 6, 8, 10, 12, and 14, and an example of a nucleotide sequence (SEQ ID NOs: 1, 3, 5, 7, 9, 11, and 13, respectively) encoding each amino acid sequence.

[0041] FIGS. 15-1 and 15-2. Sequence alignment of SEQ ID NO:2 and four other sequences. WP\_053521090.1 (SEQ ID NO:15), 1121\_HgbA\_968aa (SEQ ID NO:2), WP\_005756141.1 (SEQ ID NO:16), AAQ14873.1 (SEQ ID NO:17), WP\_061405928.1 (SEQ ID NO:18). All sequences are *P. multocida*, except WP\_053521090.1 which is *Hemophilus influenzae*.

[0042] FIGS. 16-1 and 16-2. Sequence alignment of SEQ ID NO:4 and four other sequences. ESQ71136.1 (SEQ ID NO:19), WP\_014391205.1 (SEQ ID NO:20), WP\_016534044.1 (SEQ ID NO:21), 1121\_FepA\_790aa (SEQ ID NO:4), WP\_005756883.1 (SEQ ID NO:22). All sequences are *P. multocida*.

[0043] FIGS. 17-1 and 17-2. Sequence alignment of SEQ ID NO:6 and four other sequences. WP\_071522773.1 (SEQ ID NO:23), WP\_016534444.1 (SEQ ID NO:24), WP\_005755819.1 (SEQ ID NO:25), 1121\_PfhR\_727aa (SEQ ID NO:6), EGP05580.1 (SEQ ID NO:26). All sequences are *P. multocida*.

[0044] FIGS. 18-1, 18-2 and 18-3. Sequence alignment of SEQ ID NO:8 and five other sequences. 1135\_PM0300\_964aa (SEQ ID NO:8), WP\_005753642.1 (SEQ ID NO:27), WP\_016534557.1 (SEQ ID NO:28), WP\_010906573.1 (SEQ ID NO:29), AAQ14873.1 (SEQ ID NO:30), 1121\_HgbA\_968aa (SEQ ID NO:2). All sequences are *P. multocida*.

[0045] FIGS. 19-1 and 19-2. Sequence alignment of SEQ ID NO:10 and four other sequences. WP\_017861186.1 (SEQ ID NO:31), AAU29202.1 (SEQ ID NO:32), EGP04511.1 (SEQ ID NO:33), 1135\_HasR\_848aa (SEQ ID NO:10), WP\_005752163.1 (SEQ ID NO:34). All sequences are *P. multocida*.

[0046] FIGS. 20-1 and 20-2. Sequence alignment of SEQ ID NO: 12 and four other sequences. WP\_005751557.1 (SEQ ID NO:35), 1135\_PM0741\_784aa (SEQ ID NO:12), WP\_016534554.1 (SEQ ID NO:36), WP\_064972816.1 (SEQ ID NO:37), WP\_074865020.1 (SEQ ID NO:38). All sequences are *P. multocida*.

[0047] FIGS. 21-1 and 21-2. Sequence alignment of SEQ ID NO:14 and four other sequences. WP\_050948957.1 (SEQ ID NO:39), WP\_014391043.1 (SEQ ID NO:40), WP\_016533738.1 (SEQ ID NO:41), 1135\_P1062\_0207600\_742aa (SEQ ID NO:14), WP\_025248456.1 (SEQ

ID NO:42). All sequences are *P. multocida*, except WP 050948957.1 which is *Haemophilus influenzae*.

[0048] FIGS. 22-1 and 22-2. The amino acid sequences of SEQ ID NO:44 and an example of a nucleotide sequence (SEQ ID NO:43) encoding the amino acid sequence. The first 78 nucleotides of SEQ ID NO:43 are predicted to encode the signal sequence of SEQ ID NO: 44 (amino acids 1-26).

[0049] FIG. 23. The percent survival of chickens challenged with *Pasteurella Multocida* comparing a recombinant zinc (rZinc) protein to SRP extract. The results clearly show the efficacy of both the *Pasteurella Multocida* SRP extract and the rZinc protein compared to non-vaccinated Placebo controls.

[0050] FIG. 24. Gel image showing the inclusion body preparation of *Pasteurella Multocida* recombinant zinc acquisition protein. Lane 1: Un-Induced Whole Cell, lane 2: Induced Whole Cell, Lane 3: Lysed Cell Supernatant, Lane 4: 1x Bugbuster Wash, Lane 5: ½x Bugbuster Wash, Lane 6: TBW Wash, Lane 7: Solubilized Inclusion Bodies and Lane 8: Pellet Post Solubilization showing the recombinant protein with a molecular weight of 90.85 kDa.

[0051] FIG. 25. Sequence alignment of the zinc affinity region of zinc acquisition proteins from *Mannheimia haemolytica* (ZAP, SEQ ID NO:45) and 10 other respiratory pathogens. AHG81836.1 (SEQ ID NO:46), WP\_005612269.1 (SEQ ID NO: 47), WP\_027074597.1 (SEQ ID NO:48), AHG73391.1 (SEQ ID NO:49), WP\_021114857.1 (SEQ ID NO:50), WP\_026212957.1 (SEQ ID NO:51), WP\_028858792.1 (SEQ ID NO:52), WP\_016534590.1 (SEQ ID NO:53), KDN24548.1 (SEQ ID NO:54), and WP\_027021676.1 (SEQ ID NO:55).

#### DETAILED DESCRIPTION OF ILLUSTRATIVE EMBODIMENTS

[0052] In one aspect, this disclosure provides proteins and compositions including proteins. As used herein, “protein” refers to a polymer of amino acids linked by peptide bonds. Thus, for example, the terms peptide, oligopeptide, polypeptide, and enzyme are included within the definition of protein. This term also includes proteins that may include one or more post-expression modifications of the protein such as, for example, a glycosylation, an acetylation, a phosphorylation, and the like. The term protein does not connote a specific length of a polymer of amino acids. A protein may be isolatable directly from a natural source or can be prepared with the aid of recombinant, enzymatic, or chemical techniques. In the case of a protein that is naturally occurring, such a protein is typically isolated.

[0053] An “isolated” protein is one that has been removed from its natural environment. For instance, an isolated protein is a protein that has been removed from the cytoplasm or from the membrane of a cell, and many of the proteins, nucleic acids, and other cellular material of its natural environment are no longer present.

[0054] A protein characterized as “isolatable” from a particular source is a protein that, under appropriate conditions, is produced by the identified source, although the protein may be obtained from alternate sources using, for example, conventional recombinant, chemical, or enzymatic techniques. Thus, characterizing a protein as “isolatable” from a particular source does not imply any specific source

from which the protein must be obtained or any particular conditions or processes under which the protein must be obtained.

[0055] A “purified” protein is one that is at least 60% free, preferably at least 75% free, and most preferably at least 90% free from other components with which they are naturally associated. Proteins that are produced outside the organism in which they naturally occur, e.g., through chemical or recombinant means, are considered to be isolated and purified by definition, since they were never present in a natural environment.

[0056] Generally, a protein may be characterized by molecular weight, amino acid sequence, mass fingerprint, nucleic acid that encodes the protein, immunological activity, or any combination of two or more such characteristics. The molecular weight of a protein, typically expressed in kilodaltons (kDa), can be determined using routine methods including, for instance, gel filtration, gel electrophoresis including sodium dodecyl sulfate (SDS) polyacrylamide gel electrophoresis (PAGE), capillary electrophoresis, mass spectrometry, liquid chromatography (including HPLC), and calculating the molecular weight from an observed or predicted amino acid sequence. Unless indicated otherwise, reference to molecular weight refers to molecular weight as determined by resolving a protein using an SDS polyacrylamide gel having a stacking gel of about 4% and a resolving gel of about 10% under reducing and denaturing conditions. In one embodiment, the molecular weight of a protein identified by SDS-PAGE includes molecular weights of 1, 2, 3, 4, or 5 kDa above and below the stated value.

[0057] The proteins described herein may be metal-regulated. As used herein, a “metal-regulated protein” is a protein that is expressed by a microbe at a greater level when the microbe is grown in low metal conditions compared to when the same microbe is grown in high metal conditions. Low metal and high metal conditions are described herein. For instance, certain metal-regulated proteins produced by *P. multocida* are not expressed at detectable levels during growth of the microbe in high metal conditions but are expressed at detectable levels during growth in low metal conditions. In one embodiment, a metal-regulated protein can be a siderophore receptor protein. Examples of metal-regulated proteins isolatable from a *P. multocida* after growth in low iron conditions include metal-regulated proteins having molecular weights of 104 kDa to 75 kDa. Specific examples of metal-regulated proteins isolatable from a *P. multocida* after growth in low iron conditions include proteins of 99 kDa, 81 kDa, and 80 kDa as determined by resolving a protein using an SDS polyacrylamide gel having a stacking gel of about 4% and a resolving gel of about 10% under reducing and denaturing conditions. In one embodiment, a low iron conditions is growth in the presence of 2,2'-dipyridyl. Examples of the iron regulated proteins having molecular weights of 99 kDa, 81 kDa, and 80 kDa, and nucleotide sequences encoding the proteins, are shown in FIGS. 10, 14, and 6, respectively. A specific example of metal-regulated proteins isolatable from a *P. multocida* after growth in low zinc conditions include a protein of 91 kDa as determined by resolving a protein using an SDS polyacrylamide gel having a stacking gel of about 4% and a resolving gel of about 10% under reducing and denaturing conditions. In one embodiment, a low zinc conditions is growth in the presence of N,N,N',N'-tetrakis(2-pyridylmethyl)ethylenediamine (TPEN) (Sigma-Aldrich, St. Louis MO). An example

of the zinc regulated protein having a molecular weight of 91 kDa, and nucleotide sequences encoding the protein, is shown in FIG. 22.

[0058] The proteins described herein may be expressed at detectable levels during growth of the microbe in high metal conditions but expressed at higher levels during growth in low metal conditions. The expression of such proteins is referred to herein as “enhanced” during growth in low metal conditions. Typically, the increase in expression of a protein during growth in low metal conditions is between 50% and 500% compared to the expression of the protein during growth in high metal conditions.

[0059] Examples of metal-regulated proteins having enhanced expression and isolatable from *P. multocida* after growth in low iron conditions include metal-regulated proteins having molecular weights of and 114 kDa to 82 kDa. Specific examples of metal-regulated proteins isolatable from a *P. multocida* after growth in low iron conditions include proteins of 109 kDa, 89 kDa, and 87 kDa as determined by resolving a protein using an SDS polyacrylamide gel having a stacking gel of about 4% and a resolving gel of about 10% under reducing and denaturing conditions. Examples of the proteins having molecular weights of 109 kDa and nucleotide sequences encoding them are shown in FIGS. 8-1, 8-2, and 11-1, 11-2. Examples of the proteins having molecular weights of 89 kDa and 87 kDa, and nucleotide sequences encoding the proteins, are shown in FIGS. 13 and 9, respectively.

[0060] This disclosure also describes certain proteins that are not metal-regulated. Such proteins are expressed in the presence of a metal ion such as, for example, in the presence of ferric chloride, and also expressed when grown in low iron conditions. Examples of this type of protein isolatable from *P. multocida* have molecular weights of 254 kDa to 244 kDa, 65 kDa to 55 kDa, and 47 kDa to 17 kDa. Examples of molecular weights of this type of protein include 249 kDa, 60 kDa, 42 kDa, 38 kDa, 27 kDa, 26 kDa, and 22 kDa. Additional examples of proteins include recombinantly-produced versions of proteins described herein. A recombinantly-produced protein may include the entire amino acid sequence translatable from an mRNA transcript. Alternatively, a recombinantly-produced protein can include a fragment of the entire translatable amino acid sequence. For example, a recombinantly-produced protein may lack a cleavable sequence at either terminus of the protein—e.g., a cleavable signal sequence at the amino terminus of the protein.

[0061] In one embodiment, a protein lacks one or more amino acids from the amino terminus of the protein encoded by a coding sequence obtained from a wild-type cell, e.g., the protein lacks a signal sequence. Thus, a fragment can lack at least one, at least two, at least three, at least four, at least five, at least six, at least seven, at least eight, at least nine, at least ten, at least 11, at least 12, at least 13, at least 14, at least 15, at least 16, at least 17, at least 18, at least 19, at least 20, at least 21, at least 22, at least 23, at least 24, at least 25, at least 26, at least 27, at least 28, at least 29, at least 30, at least 31, at least 32, at least 33, at least 34, at least 35, at least 36, at least 37, at least 38, at least 39, at least 40, at least 41, at least 42, at least 43, at least 44, at least 45, at least 46, at least 47, at least 48, at least 49, at least 50, at least 51, at least 52, at least 53, at least 54, at least 55, at least 56, at least 57, at least 58, at least 59, at least 60, at least 61, at least 62, or at least 63 amino acids from the amino terminus of the

protein. In one embodiment, a fragment of the protein depicted at SEQ ID NO:2 does not include amino acids 1-24. In one embodiment, a fragment of the protein depicted at SEQ ID NO:4 does not include amino acids 1-26. In one embodiment, a fragment of the protein depicted at SEQ ID NO:6 does not include amino acids 1-22. In one embodiment, a fragment of the protein depicted at SEQ ID NO:8 does not include amino acids 1-24. In one embodiment, a fragment of the protein depicted at SEQ ID NO:10 does not include amino acids 1-25. In one embodiment, a fragment of the protein depicted at SEQ ID NO:12 does not include amino acids 1-26. In one embodiment, a fragment of the protein depicted at SEQ ID NO:14 does not include amino acids 1-24. In one embodiment, a fragment of the protein depicted at SEQ ID NO:44 does not include amino acids 1-26.

[0062] Whether a protein is a metal-regulated protein, an enhanced protein, or a non-metal-regulated protein can be determined by methods useful for comparing the presence of proteins, including, for example, gel filtration, gel electrophoresis including sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE), capillary electrophoresis, mass spectrometry, isobaric tags for relative and absolute quantification (iTRAQ), and liquid chromatography including HPLC. Separate cultures of a microbe can be grown under high metal conditions and under low metal conditions, proteins may be isolated as described herein, and the proteins present in each culture can be resolved and compared. Typically, an equal amount of proteins from each culture is used. In one embodiment, the proteins can be resolved using an SDS polyacrylamide gel having a stacking gel of about 4% and a resolving gel of about 10% under reducing and denaturing conditions. For instance, 30 micrograms ( $\mu$ g) of total protein from each culture may be used and loaded into wells of a gel. After running the gel and staining the proteins with Coomassie Brilliant Blue, the two lanes can be compared. When determining whether a protein is or is not expressed at a detectable level, 30  $\mu$ g of total protein from a culture is resolved on an SDS-PAGE gel and stained with Coomassie Brilliant Blue using methods known in the art. A protein that can be visualized by eye is considered to be expressed at a detectable level, while a protein that cannot be visualized by eye is considered to not be expressed at a detectable level.

[0063] Alternatively, whether a protein is a metal-regulated protein or a non-metal-regulated protein can be determined using microarray-based gene expression analysis. Separate cultures of a microbe can be grown under high metal conditions and under low metal conditions, RNA can be extracted from cells of each culture, and differences in RNA expression in cells grown in high metal conditions versus RNA expression in cells grown in low metal conditions can be detected and compared. For example, labeled cDNA can be prepared from 8-10  $\mu$ g of bacterial RNA using established protocols. The labeled cDNA can be applied to a microarray of a microbe's genome. Such microarrays are commercially available and evaluating gene expression using such arrays is routine.

[0064] The proteins described herein can have immunological activity. “Immunological activity” refers to the ability of a protein to elicit an immunological response in an animal. An immunological response to a protein is the development in an animal of a cellular and/or antibody-mediated immune response to the protein. Usually, an

immunological response includes but is not limited to one or more of the following effects: the production of antibodies, B cells, helper T cells, suppressor T cells, and/or cytotoxic T cells, directed to an epitope or epitopes of the protein. "Epitope" refers to the site on an antigen to which specific B cells and/or T cells respond so that antibody is produced. The immunological activity may be protective. "Protective immunological activity" refers to the ability of a protein to elicit an immunological response in an animal that inhibits or limits infection by *P. multocida*. Whether a protein has protective immunological activity can be determined by methods known in the art such as, for example, methods described in Examples 6-9 and 15-16. A protein may have seroactive activity. As used herein, "seroactive activity" refers to the ability of a candidate protein to react with antibody present in convalescent serum from an animal infected with a *P. multocida*.

[0065] A protein as described herein may have the characteristics of a reference protein. The characteristics can include, for example, molecular weight, mass fingerprint, amino acid sequence, or any combination thereof. In one embodiment, a protein can be obtained from a microbe. A microbe can express one, two, three, four, five, six, seven, eight, or nine of the proteins described herein. The reference protein can be isolatable from a gram negative microbe, preferably a member of the family Pasteurellaceae, including the genus *Pasteurella*, *Mannheimia*, or *Haemophilus*. A member of the genus *Pasteurella Mannheimia*, and *Haemophilus* is also referred to herein as *Pasteurella spp.*, *Mannheimia spp.*, and *Haemophilus spp.*, respectively. In one embodiment, the reference protein is expressed by a *P. multocida*. In one embodiment, the *P. multocida* can be serotype 2,5 or serotype 3,4.

[0066] As used herein, a protein may be "structurally similar" to a reference protein if the amino acid sequence of the protein possesses a specified amount of sequence similarity and/or sequence identity compared to the reference protein. A protein also may be "structurally similar" to a reference protein if the protein exhibits a mass fingerprint possessing a specified amount of identity compared to a comparable mass fingerprint of the reference protein. Thus, a protein may be "structurally similar" to a reference protein if, compared to the reference protein, it possesses a sufficient level of amino acid sequence identity, amino acid sequence similarity, or a combination thereof. In one embodiment, a protein described herein can have an amino acid sequence that is structurally similar, as described below, to amino acids 25-968 of SEQ ID NO:2, amino acids 27-790 of SEQ ID NO: 4, amino acids 23-727 of SEQ ID NO:6, amino acids 25-964 of SEQ ID NO:8, amino acids 26-848 of SEQ ID NO: 10, amino acids 27-784 of SEQ ID NO:12, amino acids 25-742 of SEQ ID NO: 14, or amino acids 26-805 of SEQ ID NO:44.

#### Protein Sequence Similarity and Protein Sequence Identity

[0067] Structural similarity of two proteins can be determined by aligning the residues of the two proteins (for example, a candidate protein and any appropriate reference protein described herein) to optimize the number of identical amino acids along the lengths of their sequences; gaps in either or both sequences are permitted in making the alignment in order to optimize the number of identical amino acids, although the amino acids in each sequence must nonetheless remain in their proper order. A reference protein

may be a protein described herein or any known metal-regulated protein, as appropriate. Examples of reference proteins include, but are not limited to, amino acids 25-968 of SEQ ID NO:2, amino acids 27-790 of SEQ ID NO:4, amino acids 23-727 of SEQ ID NO:6, amino acids 25-964 of SEQ ID NO:8, amino acids 26-848 of SEQ ID NO:10, amino acids 27-784 of SEQ ID NO: 12, amino acids 25-742 of SEQ ID NO: 14, and amino acids 26-805 of SEQ ID NO: 44. A candidate protein is the protein being compared to the reference protein. A candidate protein can be isolated, for example, from a microbe, or can be produced using recombinant techniques, or chemically or enzymatically synthesized.

[0068] Unless modified as otherwise described herein, a pair-wise comparison analysis of amino acid sequences can be carried out using the BESTFIT algorithm in the GCG package (version 10.2, Madison WI). Alternatively, proteins may be compared using the Blastp program of the BLAST 2 search algorithm, as described by Tatiana et al. (*FEMS Microbiol Lett*, 174:247-250 (1999)), and available on the National Center for Biotechnology Information (NCBI) website. The default values for all BLAST 2 search parameters may be used, including matrix=BLOSUM62; open gap penalty=11, extension gap penalty=1, gap x\_dropoff=50, expect=10, wordsize=3, and filter on.

[0069] In the comparison of two amino acid sequences, structural similarity may be referred to by percent "identity" or may be referred to by percent "similarity." "Identity" refers to the presence of identical amino acids. "Similarity" refers to the presence of not only identical amino acids but also the presence of conservative substitutions. A conservative substitution for an amino acid in a protein may be selected from other members of the class to which the amino acid belongs. For example, it is well-known in the art of protein biochemistry that an amino acid belonging to a grouping of amino acids having a particular size or characteristic (such as charge, hydrophobicity, or hydrophilicity) can be substituted for another amino acid without altering the activity of a protein, particularly in regions of the protein that are not directly associated with activity. For example, nonpolar (hydrophobic) amino acids include alanine, leucine, isoleucine, valine, proline, phenylalanine, tryptophan, and tyrosine. Polar neutral amino acids include glycine, serine, threonine, cysteine, tyrosine, asparagine and glutamine. The positively charged (basic) amino acids include arginine, lysine and histidine. The negatively charged (acidic) amino acids include aspartic acid and glutamic acid. Conservative substitutions include, for example, Lys for Arg and vice versa to maintain a positive charge; Glu for Asp and vice versa to maintain a negative charge; Ser for Thr so that a free —OH is maintained; and Gln for Asn to maintain a free —NH<sub>2</sub>. Likewise, biologically active analogs of a protein containing deletions or additions of one or more contiguous or noncontiguous amino acids that do not eliminate a functional activity—such as, for example, immunological activity—of the protein are also contemplated.

[0070] Thus, as used herein, a protein as described herein and/or the amino acid sequence of one or more SEQ ID NOs can include a protein with at least 50%, at least 55%, at least 60%, at least 65%, at least 70%, at least 75%, at least 80%, at least 85%, at least 86%, at least 87%, at least 88%, at least 89%, at least 90%, at least 91%, at least 92%, at least 93%, at least 94%, at least 95%, at least 96%, at least 97%, at least

98%, or at least 99% amino acid sequence similarity to the reference amino acid sequence.

[0071] Alternatively, as used herein, a protein as described herein and/or the amino acid sequence of one or more SEQ ID NOs can include a protein with at least 50%, at least 55%, at least 60%, at least 65%, at least 70%, at least 75%, at least 80%, at least 85%, at least 86%, at least 87%, at least 88%, at least 89%, at least 90%, at least 91%, at least 92%, at least 93%, at least 94%, at least 95%, at least 96%, at least 97%, at least 98%, or at least 99% amino acid sequence identity to the reference amino acid sequence.

[0072] FIGS. 15-21 show Clustal Omega alignments for proteins having the amino acid sequences shown in SEQ ID NOs: 2, 4, 6, 8, 10, 12, and 14. The alignments indicate amino acids that are conserved in the variants of each protein across different *P. multocida* strains and, for FIGS. 15 and 21, *Haemophilus influenzae*. In FIGS. 15-21 an asterisk (\*) indicates positions which have a single, fully conserved residue, a colon (:) indicates conservation between groups of strongly similar properties as roughly equivalent to scoring >0.5 in the Gonnet PAM 250 matrix, and a period (.) indicates conservation between groups of weakly similar properties as roughly equivalent to scoring <0.5 and >0 in the Gonnet PAM 250 matrix. A person of ordinary skill in the art can deduce from such data regions of the protein in which substitutions, particularly conservative substitutions, may be permitted without unduly affecting activity (immunological activity, protective immunological activity, or seroactive activity of the modified protein).

[0073] Consequently, a protein as described herein can include certain variants including, for example, homologous proteins that originate—biologically and/or recombinantly—from microbial species or strains other than the microbial species or strain from which the protein was originally isolated and/or identified.

[0074] A protein described herein also can be designed to provide one or more additional sequences such as, for example, the addition of coding sequences for added C-terminal and/or N-terminal amino acids that may facilitate purification by trapping on columns or use of antibodies. Such tags include, for example, histidine-rich tags that allow purification of proteins on nickel columns. Such gene modification techniques and suitable additional sequences are well known in the molecular biology arts. A protein as described herein also may be designed so that certain amino acids at the C-terminus and/or N-terminus are deleted.

[0075] A protein described herein can include a “modification.” A modification refers to a chemical or enzymatic derivatization at one or more constituent amino acids. Such a modification can include, for example, a side chain modification, a backbone modification, an N-terminal modification, and/or a C-terminal modification such as, for example, acetylation, hydroxylation, methylation, amidation, and the attachment of a carbohydrate and/or lipid moiety, a cofactor, and the like, and combinations thereof. Modified proteins as described herein may retain the biological activity—such as, for example, immunological activity—of the unmodified protein or may exhibit a reduced or increased biological activity compared to the unmodified protein.

[0076] A protein as described herein (including a biologically active analog thereof and/or a modification thereof) can include a native (naturally occurring), a recombinant, a chemically synthesized, or an enzymatically synthesized protein. For example, a protein as described herein may be

prepared by isolating the protein from a natural source or may be prepared recombinantly by conventional methods including, for example, preparation as fusion proteins in bacteria or other host cells.

#### Polynucleotide Sequence Similarity and Polynucleotide Sequence Identity

[0077] Proteins as described herein also can be identified in terms of the polynucleotide that encodes the protein. Thus, this disclosure provides polynucleotides that encode a protein as described herein or hybridize, under standard hybridization conditions, to a polynucleotide that encodes a protein as described herein, and the complements of such polynucleotide sequences.

[0078] As used herein, a polynucleotide as described herein and/or the nucleic acid sequence of one or more SEQ ID NOs: 1, 3, 5, 7, 9, 11, 13, or 43, or a fragment thereof, can include polynucleotides having a sequence identity of at least 50%, at least 55%, at least 60%, at least 65%, at least 70%, at least 75%, at least 80%, at least 85%, at least 86%, at least 87%, at least 88%, at least 89%, at least 90%, at least 91%, at least 92%, at least 93%, at least 94%, at least 95%, at least 96%, at least 97%, at least 98%, or at least 99% sequence identity to an identified reference polynucleotide sequence.

[0079] In this context, “sequence identity” refers to the identity between two polynucleotide sequences. Sequence identity is generally determined by aligning the bases of the two polynucleotides (for example, aligning the nucleotide sequence of the candidate sequence and a nucleotide sequence that includes, for example, a nucleotide sequence disclosed herein, such as SEQ ID NO:1, 3, 5, 7, 9, 11, 13, or 43, or a fragment thereof) to optimize the number of identical nucleotides along the lengths of their sequences; gaps in either or both sequences are permitted in making the alignment in order to optimize the number of shared nucleotides, although the nucleotides in each sequence must nonetheless remain in their proper order. A candidate sequence is the sequence being compared to a known sequence—e.g., a nucleotide sequence that includes a nucleotide sequence described herein, for example, SEQ ID NO: 1, 3, 5, 7, 9, 11, or 13, or 43, or a fragment thereof. For example, two polynucleotide sequences can be compared using the Blastn program of the BLAST 2 search algorithm, as described by Tatusova et al., (*FEMS Microbiol Lett.*, 174:247-250 (1999)), and available on the world wide web at ncbi.nlm.nih.gov/BLAST/. The default values for all BLAST 2 search parameters may be used, including reward for match=1, penalty for mismatch=-2, open gap penalty=5, extension gap penalty=2, gap x dropoff=50, expect=10, wordsize=11, and filter on.

[0080] Finally, a polynucleotide as described herein can include any polynucleotide that encodes a protein as described herein. Thus, the nucleotide sequence of the polynucleotide may be deduced from the amino acid sequence that is to be encoded by the polynucleotide.

[0081] This disclosure also provides whole cell preparations of microbes. In one embodiment, the preparation includes a microbe that has been engineered to express the proteins described herein. For instance, a microbe can be engineered to express proteins having structural similarity with SEQ ID NOs: 2, 4, 6, 8, 10, 12, 14, or 44, or a fragment thereof, or any subcombination of those eight proteins. The microbe can be any microbial cell that is amenable to genetic

manipulation, including a gram negative or a gram positive microbe. Examples of gram negative microbes include, but are not limited to, *E. coli*, *Salmonella* spp., and *Pasteurella* spp., such as *P. multocida*.

[0082] In one embodiment, the preparation includes two or more populations of microbes. Each of the populations do not express all of SEQ ID NOs: 2, 4, 6, 8, 10, 12, and 14, or a fragment thereof. Rather, each of the populations express a subset of the seven proteins, and the two or more populations when considered as a whole express the seven proteins. For instance, in one embodiment one population of microbe expresses SEQ ID NOs: 2, 4, and 6, and a second population of microbe expresses SEQ ID NOs: 8, 10, 12, and 14, or a fragment thereof. A population can be a wild-type microbe or an engineered microbe. A preparation can include one or more wild-type microbes, one or more engineered microbes, or a combination of wild-type and engineered microbes. Examples of wild-type cells include members of the genus *Pasteurella*, such as *P. multocida*. In one embodiment, the cell is an attenuated *P. multocida*. The inventors have determined that administering certain types of *Pasteurella* strains as whole cells is expected to result in immunological properties that are different than the result of administering the individual strains separately. The different immunological properties include the ability to protect against *P. multocida* strains having a different serotype than the *P. multocida* cells administered to the animal. In one embodiment, one whole cell is *P. multocida* serotype 2,5 and a second whole cell is *P. multocida* serotype 3,4, and one expresses three of the seven proteins and the other expresses four of the seven proteins. In one embodiment, one of more of the populations of microbes can express SEQ ID NO:44, or a fragment thereof.

[0083] The cells present in a whole cell preparation may be inactivated such that the cells cannot replicate but the immunological activity of the proteins as described herein expressed by the microbe is maintained. Typically, the cells may be killed by exposure to agents such as glutaraldehyde, formalin, or formaldehyde.

#### Compositions

[0084] A composition can include one protein isolated described herein, at least two isolated proteins described herein, or a number of proteins that is an integer greater than two (e.g., at least two, at least three, at least four, at least five, at least six, at least seven, at least eight, at least nine, at least 10, at least 11, at least 12, at least 13, at least 14, at least 15, at least 16, at least 17, at least 18, at least 19, or at least 20, and so on). In one embodiment, a composition including one protein described herein includes a protein identical to or having structural similarity with SEQ ID NO:44, or a fragment thereof. Unless a specific level of sequence similarity and/or identity is expressly indicated herein (e.g., at least 80% sequence similarity, at least 90% sequence identity, etc.), reference to the amino acid sequence of an identified SEQ ID NO includes variants having the levels of sequence similarity and/or the levels of sequence identity described herein in the section headed "Protein sequence similarity and protein sequence identity."

[0085] A recombinantly-produced protein may be expressed from a vector that permits expression of the protein when the vector is introduced into an appropriate host cell. A host cell may be constructed to produce one or more recombinantly-produced proteins as described herein

and, therefore, can include one or more vectors that include at least one polynucleotide encoding a protein described herein. Thus, each vector can include one or more polynucleotides as described herein—i.e., a polynucleotide that encodes a protein as described herein. Methods for the genetic manipulation of microbes, such as *P. multocida*, are known and routine in the art.

[0086] Certain compositions such as, for example, those including recombinantly-produced proteins, can include a maximum number of proteins. In some embodiments, the maximum number of proteins can refer to the maximum total number of proteins. Certain compositions can include, for example, no more than 50 proteins such as, for example, no more than 40 proteins, no more than 30 proteins, no more than 25 proteins, no more than 20 proteins, no more than 17 proteins, no more than 16 proteins, no more than 15 proteins, no more than 14 proteins, no more than 13 proteins, no more than 12 proteins, no more than 11 proteins, no more than 10 proteins, no more than nine proteins, no more than eight proteins, no more than seven proteins, no more than six proteins, no more than five proteins, no more than four proteins, no more than three proteins, no more than two proteins, or no more than one protein. In other embodiments, a maximum number of recombinantly-produced proteins may be specified in a similar manner. In still other embodiments, a maximum number of non-recombinantly-produced proteins may be specified in a similar manner.

[0087] In one embodiment, a composition can include proteins isolatable from one microbe when the microbe is engineered to express one or more proteins identical to or having structural similarity with a protein described herein. In one embodiment, the microbe is engineered to express proteins identical to or having structural similarity with one of more of SEQ ID NOs: 2, 4, 6, 8, 10, 12, 14, and 44, or a fragment thereof. In one embodiment, the microbe expresses a protein identical to or having structural similarity with SEQ ID NO:44, or a fragment thereof. In one embodiment, a composition can include proteins isolatable from two or more microbes. For instance, a composition can include proteins isolatable from two or more wild-type *P. multocida*.

[0088] In certain embodiments, a composition can include a preparation of whole cells. In one embodiment, the preparation is a whole cell that has been engineered to express one or more proteins identical to or having structural similarity with a protein described herein. In one embodiment, the microbe is engineered to express proteins identical to or have structural similarity with SEQ ID NOs: 2, 4, 6, 8, 10, 12, and 14, or a fragment thereof. In one embodiment, the microbe is engineered to express proteins are identical to or have structural similarity with SEQ ID NOs: 2, 4, 6, 8, 10, 12, 14, and 44, or a fragment thereof. In one embodiment, the microbe expresses a protein identical to or having structural similarity with SEQ ID NO: 44, or a fragment thereof. In one embodiment, the preparation is two or more populations of microbes where each of the populations express a subset of the proteins, e.g., the seven proteins (proteins identical to or having structural similarity with SEQ ID NOs: 2, 4, 6, 8, 10, 12, and 14, or a fragment thereof), or the eight proteins (proteins identical to or having structural similarity with SEQ ID NOs: 2, 4, 6, 8, 10, 12, 14, and 44, or a fragment thereof) and the two or more populations when considered as a whole express the seven proteins. In some of these embodiments, the whole cell can

be a *P. multocida*, such as a wild-type *P. multocida*. In some embodiments, a composition can include whole cell preparations from two, three, four, five, or six strains.

[0089] Specific examples of compositions include, but are not limited to, the following. In one embodiment, a composition includes metal regulated proteins and enhanced proteins. The metal regulated proteins are isolatable from *P. multocida* after growth in low iron conditions and have molecular weights of 104 kDa to 75 kDa, such as 99 kDa, 81 kDa, and 80 kDa. The proteins can be, or have structural similarity with, a fragment of SEQ ID NOs: 6, 10, or 14. The enhanced proteins are isolatable from *P. multocida* after growth in low iron conditions and have molecular weights of 156 kDa to 146 kDa, such as 151 kDa, and 114 kDa to 82 kDa, such as 109 kDa, 89 kDa, and 87 kDa. The proteins can be, or have structural similarity with, a fragment of SEQ ID NOs: 2, 4, 8, or 12. Optionally, a composition can include non-metal regulated proteins that are isolatable from *P. multocida*. The non-metal regulated proteins can have molecular weights of 254 kDa to 244 kDa, such as 249 kDa, 65 kDa to 55 kDa, such as 60 kDa, and 47 kDa to 17 kDa, such as 42 kDa, 38 kDa, 27 kDa, 26 kDa, and 22 kDa. Another specific example of a composition is one that includes a protein that is, or has structural similarity with, a fragment of SEQ ID NO:44, or a fragment thereof.

[0090] Optionally, a protein described herein can be covalently bound to a carrier protein to improve the immunological properties of the protein. Useful carrier proteins are known in the art. The chemical coupling of a protein described herein can be carried out using known and routine methods. For instance, various homobifunctional and/or heterobifunctional cross-linker reagents such as bis(sulfosuccinimidyl) suberate, bis(diazobenzidine), dimethyl adipimidate, dimethyl pimelimidate, dimethyl superimidate, disuccinimidyl suberate, glutaraldehyde, m-maleimidobenzoyl-N-hydroxysuccinimide, sulfo-m-maleimidobenzoyl-N-hydroxysuccinimide, sulfosuccinimidyl 4-(N-maleimidomethyl)cycloheane-1-carboxylate, sulfosuccinimidyl 4-(p-maleimido-phenyl) butyrate and (1-ethyl-3-(dimethyl-amino-propyl) carbodiimide can be used (Harlow and Lane, Antibodies, A Laboratory Manual, generally and Chapter 5, Cold Spring Harbor Laboratory, Cold Spring Harbor, New York, NY (1988)).

[0091] A compositions described herein can include low concentrations of lipopolysaccharide (LPS). LPS is a component of the outer membrane of most gram negative microbes (see, for instance, Nikaido and Vaara, Outer Membrane, In: *Escherichia coli* and *Salmonella typhimurium*, Cellular and Molecular Biology, Neidhardt et al., (eds.) American Society for Microbiology, Washington, D.C., pp. 7-22 (1987), and typically includes polysaccharides (O-specific chain, the outer and inner core) and the lipid A region. The lipid A component of LPS is the most biologically active component of the LPS structure and together induces a wide spectrum of pathophysiological effects in mammals. The most dramatic effects are fever, disseminated intravascular coagulation, complement activation, hypotensive shock, and death. The non-specific immunostimulatory activity of LPS can enhance the formation of a granuloma at the site of administration of compositions that include LPS. Such reactions can result in undue stress on the animal by which the animal may back off feed or water for a period of time, and exasperate infectious conditions in the animal. In addition, the formation of a granuloma at the site of injection can

increase the likelihood of possible down grading of the carcass due to scarring or blemishes of the tissue at the injection site.

[0092] The concentration of LPS can be determined using routine methods known in the art. Such methods typically include measurement of dye binding by LPS (see, for instance, Keler and Nowotny, *Analyt. Biochem.*, 156, 189 (1986)) or the use of a *Limulus* amebocyte lysate (LAL) test (see, for instance, Endotoxins and Their Detection With the Limulus Amebocyte Lystate Test, Alan R. Liss, Inc., 150 Fifth Avenue, New York, NY (1982)). There are four basic commercially available methods that are typically used with an LAL test: the gel-clot test; the turbidimetric (spectrophotometric) test; the colorimetric test; and the chromogenic test. An example of a gel-clot assay is available under the tradename E-TOXATE (Sigma Chemical Co., St. Louis, MO; see Sigma Technical Bulletin No. 210), and PYROTELL (Associates of Cape Cod, Inc., East Falmouth, MA). Typically, assay conditions include contacting the composition with a preparation containing a lysate of the circulating amebocytes of the horseshoe crab, *Limulus polyphemus*. When exposed to LPS, the lysate increases in opacity as well as viscosity and may gel. About 0.1 milliliter of the composition is added to lysate. Typically, the pH of the composition is between 6 and 8, preferably, between 6.8 and 7.5. The mixture of composition and lysate is incubated for 1 hour undisturbed at 37° C. After incubation, the mixture is observed to determine if there was gelation of the mixture. Gelation indicates the presence of endotoxin. To determine the amount of endotoxin present in the composition, dilutions of a standardized solution of endotoxin are made and tested at the same time that the composition is tested. Standardized solutions of endotoxin are commercially available from, for instance, Sigma Chemical (Catalog No. 210-SE), U.S. Pharmacopeia (Rockville, MD, Catalog No. 235503), and Associates of Cape Cod, Inc., (Catalog No. E0005). In general, when a composition of the present invention is prepared by isolating proteins from a microbe, such as *P. multocida*, by a method as described herein (e.g., a method that includes disrupting and solubilizing the cells, and collecting the insoluble proteins), the amount of LPS in a composition of the present invention is less than the amount of LPS present in a mixture of same amount of the microbe that has been disrupted under the same conditions but not solubilized. Typically, the level of LPS in a composition of the present invention is decreased by, in increasing order of preference, at least 50%, at least 60%, at least 70%, at least 80%, or at least 90% relative to the level of LPS in a composition prepared by disrupting, but not solubilizing, the same microbe.

[0093] A composition described herein optionally further includes a pharmaceutically acceptable carrier. "Pharmaceutically acceptable" refers to a diluent, carrier, excipient, salt, etc., that is compatible with the other ingredients of the composition, and not deleterious to the recipient thereof. Typically, the composition includes a pharmaceutically acceptable carrier when the composition is used as described herein. Exemplary pharmaceutically acceptable carriers include buffer solutions and generally exclude blood products such as, for example, whole blood and/or plasma. The compositions as described herein may be formulated in pharmaceutical preparations in a variety of forms adapted to the chosen route of administration, including routes suitable for stimulating an immune response to an antigen. Thus, a

composition as described herein can be administered via known routes including, for example, oral; parenteral including intradermal, transcutaneous and subcutaneous, intramuscular, intravenous, intraperitoneal, etc. and topically, such as, intranasal, intrapulmonary, intramammary, intravaginal, intrauterine, intradermal, transcutaneous and rectally, etc. It is foreseen that a composition can be administered to a mucosal surface, such as by administration to the nasal or respiratory mucosa (e.g., via a spray or aerosol), in order to stimulate mucosal immunity, such as production of secretory IgA antibodies, throughout the animal's body.

[0094] A composition as described herein can also be administered via a sustained or delayed release implant. Implants suitable for use according to the invention are known and include, for example, those disclosed in International Publication No. WO 2001/037810 and/or International Publication No. WO 1996/001620. Implants can be produced at sizes small enough to be administered by aerosol or spray. Implants also can include nanospheres and microspheres.

[0095] A composition of the present invention is administered in an amount sufficient to provide an immunological response to proteins or whole cells described herein. The amount of protein present in a composition can vary. For instance, the dosage of protein can be between 0.01 micrograms ( $\mu\text{g}$ ) and 3000 milligrams (mg), typically between 100  $\mu\text{g}$  and 2000  $\mu\text{g}$ . When the composition is a whole cell preparation, the cells can be present at a concentration of 10<sup>6</sup> bacteria/ml, 10<sup>7</sup> bacteria/ml, 10<sup>8</sup> bacteria/ml, or 10<sup>9</sup> bacteria/ml. When a mixture of whole cells is administered (e.g., one population of cells expressing a subset of proteins and a second population expressing another subset of proteins) the ratio of populations can be 1:1. For an injectable composition (e.g., subcutaneous, intramuscular, etc.) the protein is preferably present in the composition in an amount such that the total volume of the composition administered is 0.5 ml to 5.0 ml, typically 1.0-3.0 ml. When the composition is a whole cell preparation, the cells are preferably present in the composition in an amount that the total volume of the composition administered is 0.5 ml to 5.0 ml, typically 1.0-2.0 ml. The amount administered will vary depending on various factors including, but not limited to, the specific proteins or cells chosen, the weight, physical condition and age of the animal, and the route of administration. Thus, the absolute weight of the protein or number of cells included in a given unit dosage form can vary, and depends upon factors such as the species, age, weight and physical condition of the animal, as well as the method of administration. Such factors can be determined by one skilled in the art. Other examples of dosages suitable for the invention are disclosed in Emery et al. (U.S. Pat. No. 6,027,736).

[0096] The formulations may be conveniently presented in unit dosage form and may be prepared by methods well known in the art of pharmacy. All methods of preparing a composition including a pharmaceutically acceptable carrier include the step of bringing the active compound (e.g., a protein or whole cell described herein) into association with a carrier that constitutes one or more accessory ingredients. In general, the formulations are prepared by uniformly and intimately bringing the active compound into association with a liquid carrier, a finely divided solid carrier, or both, and then, if necessary, shaping the product into the desired formulations.

[0097] A composition including a pharmaceutically acceptable carrier can also include an adjuvant. An "adjuvant" refers to an agent that can act in a nonspecific manner to enhance an immune response to a particular antigen, thus potentially reducing the quantity of antigen necessary in any given immunizing composition, and/or the frequency of injection necessary in order to generate an adequate immune response to the antigen of interest. Adjuvants may include, for example, IL-1, IL-2, emulsifiers, muramyl dipeptides, dimethyldioctadecylammonium bromide (DDA), avridine, aluminum hydroxide, oils, saponins, alpha-tocopherol, polysaccharides, emulsified paraffins (available from under the tradename EMULSIGEN from MVP Laboratories, Ralston, Nebraska), ISA-70, RIBI and other substances known in the art.

[0098] In another embodiment, a composition including a pharmaceutically acceptable carrier can include a biological response modifier, such as, for example, IL-2, IL-4 and/or IL-6, TNF, IFN-alpha, IFN-gamma, and other cytokines that effect immune cells. A composition can also include an antibiotic, preservative, anti-oxidant, chelating agent, etc. Such components are known in the art.

#### Methods of Making

[0099] This disclosure also provides methods for obtaining the proteins and whole cells described herein. Proteins and whole cell preparations described herein may be obtained by incubating a microbe, such as *P. multocida*, under conditions that promote expression of one or more of the proteins described herein. The proteins and whole cells as described herein may be isolatable from a microbe engineered to recombinantly express one or more of the proteins. In one embodiment, a *P. multocida* of serotype 2,5 or serotype 3,4 is used. In addition, such microbes are readily obtainable by techniques routine and known in the art. The microbes may be derived from an infected animal as a field isolate, and used to obtain the proteins and/or the whole cell preparations as described herein, or stored for future use, for example, in a frozen repository at from -20° C. to -95° C., or from -40° C. to -50° C., in bacteriological media containing 20% glycerol, and other like media.

[0100] The present invention also includes compositions prepared by the processes disclosed herein. Typically, such conditions are low metal conditions. As used herein, the phrase "low metal conditions" refers to an environment, typically a bacteriological medium that contains amounts of a free metal that cause a microbe to express a metal regulated protein at a detectable level. As used herein, the phrase "high metal conditions" refers to an environment that contains an amount of a free metal that causes a microbe to express a metal-regulated protein at a decreased level compared to expression of the metal-regulated protein under low metal conditions. In some cases, "high metal conditions" can refer to an environment that causes a cell to fail to express one or more of the metal-regulated proteins described herein at a detectable level. In some cases, "high metal conditions" can include a metal-rich natural environment and/or culture in a metal-rich medium without a metal chelator. In contrast, in some cases, "low metal conditions" can include culture in a medium that includes a metal chelator, as described in more detail below. Metals are those present in the periodic table under Groups 1 through 17 (IUPAC notation; also referred to as Groups I-A, II-A, III-B, IV-B, V-B, VI-B, VII-B, VIII, I-B, II-B, III-A, IV-A, V-A, VI-A, and VII-A,

respectively, under CAS notation). Preferably, metals are those in Groups 2 through 12, more preferably, Groups 3-12. Even more preferably, the metal is iron, zinc, copper, magnesium, nickel, cobalt, manganese, molybdenum, or selenium, most preferably, iron, copper, or zinc.

[0101] Low metal conditions are generally the result of the addition of a metal chelating compound to a bacteriological medium, the use of a bacteriological medium that contains low amounts of a metal, or a combination thereof. High metal conditions are generally present when a chelator is not present in the medium, when a metal is added to the medium, or a combination thereof. Examples of metal chelators include natural and synthetic compounds. Examples of natural compounds include plant phenolic compounds, such as flavonoids. Examples of flavonoids include the iron chelator myricetin. Examples of synthetic iron chelators include 2,2'-dipyridyl (also referred to in the art as 2,2'-bipyridyl), 8-hydroxyquinoline, ethylenediamine-di-O-hydroxyphenylacetic acid (EDDHA), desferrioxamine methanesulphonate (desferrioxamine), transferrin, lactoferrin, ovotransferrin, biological siderophores, such as the catecholates and hydroxamates, and citrate.

[0102] In one embodiment, 2,2'-dipyridyl is used for the chelation of iron. Typically, 2,2'-dipyridyl is added to the media at a concentration of at least 0.0025 micrograms/milliliter ( $\mu\text{g}/\text{ml}$ ), at least 0.025  $\mu\text{g}/\text{ml}$ , or at least 0.25  $\mu\text{g}/\text{ml}$ . High levels of 2,2'-dipyridyl can be 10  $\mu\text{g}/\text{ml}$ , 20  $\mu\text{g}/\text{ml}$ , or 30  $\mu\text{g}/\text{ml}$ .

[0103] Typically, TPEN is added to the media at a concentration of at least 25 micromolar ( $\mu\text{M}$ ), at least 50  $\mu\text{M}$ , or at least 70  $\mu\text{M}$ . In one embodiment, TPEN can be 70  $\mu\text{M}$  for expression of the polypeptide described herein, and higher levels may also be used.

[0104] It is expected that a *P. multocida* with a mutation in a fur gene will result in the constitutive expression of many, if not all, of the metal regulated proteins of the present invention. The production of a fur mutation in a *P. multocida* can be produced using routine methods including, for instance, electroporation and genetic constructs useful for gene knock-out in gram negative bacteria.

[0105] In one embodiment, the microbe, such as *P. multocida* used to make a composition described herein, e.g., a composition including isolated proteins or a composition including whole cells, may be produced using a *P. multocida* that has been engineered to recombinantly express one or more of the proteins described herein. In one embodiment, a *P. multocida* that expresses one or more of the proteins can be engineered to express the others. In one embodiment, such a *P. multocida* is incubated in the presence of low iron conditions, and the one or more recombinant proteins are expressed during the incubation in the low iron conditions. The result is a *P. multocida* that expresses iron-regulated proteins and the one or more recombinant proteins.

[0106] Many *Pasteurella* spp. are able to grow in low metal conditions in vitro in artificial media only after adaptation. For instance, a *Pasteurella* spp. can be adapted to low iron conditions in vitro by growth in the presence of low concentrations of an iron chelator and, after growth in a medium containing the chelator, gradually increasing the concentration of the chelator. For instance, a *Pasteurella* spp. can be adapted to growth in low iron conditions by adding 0.0025  $\mu\text{g}/\text{ml}$  of 2,2'-dipyridyl to a medium, and gradually increasing the concentration of the chelator to a greater concentration, for instance 25  $\mu\text{g}/\text{ml}$ .

[0107] The medium used to incubate the microbe is not critical, and conditions useful for the culture of *P. multocida* are known to the skilled person. The volume of medium used to incubate the microbe can vary. When a *P. multocida* microbe is being evaluated for the ability to produce the proteins described herein, the microbe can be grown in a suitable volume, for instance, 10 milliliters to 1 liter of medium. When a microbe is being grown to obtain proteins for use in, for instance, administration to animals, the microbe may be grown in a bioreactor to allow the isolation of larger amounts of proteins. Methods for growing microbes in a bioreactor are routine and known in the art. The conditions used for growing a microbe preferably include a metal chelator, more preferably an iron chelator, for instance 2,2'-dipyridyl, TPEN, or quercetin, a pH of between 6.5 and 7.5, preferably between 6.9 and 7.1, and a temperature of 37° C. When a fermentor is used, the culture may be purged with an appropriate gas to reduce dissolved oxygen content. Nitrogen is an example of such a gas. Dissolved oxygen may be regulated automatically or manually by agitation, the introduction of sterile air or pure oxygen to the culture.

[0108] In some aspects of the invention, a *P. multocida* may be harvested after growth. Harvesting includes concentrating the microbe into a smaller volume and suspending in a media different than the growth media. Methods for concentrating a microbe are routine and known in the art, and include, for example, filtration and/or centrifugation. Typically, the concentrated microbe is suspended in decreasing amounts of buffer. Preferably, the final buffer includes a metal chelator, preferably, ethylenediaminetetraacetic acid (EDTA). An example of a buffer that can be used contains Tris-base (7.3 grams/liter) and EDTA (0.9 grams/liter), at a pH of 8.5. Optionally, the final buffer also minimizes proteolytic degradation. This can be accomplished by having the final buffer at a pH of greater than 8.0, preferably, at least 8.5, and/or including one or more proteinase inhibitors (e.g., phenylmethanesulfonyl fluoride). Optionally and preferably, the concentrated microbe is frozen at -20° C. or below until disrupted. In one embodiment, bacterial cells may be concentrated into a pellet by, for instance, centrifugation, and the concentrated cells suspended in osmotic shock buffer (OMS; 7.26 grams/liter Tris-base and 0.93 grams/liter EDTA adjusted to a pH of 8.5). The ratio of cells to OMS may be 50 grams cell pellet, 60 grams cell pellet, or 70 grams cell pellet to 1 liter of OMS. The suspension of cells in OMS can be incubated at 2-8° C. for at least 24 hours, at least 48 hours, or at least 60 hours to remove excess endotoxin from the cells. In one embodiment, the incubation is for no greater than 72 hours. After the incubation the suspension is centrifuged again and the supernatant discarded to remove free endotoxin and any extracellular material, e.g., secreted proteins.

[0109] When the *P. multocida* is to be used as a whole cell preparation, the harvested cells may be processed using routine and known methods to inactivate the cells. Alternatively, when a *P. multocida* is to be used to prepare proteins of the present invention, the *P. multocida* may be disrupted using chemical, physical, or mechanical methods routine and known in the art, including, for example, french press, sonication, or homogenization. Preferably, homogenization is used. As used herein, "disruption" refers to the breaking up of the cell. Disruption of a microbe can be measured by methods that are routine and known in the art, including, for

instance, changes in optical density. Typically, a microbe is subjected to disruption until the percent transmittance is increased by 20% when a 1:100 dilution is measured. The temperature during disruption is typically kept at 4° C., to further minimize proteolytic degradation.

[0110] The disrupted microbe is solubilized in a detergent, for instance, an anionic, zwitterionic, nonionic, or cationic detergent. Preferably, the detergent is sarcosine, more preferably, sodium lauroyl sarcosinate. As used herein, the term "solubilize" refers to dissolving cellular materials (e.g., proteins, nucleic acids, carbohydrates) into the aqueous phase of the buffer in which the microbe was disrupted, and the formation of aggregates of insoluble cellular materials. The conditions for solubilization preferably result in the aggregation of proteins of the present invention into insoluble aggregates that are large enough to allow easy isolation by, for instance, centrifugation.

[0111] Preferably, the sarcosine is added such that the final ratio of sarcosine to gram weight of disrupted microbe is between 1.0 gram sarcosine per 4.5 grams pellet mass and 6.0 grams sarcosine per 4.5 grams pellet mass, preferably, 4.5 gram sarcosine per 4.5 grams pellet mass. The solubilization of the microbe may be measured by methods that are routine and known in the art, including, for instance, changes in optical density. Typically, the solubilization is allowed to occur for at least 24 hours, more preferably, at least 48 hours, most preferably, at least 60 hours. The temperature during disruption is typically kept low, preferably at 4° C.

[0112] The insoluble aggregates that include the proteins described herein may be isolated by methods that are routine and known in the art, such as centrifugation, filtration, or a combination thereof. In one embodiment, the insoluble aggregates are isolated by filtration, such as tangential or crossflow filtration. Examples of a molecular weight cutoff to use with tangential filtration are 100 kDa or 300 kDa. In one embodiment, a tangential filtration system has a pore size of 0.2 microns. Tangential filtration may aid in removal of residual sarcosine from the protein suspension. Tangential filtration results in concentration of the protein suspension. Thus, the insoluble aggregates can be isolated at a significantly lower cost.

[0113] In one embodiment, the sarcosine is removed from the isolated proteins. Methods for removing sarcosine from the isolated proteins are known in the art, and include, for instance, diafiltration, precipitation, hydrophobic chromatography, ion-exchange chromatography, affinity chromatography, and ultra-filtration and washing the proteins in alcohol, such as isopropyl alcohol, by diafiltration. After isolation, the proteins suspended in buffer and stored at low temperature, for instance, -20° C. or below. In those aspects of the present invention where a whole cell preparation is to be made, after growth the microbe can be killed with the addition of an agent such as glutaraldehyde, formalin, or formaldehyde, at a concentration sufficient to inactivate the cells in the culture. For instance, formalin can be added at a concentration of 0.3% (vol: vol). After a period of time sufficient to inactivate the cells, the cells can be harvested by, for instance, diafiltration and/or centrifugation, and washed.

[0114] In other aspects, an isolated protein of the invention may be prepared recombinantly. When prepared recombinantly, a polynucleotide encoding the protein may be identified and cloned into an appropriate expression host. The

recombinant expression host may be grown in an appropriate medium, disrupted, and the proteins isolated as described above.

#### Methods of Use

[0115] Also provided are methods of using the compositions described herein. The methods include administering to an animal an effective amount of a composition described herein. As used herein, an "effective amount" of a composition described herein is the amount able to elicit the desired response in the recipient. The composition can be administered at a time that maternal antibody may be present, for instance, as early as one day of age, or at a later time during the life of the animal. In one embodiment, the composition is administered before weaning, or at the time the animal is weaned (when the animal begins to be accustomed to food other than its mother's milk). The animal can be, for instance, fish, avian (including, for instance, chickens or turkeys), bovine (including, for instance, cattle), caprine (including, for instance, goats), ovine (including, for instance, sheep), porcine (including, for instance, swine), bison (including, for instance, buffalo), equine (including, for instance, horses), a companion animal (including, for instance, a dog or a cat), members of the family Cervidae (including, for instance, deer, elk, moose, caribou and reindeer), or a human. Examples of companion animals include dogs and cats. In one embodiment, an animal is a mouse. In one embodiment, an animal is a hooved animal.

[0116] The methods described herein refer to gram negative microbes. As used herein, a gram negative microbe includes, but is not limited to, members of the family Pasteurellaceae, such as *Pasteurella* spp. (including, for instance, *P. multocida* and *P. haemolytica*), *Photobacterium damsela* subsp., *piscicida* formerly known as *Pasteurella piscicida*, *Mannheimia* spp., and *Haemophilus* spp., members of the family Vibrionaceae (including, for instance, *Vibrio cholerae*), *Campylobacter* spp. (including, for instance, *C. jejuni*), members of the family Enterobacteriaceae (including, for instance, *Klebsiella* spp., *E. coli*, *Shigella* spp., *Salmonella* spp., *Proteus* spp., *Serratia* spp., and *Yersinia* spp.), and members of the family Pseudomonadaceae, preferably *Pseudomonas* spp., (including, for instance, *Pseudomonas aeruginosa*). Examples of *Klebsiella* spp. include *K. pneumoniae* and *K. oxytoca*. Examples of *Salmonella* spp. include *Salmonella enterica* serovars., *Bredeney*, *Dublin*, *Agona*, *Blockley*, *Enteriditis*, *Typhimurium*, *Hadar*, *Heidelberg*, *Montevideo*, *Muenster*, *Newport*, *senftenberg*, *Salmonella cholerasuis*, and *S. typhi*. Examples of strains of *P. multocida* include, for example, *P. multocida* serotypes 1 through 16 (serotype 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, or 16) or combinations thereof (for instance, 2,5, also referred to as 2x5 [both serotypes 2 and 5 are expressed by a *P. multocida*], or 3, 4 also referred to as 3x4 [both serotypes 3 and 4 are expressed by a *P. multocida*]). Examples of strains of *E. coli* include, for example, *E. coli* serotypes O1a, O2a, O78, and O157; different O:H serotypes including O104, O111, O26, O113, O91; hemolytic strains of enterotoxigenic *E. coli* such as K88<sup>+</sup>, F4<sup>+</sup>, F18ab<sup>+</sup>, and F18ac<sup>+</sup>; enteropathogenic (EPEC), enterohemorrhagic (EHEC), enteroinvasive (EIEC) and enteroaggregative (EAEC) strains of *E. coli*; and *E. coli* able to cause extra-intestinal infections, such as uropathogenic strains. In one embodiment, the gram negative microbe is a

pathogenic microbe. Respiratory pathogens such as *Bordetella* such species as *B. bronchiseptica*, *B. pertussis* and *B. parapertussis*, and *B. avium*.

[0117] In some embodiments, a method may further include additional administrations (e.g., one or more booster administrations) of the composition to the animal to enhance or stimulate a secondary immune response. A booster can be administered at a time after the first administration, for instance, one to eight weeks, preferably two to four weeks, after the first administration of the composition. Subsequent boosters can be administered one, two, three, four, or more times annually. Without intending to be limited by theory, it is expected that in some embodiments annual boosters will not be necessary, as an animal will be challenged in the field by exposure to microbes expressing proteins having epitopes that are structurally related to epitopes present on proteins of the composition administered to the animal.

[0118] In one embodiment, a method includes making antibody to a protein described herein, for instance by inducing the production of antibody in an animal, or by recombinant techniques. The antibody produced includes antibody that specifically binds at least one protein present in the composition. In this embodiment, an “effective amount” is an amount effective to result in the production of antibody in the animal. Methods for determining whether an animal has produced antibodies that specifically bind a protein present in a composition of described herein can be determined using routine methods. Also provided is antibody that specifically binds to a protein described herein, and compositions including such antibodies.

[0119] As used herein, an antibody that can “specifically bind” a protein is an antibody that interacts with the epitope of the antigen that induced the synthesis of the antibody, or interacts with a structurally related epitope. At least some of the epitopes present in the proteins described herein are epitopes that are conserved in the proteins of different species and different genera of microbes. Accordingly, antibody produced using a protein described herein is expected to bind to proteins expressed by more than one species of microbe, and provide broad spectrum protection against gram negative microbes.

[0120] In one embodiment, a method includes treating an infection in an animal, caused by a gram negative microbe. As used herein, the term “infection” refers to the presence of a gram negative microbe in an animal’s body, which may or may not be clinically apparent. Treating an infection can be prophylactic or, alternatively, can be initiated after the animal is infected by the microbe. Treatment that is prophylactic—e.g., initiated before a subject is infected by a microbe or while any infection remains subclinical—is referred to herein as treatment of a subject that is “at risk” of infection. As used herein, the term “at risk” refers to an animal that may or may not actually possess the described risk. Thus, typically, an animal “at risk” of infection by a microbe is an animal present in an area where animals have been identified as infected by the microbe and/or is likely to be exposed to the microbe even if the animal has not yet manifested any detectable indication of infection by the microbe and regardless of whether the animal may harbor a subclinical amount of the microbe. Accordingly, administration of a composition can be performed before, during, or after the animal has first contact with the microbe. Treatment initiated after the animal’s first contact with the microbe may result in decreasing the severity of symptoms and/or

clinical signs of infection by the microbe, completely removing the microbe, and/or decreasing the likelihood of experiencing a clinically evident infection compared to an animal to which the composition is not administered. The method includes administering an effective amount of a composition described herein to an animal having, or at risk of having, an infection caused by a gram negative microbe, and determining whether the number of microbes causing the infection has decreased. In this embodiment, an “effective amount” is an amount effective to reduce the number of the specified microbes in an animal or reduce the likelihood that the animal experiences a clinically-evident infection compared to an animal to which the composition is not administered. Methods for determining whether an infection is caused by a gram negative microbe are routine and known in the art, as are methods for determining whether the infection has decreased. The successful treatment of a gram negative microbial infection in an animal is disclosed in Examples 15-16, which demonstrates that a composition described herein made from two *P. multocida* strains, one serotype 2×5 and one serotype 3×4, protected chickens from challenge with a *P. multocida* of serotype 1.

[0121] In another embodiment, a method includes treating one or more symptoms or clinical signs of certain conditions in an animal that may be caused by infection by a gram negative microbe. The method includes administering an effective amount of a composition described herein to an animal having or at risk of having a condition, or exhibiting symptoms and/or clinical signs of a condition, and determining whether at least one symptom and/or clinical sign of the condition is changed, preferably, reduced. In one embodiment, the animal has a condition caused by a member of the family Pasteurellaceae, such as *Pasteurella* spp. (including, for instance, *P. multocida*, *P. haemolytica*, and *P. anatipesfizer*), *Mannheimia* spp., or *Haemophilus* spp. Examples of conditions include, but are not limited to, fowl cholera, new duck disease, Bovine Respiratory Disease Complex (also referred to as shipping fever pneumonia, or simply pneumonia), hemorrhagic septicemia in cattle, buffalo and bison in tropical and subtropical areas; pneumonia and atrophic rhinitis in swine; snuffles in rabbits; and mastitis and pneumonia in sheep. *Mannheimia haemolytica*, also known as *Pasteurella haemolytica*, can cause fibrinous pleuropneumonia involved in the shipping fever complex in cattle, septicemia in newborn and pneumonia and mastitis in adult sheep. *Haemophilus* spp. can cause Glasser’s disease in swine, fowl coryza in chickens, contagious equine metritis. *P. multocida* can cause fowl cholera which, in the peracute form, is one of the most virulent and highly infectious diseases of poultry. *P. anatipesfizer* can cause new duck disease (also known as duck septicaemia or infectious serositis).

[0122] Treatment of symptoms and/or clinical signs associated with conditions caused by infection by a gram negative microbe can be prophylactic or, alternatively, can be initiated after the development of a condition described herein. As used herein, the term “symptom” refers to subjective evidence of a disease or condition experienced by the patient and caused by infection by a microbe. As used herein, the term “clinical sign” or, simply, “sign” refers to objective evidence of disease or condition caused by infection by a microbe. Symptoms and/or clinical signs associated with conditions referred to herein and the evaluations of such symptoms are routine and known in the art. Treatment

that is prophylactic, for instance, initiated before a subject manifests symptoms or signs of a condition caused by a microbe, is referred to herein as treatment of a subject that is “at risk” of developing the condition. Thus, typically, an animal “at risk” of developing a condition is an animal present in an area where animals having the condition have been diagnosed and/or is likely to be exposed to a microbe causing the condition even if the animal has not yet manifested symptoms or signs of any condition caused by the microbe. Accordingly, administration of a composition can be performed before, during, or after the occurrence of the conditions described herein. Symptoms and/or clinical signs caused by a gram negative microbial infection are known to the person skilled in the art. Examples of symptoms and/or clinical signs include, but are not limited to, pneumonia, depression and toxemia, fever, serious to mucopurulent nasal discharge, moist cough, a rapid, shallow respiratory rate, abscesses and lesions in the lungs of cattle and swine; acute septicemia, joint infections and arthritis in poultry and wildfowl; turbinate atrophy in swine; superficial abscesses in cats; bite wound infections in humans, usually from dog or cat bites. Treatment initiated after the development of a condition may result in decreasing the severity of the symptoms or signs of one of the conditions, or completely removing the symptoms or signs. In this embodiment, an “effective amount” is an amount effective to prevent the manifestation of symptoms or signs of a disease, decrease the severity of the symptoms or signs of a disease, and/or completely remove the symptoms or signs.

[0123] Also provided is a method for decreasing colonization by a gram negative microbe, for instance blocking the attachment sites of a gram negative microbe, including tissues of the skeletal system (for instance, bones, cartilage, tendons and ligaments), muscular system, (for instance, skeletal and smooth muscles), circulatory system (for instance, heart, blood vessels, capillaries and blood), nervous system (for instance, brain, spinal cord, and peripheral nerves), respiratory system (for instance, nose, trachea lungs, bronchi, bronchioles, alveoli), digestive system (for instance, mouth, salivary glands, esophagus, liver, stomach, large and small intestine), excretory system (for instance, kidney, ureter, bladder, and urethra), endocrine system (for instance, hypothalamus, pituitary, thyroid, pancreas and adrenal glands), reproductive system (for instance, ovaries, oviduct, uterus, vagina, mammary glands, testes, and seminal vesicles), lymphatic/immune systems (for instance, lymph, lymph nodes and vessels, mononuclear or white blood cells, such as macrophages, neutrophils, monocytes, eosinophils, basophils, and lymphocytes, including T cells and B cells), and specific cell lineages (for instance, precursor cells, epithelial cells, stem cells), and the like. Decreasing colonization in an animal may be performed prophylactically or, alternatively, can be initiated after the animal is colonized by the microbe. Treatment that is prophylactic—e.g., initiated before a subject is colonized by a microbe or while any colonization remains undetected—is referred to herein as treatment of a subject that is “at risk” of colonization by the microbe. Thus, typically, an animal “at risk” of colonization by a microbe is an animal present in an area where animals have been identified as colonized by the microbe and/or is likely to be exposed to the microbe even if the animal has not yet manifested any detectable indication of colonization by the microbe and regardless of whether the animal may harbor a sub-colonization number

of the microbe. Accordingly, administration of a composition can be performed before, during, or after the animal has first contact with the microbe. Treatment initiated after the animal’s first contact with the microbe may result in decreasing the extent of colonization by the microbe, completely removing the microbe, and/or decreasing the likelihood that the animal becomes colonized by the microbe compared to an animal to which the composition is not administered. Thus, the method includes administering an effective amount of a composition described herein to an animal colonized by, or at risk of being colonized by, a gram negative microbe. In this embodiment, an “effective amount” is an amount sufficient to decrease colonization of the animal by the microbe, where decreasing colonization refers to one or more of: decreasing the extent of colonization by the microbe, completely removing the microbe, and/or decreasing the likelihood that the animal becomes colonized by the microbe compared to an animal to which the composition is not administered. Methods for evaluating the colonization of an animal by a microbe are routine and known in the art. For instance, colonization of an animal’s respiratory tract by a microbe can be determined by measuring the presence of the microbe in the animal’s specimens from the lower respiratory tract by tracheal swab, transtracheal wash, or bronchoalveolar lavage. It is expected that decreasing the colonization of an animal by a microbe will reduce transmission of the microbe to other animals of the same or different species.

[0124] Also provided is the use of such antibody to target a microbe expressing a protein having an epitope structurally related to an epitope present on a protein described herein. A composition described herein can be used to provide for active or passive immunization against bacterial infection. Generally, the composition can be administered to an animal to provide active immunization. However, the composition can also be used to induce production of immune products, such as antibodies, which can be collected from the producing animal and administered to another animal to provide passive immunity. Immune components, such as antibodies, can be collected to prepare compositions (preferably containing antibody) from serum, plasma, blood, colostrum, etc. for passive immunization therapies. Antibody compositions including monoclonal antibodies and/or anti-idiotypes can also be prepared using known methods. Chimeric antibodies include human-derived constant regions of both heavy and light chains and murine-derived variable regions that are antigen-specific (Morrison et al., Proc. Natl. Acad. Sci. USA, 1984, 81(21):6851-5; LoBuglio et al., Proc. Natl. Acad. Sci. USA, 1989, 86(11):4220-4; Boulianne et al., Nature, 1984, 312(5995):643-6.). Humanized antibodies substitute the murine constant and framework (FR) (of the variable region) with the human counterparts (Jones et al., Nature, 1986, 321(6069):522-5; Riechmann et al., Nature, 1988, 332(6162):323-7; Verhoeyen et al., Science, 1988, 239(4847):1534-6; Queen et al., Proc. Natl. Acad. Sci. USA, 1989, 86(24):10029-33; Daugherty et al., Nucleic Acids Res., 1991, 19(9):2471-6.). Alternatively, certain mouse strains can be used that have been genetically engineered to produce antibodies that are almost completely of human origin; following immunization the B cells of these mice are harvested and immortalized for the production of human monoclonal antibodies (Bruggeman and Taussig, Curr. Opin. Biotechnol., 1997, 8(4):455-8; Lonberg and Huszar, Int. Rev. Immunol., 1995; 13(1):65-93;

Lonberg et al., *Nature*, 1994, 368:856-9; Taylor et al., *Nucleic Acids Res.*, 1992, 20:6287-95.). Passive antibody compositions and fragments thereof, e.g., scFv, Fab, F(ab')<sub>2</sub> or Fv or other modified forms thereof, may be administered to a recipient in the form of serum, plasma, blood, colostrum, and the like. However, the antibodies may also be isolated from serum, plasma, blood, colostrum, and the like, using known methods for later use in a concentrated or reconstituted form such as, for instance, lavage solutions, impregnated dressings and/or topical agents and the like. Passive immunization preparations may be particularly advantageous for the treatment of acute systemic illness, or passive immunization of young animals that failed to receive adequate levels of passive immunity through maternal colostrum. Antibodies useful for passive immunization may also be useful to conjugate to various drugs or antibiotics that could be directly targeted to bacteria expressing during a systemic or localized infection a protein having an epitope structurally related to an epitope present on a protein described herein.

[0125] Animal models are available for experimentally evaluating the compositions described herein. These models are commonly accepted models for the study of disease caused by gram negative microbes. In those cases where a gram negative microbe causes disease in an animal, for instance a cow or a chicken, the natural host can be used to experimentally evaluate the compositions described herein.

[0126] However, protection in an animal model is not the only way to assess whether a composition can confer protection to an animal against infection by a gram negative microbe. The adaptive immune response consists of two primary divisions: the humoral (antibody) response and the cellular (T cell) response. Following infection by a bacterial pathogen, dendritic cells at the infection site encounter microbial antigens and produce signaling molecules such as, for example, surface receptors and cytokines in response to conserved molecular patterns associated with the specific bacterium. These signals are shaped by the nature of the pathogen and ideally lead to the appropriate antibody and T cell responses that protect the host from disease. While some bacterial diseases are controlled primarily through antibody functions, others require T cell responses or both antibody and T cell responses for protection. The goal of vaccine biology is to identify the immune responses that provide protection and then design a vaccine to reproduce one or more of these responses in humans.

[0127] Antibodies can have many different functions in conferring protection against infection such as, for example, complement fixation, opsonization, neutralization, and/or agglutination. Moreover, some subclasses of antibodies are better than others at specific functions; for example, or complement fixation the following hierarchy exists for human IgG subclasses: IgG3>IgG1>IgG2>IgG4.

[0128] Antibody immunological functions can be studied in a variety of ways. For instance, Western blots are used to identify antigen-specific binding based on size of separated proteins, while the standard enzyme-linked immunosorbant assay (ELISA) is used to produce quantitative information about antibody titers within serum. Antibody surface binding studies are used to determine whether antibody in serum are able to recognize antigens on the surface of intact bacteria, an important indicator of whether the antibodies have the potential to work in vivo. Thus, one skilled in the art recognizes that antibody binding assays such as a West-

ern blot, ELISA (e.g., using human antisera), and/or surface binding correlate positively with the specifically-bound antigens providing immunological activity against microbial infection. However, one skilled in the art further recognizes that a lack of antibody binding in an assay such as, for example, a Western blot, ELISA, or surface binding assay does not mean that the assayed antigen fails to provide immunological activity against microbial infection.

[0129] Antibodies can mediate bacterial death by blocking the acquisition of nutrients or initiating complement-mediated membrane perforation that leads to osmotic lysis. Bactericidal antibodies can be assayed by mixing serum with live cultures and measuring for the presence of viable bacteria under appropriate conditions known to those skilled in the art. Techniques such as opsonophagocytosis assays (OPA), in which antibody and complement-bound bacteria are combined with human or mouse phagocytes to determine levels of bacterial killing, are useful for studying antibody function. A similar oxidative burst assay can be used to assess the level of reactive oxygen species (ROS) by fresh human or mouse neutrophils following interaction with antibody and complement-bound bacteria.

[0130] In some cases, one can determine that a protein described herein possesses cell-mediated immunological activity against a gram negative microbe and, therefore, the protein may exhibit immunological activity in the absence of inducing the production of antibodies. Cytotoxic or CD8 T cells primarily kill infected cells directly through various effector mechanisms, while helper CD4 T cells function to provide important signaling in the way of cytokines. These T cell classes can be further subdivided based on the cytokines they produce, and different subclasses are effective against different bacterial pathogens. T cells are often studied by assessing their phenotypes with flow cytometry, where antibodies are used to visualize the levels of specific surface markers that enable classification of the T cells as, for example, a recently activated CD4 T cell, a memory CD8 T cell, etc. In addition, cytokines and other products of T cells can be studied by isolating the T cells from lymphoid tissue and restimulating them with cognate antigen. Following antigen stimulation the T cells produce cytokines that may be visualized by, for example, intracellular cytokine staining coupled with flow cytometry, or collecting the cell supernatants and using Luminex bead technology to measure 15-25 cytokines simultaneously.

[0131] Thus, in addition to animal models, those of ordinary skill in the art recognize that immunological activity commensurate with the methods described herein may correlate with any one or more of the following: Western blot data showing that serum from animals exposed to a microbial pathogen contains antibody that specifically binds to a protein described herein, Western blot data showing that serum from animals exposed to protein described herein contains antibody that specifically binds to a gram negative microbe, cell surface binding assays demonstrating that antibody that specifically binds to a protein described herein specifically binds to a gram negative microbe, opsonophagocytosis data, and cytokine induction.

[0132] Also provided is a method for detecting antibody that specifically binds proteins described herein. These methods are useful in, for instance, detecting whether an animal has antibody that specifically binds a protein described herein, and diagnosing whether an animal may have a condition caused by a microbe expressing proteins

that share epitopes with the proteins described herein. Such diagnostic systems may be in kit form. The methods include contacting an antibody with a preparation that includes a protein described herein to result in a mixture. The antibody may be present in a biological sample, for instance, blood, milk, or colostrum. The method further includes incubating the mixture under conditions to allow the antibody to specifically bind the protein to form a protein: antibody complex. As used herein, the term "protein: antibody complex" refers to the complex that results when an antibody specifically binds to a protein. The preparation that includes the proteins described herein may also include reagents, for instance a buffer, that provide conditions appropriate for the formation of the protein: antibody complex. The protein: antibody complex is then detected. The detection of antibodies is known in the art and can include, for instance, immunofluorescence or peroxidase. The methods for detecting the presence of antibodies that specifically bind to proteins described herein can be used in various formats that have been used to detect antibody, including radioimmunoassay and enzyme-linked immunosorbent assay.

#### Kits

[0133] Also provided are kits. In one embodiment, a kit is for detecting antibody that specifically binds a protein described herein. The antibody detected may be obtained from an animal suspected of having an infection caused by a gram negative microbe. In another embodiment, a kit is for detecting a protein described herein. In yet another embodiment, a kit is for using a protein described herein, such as using a protein to produce antibody, treat a condition, or treat an infection.

[0134] The kit includes at least one of the proteins described herein (e.g., one, at least two, at least three, etc.), or an antibody described herein in a suitable packaging material in an amount sufficient for at least one assay or use. Optionally, other reagents such as buffers and solutions are also included. For instance, a kit may also include a reagent to permit detection of an antibody that specifically binds to a protein described herein, such as a detectably labeled secondary antibody designed to specifically bind to an antibody obtained from an animal. Instructions for use of the packaged antibody or protein are also typically included. As used herein, the phrase "packaging material" refers to one or more physical structures used to house the contents of the kit. The packaging material is constructed by routine methods, generally to provide a sterile, contaminant-free environment. The packaging material may have a label which indicates that the proteins can be used for detecting antibody that specifically binds a protein described herein, or using a protein described herein. In addition, the packaging material contains instructions indicating how the materials within the kit are employed to detect the antibody or administer a protein to an animal. As used herein, the term "package" refers to a container such as glass, plastic, paper, foil, and the like, capable of holding within fixed limits the proteins, and other reagents, for instance a secondary antibody. Thus, for example, a package can be a microtiter plate well to which microgram quantities of proteins have been affixed. A package can also contain a secondary antibody. "Instructions for use" typically include a tangible expression describing the reagent concentration or at least one assay method parameter, such as the relative amounts of reagent and sample to

be admixed, maintenance time periods for reagent/sample admixtures, temperature, buffer conditions, and the like.

[0135] The present invention is illustrated by the following examples. It is to be understood that the particular examples, materials, amounts, and procedures are to be interpreted broadly in accordance with the scope and spirit of the invention as set forth herein.

#### Example 1

##### Selection and Preparation of Type Strains

[0136] *Pasteurella multocida* strains X-73, P-1059 and P-1662 are reference strains for serotype A: 1, A: 3 and A: 4, respectively. Serotypes A: 1, A: 3, and A: 4 are also referred to as serotypes 1, 3, and 4, respectively. Each of these strains was obtained from the USDA National Veterinary Services Laboratory (NVSL) (Ames, IA) as a lyophilized culture. The cultures were resuspended in sterile saline, streaked for isolation using a sterile inoculating loop onto trypticase soy agar (TSA)+5% sheep blood (Becton Dickinson, Sparks MD), and incubated overnight at 37° C. Several isolated colonies were picked from the plate with a sterile inoculating loop and transferred to a cryobank tube (Copan diagnostics, Murrietta CA) containing polystyrene beads for the purpose of cryopreservation. The cryobank tubes were then stored at <-60° C.

#### Example 2

##### Preparation of *Pasteurella Multocida* Isolate

[0137] A *Pasteurella multocida*, serotype 3x4 was originally isolated from the liver and lungs of a 44 week old turkey breeder hen under natural field conditions showing clinical signs of Pasteurellosis. The isolate was plated on a tryptic soy agar (TSA) plus 5% sheep blood. After initial isolation, the organism was passed 5 times in non-animal-based Trypticase Soy Broth (naTSB) (EM Science, Darmstadt, Germany) (supplemented with 18 µg/ml 2,2' Dipyridyl (DP) (Sigma Aldrich, St. Louis, MO) and 6 g/l Yeast Extract (Becton Dickinson, Sparks, MD). Briefly, colonies were picked and inoculated into 500 ml naTSB and the culture was incubated for 16 hours and 25 minutes at 37° C. on a table-top shaker set at 105 revolutions per minute (RPM). The following morning, 25 ml of the overnight culture was transferred to 500 ml fresh naTSB and incubated for 8 hours at the same temperature and agitation. Five ml of the culture was transferred to 500 ml fresh naTSB and incubated for 15 hours in the same conditions as above. Fifty ml of this culture was transferred to 500 ml fresh naTSB and incubated in the same manner for 4 hours. Four bottles containing 900 ml naTSB each were each inoculated with 100 ml of the previous media and incubated for 3 hours and 45 minutes. The four liters of culture was aseptically dispensed into sterile 500 ml Nalgene jars and centrifuged in a Beckman model J2-21 centrifuge (Beckman Coulter, Brea CA) at 3500 rpm or approximately 1350×g for 20 minutes to pellet the bacterial cells. The supernatant was aseptically removed by aspiration and the cell pellets were resuspended in a total of 500 ml of naTSB containing 20% glycerol as a cryopreservative. The cell suspension was then dispensed into 243 vials at 2 ml per vial and frozen at <-60° C. This frozen Master Seed was designated MS061130.

[0138] A working seed was prepared from the above frozen Master Seed. Briefly, 1-25 µl of the frozen seed was

inoculated into 100 ml modified tryptose phosphate broth (mTPB) containing 6 g/l yeast extract. The culture was incubated for 13 hours at 37° C. on a table-top shaker until the culture reached an optical density (O.D.) of 0.6 to 0.8 on a Spectronic 20D spectrophotometer (Thermo Fisher Scientific, Mississauga, Ontario, Canada) set at 540 or 630 nm. The culture was transferred, 10 ml into 90 ml mTPB with 18 µg/ml 2, 2' dipyridyl and incubated for 2-3 hours at 37° C. with agitation. The culture was transferred and grown in the same manner two additional times as above. When the culture reached an O.D. of 0.8, the cells were centrifuged at approx. 3500 rpm for 20 minutes, the supernatant was aseptically removed by aspiration, and the culture was resuspended in the original volume of fresh mTPB. Sterile glycerol was added to reach 20% final concentration. The culture was mixed well, dispensed into sterile cryogenic vials at 2 ml per vial, and stored frozen at <-60° C.

#### Example 3

##### Production of Metal Regulated Proteins

[0139] *Pasteurella multocida* Type strains (X-73, P-1059, and P-1662) and the vaccine candidate strain (MS061130) of example 2 were grown in iron limiting and iron replete conditions in order to show differential membrane protein expression.

[0140] Each of the frozen type strain cultures from example 1 and the working seed isolate from example 2 were inoculated into 10 ml of porcine brain heart infusion broth (BD-Difco, Sparks MD)+6 g yeast extract (pBHI) and incubated overnight at 37 C with agitation. Two of the cultures did not grow directly in broth and so were first inoculated onto TSA+blood agar plates and incubated overnight at 37 C prior to transferring colonies to pBHI. Five ml of each overnight culture was transferred to 100 ml fresh pBHI containing either 20 µg/ml FeCl<sub>3</sub> or DP for iron-replete or iron-deplete conditions, respectively. The cultures were incubated at 37° C. for 7 hours and 30 minutes with agitation on a table top shaker (Barnstead, Dubuque IA). Bottles displayed turbidity indicating strong growth after incubation, and the cultures were then transferred, 50 ml into 500 ml of fresh pBHI containing either FeCl<sub>3</sub> or DP and incubated for 15 hours at 37° C., with agitation.

[0141] The cultures were processed to isolate outer membrane proteins. Briefly, the cultures were centrifuged for 20 minutes at approximately 11,000×g in a Beckman floor model centrifuge equipped with a JA-10 rotor (Beckman Coulter, Brea CA). The supernatant was decanted and discarded. The cell pellet was resuspended in 35 ml of Osmotic Shock Buffer (OMS) containing 7.26 grams/liter Tris-base and 0.93 grams/liter EDTA adjusted to a pH of 8.5, and the cell suspension was frozen at -80° C. for 3 hours to help weaken the cell wall structure. The cell suspension was then thawed and sonicated (Branson sonifier, model 102c) (Branson Ultrasonics, Danbury, CT) for 90 seconds in an ice water bath to prevent excessive heating of the disrupted cell suspension. Cell debris was removed by centrifugation at approximately 37,000×g for 20 minutes. Sodium lauroyl sarcosyl was added to the retained supernatant to reach a concentration of 3% v/v and the protein suspension was allowed to solubilize for 18-24 hours at 4° C. with slight agitation. Insoluble membrane proteins were collected by centrifugation at 37,000×g for at least two hours. The pellet was resuspended in tris buffered water and the resulting

antigen was measured for protein content via the bicinchoninic acid (BCA) assay (Pierce BCA protein assay, ThermoFisher Scientific, Waltham MA) according to the manufacturer instructions.

#### Example 4

##### Test for Metal Regulatged Protein Pattern Coverage by SDS-Page and Analysis of Protein Banding

[0142] Cell extracts, derived from each isolate grown under iron restriction, were size-fractionated on SDS-PAGE gels using a 4% stacking gel and 10% resolving gel. Samples for electrophoresis were prepared by combining 30 µg of sample with 30 µl of SDS reducing sample buffer (62.5 mM Tris-HCL pH 6.8, 20% glycerol, 2% SDS, 5% beta-mercaptoethanol) boiled for 4 minutes. A sample of each extract was resolved on a 10% SDS-PAGE gel per standard methods and visualized by Coomassie Blue staining (FIG. 1). The vaccine candidate strain (MS061130) was compared to the type strains on SDS-PAGE gels. Specific attention was given to the banding patterns in the regions between 66 and 116 kDa. The vaccine candidate strain demonstrated bands overlapping with type strains, as shown in FIG. 1, and was considered a strong candidate for further protection studies.

#### Example 5

##### Production of Metal Regulated Proteins

[0143] Two separate antigen compositions were prepared at production scale using a single Master Seed MS061130 strain of *Pasteurella multocida* designated as antigen A and antigen B. This was done to evaluate the consistency of the manufacturing process.

##### Fermentation

[0144] A cryogenic vial of the working seed of example 2 was thawed at 30-35° C. for 10-15 minutes and 0.1 ml of the thawed cell suspension was added to 250 ml of 37° C. mTPB. The culture was incubated for 6.5 hours at 37° C. on a tabletop shaker set at 70 rpm. After incubation, 25 ml of the culture was transferred to 300 ml of mTPB plus 18 µg/ml DP to restrict iron. This second culture was incubated for approximately 2 hours at 37° C. shaking at 70 rpm. The culture was transferred 250 ml into 3 liters of the above media, and the culture was incubated for approximately 3 hours at 37 C shaking at 70 rpm until an OD of 1 was reached (as measured at 540 nm). The entire culture was used to inoculate a 130 liter fermentor (Bio-Service, Easton, PA) charged with 70 liters of mTPB+18 µg/ml DP and 0.5 ml/l antifoam (MAZU DF 204 Chem/Serv, Minneapolis, MN). The parameters of the fermentation were as follows: Dissolved oxygen (DO) was maintained at 20% (target 15-30%) by automatic agitation control and airflow sparge set at 60 liters per minute (LPM) and 5 pounds per square inch (PSI) back pressure. The culture was incubated with agitation controlled automatically by controlling dissolved oxygen at 20%. The pH was controlled automatically between 6.9 and 7.1 by the automatic titration of 50% sodium hydroxide or 10% hydrochloric acid. The temperature was maintained at 37 C. The culture was allowed to grow in this manner for approximately 5 hours until an optical density of 4.21 at 540 nm was reached as measured by a Beckman DU600 spectrophotometer (Beckman Instru-

ments, Fullerton, CA). The entire expansion culture was transferred to a 1200 liter fermentation vessel (New Brunswick Scientific, Edison, NJ) charged with 989 l of mTPB plus DP and antifoam as listed above. The fermentation conditions were as follows: Airflow was set to 300 LPM and back pressure set at 5 PSI. Agitation was initially set to 100 RPM and then controlled by dissolved oxygen set point of 20% with a range of 15-30%. The culture was allowed to grow for another 5 hours until growth plateaued, as measured by optical density measurements being steady at approximately 2.5 at 540 nm at a 1:100 dilution for 1.5 hours. Growth was suspended by adjusting the culture pH to 8.7 and the temperature to 23° C. until harvest.

#### Harvest

The Bacterial Cells were Collected by Tangential Flow Filtration.

[0145] The bacterial fermentation was concentrated and washed using a Pall Filtron Tangential Flow Maxisette-25 (Pall Filtron Corporation, Northboro, MA) equipped with two 30 ft<sup>2</sup> Alpha 300-K open channel filters, catalog No. AS300C5, (Pall Filtron) connected to a Waukesha Model U-60 feed pump (Waukesha Chemy-Burrell, Delevan, Wis.) set at about 44 Hz. The original culture volume of 1078 liters was reduced to approximately 100 liters into a process tank (Lee Industries, model 2000LBDT) using a filter inlet pressure of 30 psi and a retentate pressure of 12-14 psi. The retentate (100 liters) was adjusted to 200 liters using sterile Osmotic Shock Buffer (OMS) containing 7.26 grams/liter Tris-base and 0.93 grams/liter EDTA adjusted to a pH of 8.5. The EDTA in the OMS serves to remove LPS from the cell wall, while the elevated pH prevents much of the proteolytic degradation after freezing and disruption. Protease inhibitors may be used instead, or in addition to, an elevated pH. The retentate was concentrated back down to approximately 65 liters and the system was flushed with 20 liters of OMS. The retentate was mixed thoroughly while in the 200-liter tank using a bottom mount magnetically driven mixer.

[0146] The retentate was steriley dispensed (3.5 liters) into sterile 4 liter NALGENE containers No. 2122 in a biosafety cabinet and placed into a -20° C. freezer for storage. Freezing the bacterial pellet serves to weaken the cell wall structure making downstream disruption more efficient. The pellet mass was calculated by centrifuging 30 ml samples of the fermented culture and final harvest. Briefly, pre-weighted 50 ml NALGENE conical tubes were centrifuged at 39,000 $\times g$  for 90 minutes in a BECKMAN J2-21 centrifuge using a JA-21 rotor (Beckman Instruments, Palo Alto Calif.). At the end of the run, the supernatant was poured off and the tubes were weighed again. The pellet mass was calculated for each stage. The fermentation process yielded a wet pellet mass of 3.7 kilograms.

[0147] Alternative methods for bacterial harvest can be used. Bacterial harvest may be performed by the use of hollow fiber filter methods. Bacterial culture is harvested using filter cartridges ranging in size from 0.6  $\mu$ M to 750 kDa; preferably with a 750 kDa cartridge. Culture is reduced in volume from 2-20 $\times$  and subsequently washed 1-5 $\times$  by diafiltration with buffer prior to storage at 40 C or freezing at -20° C. In this manner, undesired media proteins, bacterial proteins and LPS are removed from the culture. In another alternative, bacterial harvest may be performed by the use of industrial scale centrifugation, for example, by use of a disc-stack centrifuge.

#### Disruption

[0148] Sixty five Liters of frozen bacterial cell slurry in OMS were thawed at 4° C. (3.7 kg of pellet mass). The liquid culture suspension from each container was aseptically aspirated into a steam in place 150 liter jacketed process tank (Lee, Model Style U). The cell suspension was disrupted by homogenization. Briefly, the 150 liter tank containing the bacterial suspension was connected to an Avestin model EF-C500B Homogenizer (Avestin, Inc. Ottawa, ON, Canada). A 250 liter jacketed process tank (empty) (Lee, Model 259LU) with a top mounted mixer (Eastern, Model TME-1/2, EMI Incorporated, Clinton, CT) was connected to the homogenizer such that the fluid in the process tank could be passed through the homogenizer, into the empty tank and back again, allowing for multiple homogenizing passes while still maintaining a closed system. The temperature during homogenization was kept at 4° C. At the start of the pass, fluid was circulated at 60 psi via a Waukesha model 10DO pump (Waukesha) through the homogenizer (500 gallons/hour) and back to the tank of origin, while the homogenizer pressure was adjusted to 15,000 psi. Prior to homogenization, two pre-homogenizing samples were withdrawn from the homogenizer to establish a baseline for determining the degree of disruption and monitoring of pH. The degree of disruption was monitored by transmittance (% T at 540 nm at 1:100 dilution) compared to the non-homogenized sample. The starting % T was 86.65 at a 1:100 dilution. The cell slurry was passed once through the homogenizer, and the resulting % T was 89.9 at a 1:100 dilution.

[0149] After homogenization, 1 liter Sodium Lauroyl Sarcosinate (Hamposyl L-30, Chem Serv, Minneapolis, MN) was aseptically passed to the 81 liters of homogenized slurry for solubilization. The jacketed vessel removed from the homogenizer and was kept on a chiller loop at 5 C mixing at 30% mixer speed. After 29 hours, 121 ml of formaldehyde was added as a preservative, followed by a 2 liter wash of OMS. Solubilization continued for a total of 46 hours and 16 minutes. This time period was important for complete solubilization. It was discovered that increasing the solubilization time in OMS at an elevated pH (8.0-8.5) that metal regulated proteins aggregated together forming large insoluble aggregates that were easily removed by centrifugation.

#### Protein Harvest

[0150] The aggregated metal regulated proteins within the solubilized process fluid were collected by centrifugation using T-1 SHARPLES, (Alfa Laval Separations, Warminster, Pa.). Briefly, the tank of solubilized homogenate was fed into twelve Sharples with a feed rate of 170 ml/minute at 11 psi. The first pass protein collected on the bowls of the Sharples was discarded. This consists of large cell debris and cell wall material. The effluent was collected into a second 250 liter jacketed process tank through a closed sterile loop allowing for multiple passes through the centrifuges while maintaining a closed system. The temperature during centrifugation was kept at 4° C. The solubilized homogenate was passed an additional 7 times across the centrifuges, at a feed rate of 150 ml/minute and a pressure of 21 PSI. Protein was collected after each pass. The protein was collected, resuspended and dispensed in 7.46 liters Tris-buffer pH 8.5.

Twenty five ml of formaldehyde (Sigma) was added, to reach 0.3% concentration, as a preservative.

[0151] Alternative methods for protein harvest can be used. For example, the desired proteins can be harvested by the use of hollow fiber filter methods. Proteins can be harvested using filter cartridges ranging in size from 5 kDa to 0.2  $\mu$ M; preferably with a 50 kDa to 750 kDa cartridge. Culture is reduced in volume from 2-20x and subsequently washed 1-5x by diafiltration with buffer prior to storage at 40 °C or freezing at -20° C.

#### Diafiltration

[0152] The protein suspension was washed by diafiltration at 4° C. to remove any contaminating sarcosine that may have been bound to the protein. Briefly, the 8640 ml of protein was sterilely aspirated into a 200 liter process tank containing 50 liters sterile Tris-buffer, pH 8.5, equipped with a bottom mount Dayton mixer, Model 2Z846 (Dayton Electric, Chicago, Ill.) rotating at 125 rev/minute. The process tank was steriley connected to a MILLIPORE PELLICON TANGENTIAL FLOW FILTER assembly (Millipore Corporation), equipped with a 25 ft<sup>2</sup> screen-channel series Alpha 10K Centrasette filter (Pall Filtron) connected to a Waukesha Model U30 feed pump. The 59 liter protein solution was brought to 150 liters by adding TBW, concentrated down to 35 liters, brought up to a volume of 110 liters of TBW and then concentrated down to 10.4 L final antigen volume. The protein concentrate was aseptically dispensed (3.5 liters) into sterile 4 liter NALGENE containers and placed into a -20° C. freezer for storage.

[0153] This process produced a composition containing metal regulated proteins with a decrease in the amount of LPS and very little to no sarcosine residue. The protein was examined by SDS-PAGE for purity and banding profile, and also examined for bacterial contamination, residual sarcosine and LPS. The banding profile of the finished product showed consistent patterns as examined by electrophoresis. The composition was tested for sarcosine by the use of a modified agar gel diffusion test in which sheep red blood cells (5%) were incorporated into an agar base (1.5%). Wells were cut into the agar and samples of the finished product along with control samples of known concentrations of sarcosine at 0.05, 0.1, 0.2, 0.3, 0.4, 0.5, 1.0 and 2.0% were placed into the wells. The gel was incubated at 25° C. for 24 hours and the degree of hemolysis was determined compared to the controls. The process removes the level of detectable sarcosine below 0.05%, which at this concentration showed minimal hemolysis in control samples. The concentration of LPS is examined by a Limulus amebocyte lysate (LAL) test available under the tradename PYRO-TELL (Associates of Cape Cod, Inc., East Falmouth, Mass.).

#### Example 6

##### Preparation of Immunizing Compositions Derived from *Pasteurella Multocida*

[0154] Three vaccine serials consisting of the same antigen was prepared from the Master Seed MS061130 of Example 5. Please Note: two antigen preparation representing two separate fermentation processes were used to prepare two antigen lots designated as antigen A and antigen B. Thus, The first vaccine serial was prepared using a mixture of antigen A and antigen B, the second vaccine serial was

prepared only using antigen A, and the third vaccine serial was prepared using only antigen B.

[0155] The vaccine compositions consisting of antigen A, B and A+B were prepared in the following constituents: 44.44% aqueous protein suspension in 0.1% formalized saline, (standardized to 150  $\mu$ g protein per chicken dose) 50% Drakeol 6 mineral oil (VOPAK USA, Inc, Kirkland, WA), 3.0% Span 85 and 2.56% Tween 85 (Ruger Chemicals, Hillside, NJ). Briefly the Mineral Oil and Span 85 were combined and dispensed into a vessel equipped with a high-speed emulsifier (IKA model Process pilot 2000/4 or equivalent). Antigen, Tween 85 and 0.1% formalized saline was dispensed into a second vessel. The emulsifier was set at 60 hz, and the aqueous solution was pumped into the oil, which was pre-cooled to 7° C. The vaccine was continuously stirred as it was pumped into sterile bottles.

#### Example 7

##### Vaccination of Chickens

[0156] Leghorn type Specific Pathogen Free (SPF) chickens (Hy-Vac, Adel, IA) were vaccinated with the compositions of example 6. Briefly, 74 specific pathogen free (SPF) leghorn chickens were obtained at 11 weeks of age (WOA) and were allowed to acclimate for one week prior to vaccination. Birds were divided into 3 groups designated as Antigen A+B (21 Birds), Antigen A (21 birds), Antigen B (21 Birds) and a group of non-vaccinated controls (11 Birds). Birds were identified by colored leg bands. At 12 WOA, all birds in groups Antigen A+B, Antigen A and Antigen B were vaccinated with 0.25 ml of the appropriate vaccine intramuscularly in the breast. A group of 11 chickens received no vaccine and served as the control group. At 15 weeks of age or twenty one days post second vaccination all vaccinated birds received a second vaccination

#### Example 8

##### Preparation and Administration of Challenge Organism

[0157] One day prior to challenge, one cryogenic bead of *Pasteurella multocida*, strain X-73 of example 1, was aseptically removed from the frozen stock and spread onto two TSA II plates with 5% sheep blood (Becton Dickenson) which were then streaked for isolation of colonies. The plates were incubated for 16-19 hours at 37° C. On day 14 or 14 days post second vaccination several colonies from the sheep blood agar plate were scraped from the blood plate and suspended in a sterile 13x100 test tube. Colonies were added or the suspension diluted with tryptose broth to obtain a % T of 71-75% at 630 nm using a Spectronics 20D (ThermoFisher) spectrophotometer. The suspension was serially diluted in tryptose broth to 1:100,000 dilution. This dilution was dispensed into serum vials for challenge.

[0158] Twenty chickens of each vaccinated group and 10 chickens of the control group were inoculated intramuscularly in the breast with 0.5 ml of the above diluted challenge. The extra chickens from the vaccinated and the control group were removed at this time.

#### Example 9

##### Challenge Results

[0159] The results, shown in FIG. 2, are an average of three separate challenge studies conducted with the three

serials. The term Prevented Fraction (PF) is the complement of the Risk Ratio (RR) and has long been used by epidemiologists as an expression of the preventative role of a protective factor, or the preventative success of an implemented program of intervention (Miettinen, 1974, Am J Epidemiol 99:325-332) (Field Epidemiology, Oxford University Press, 2002, pp 147-7). Prevented Fraction PF=1-p<sub>2</sub>/p<sub>1</sub>, where p<sub>2</sub>=affected fraction in the vaccinated group, and p<sub>1</sub>=affected fraction in the unvaccinated or placebo group. Veterinary fowl cholera vaccines are expected to routinely demonstrate a PF of 53% or greater, preferably 62.5% or greater. Vaccines that demonstrate a PF of less than 53% demonstrate protection.

[0160] Table 1 presents mortality data from FIG. 2 and the calculations for Prevented Fraction for Antigen B of FIG. 2. Vaccine efficacy or prevented fraction was calculated to be 30%, indicating that vaccination prevented 30% of the cases that might have otherwise occurred among vaccinated birds had they not been vaccinated. These calculations were repeated for antigens A and A+B to obtain the prevented fraction for each vaccine.

TABLE 1

	DEAD	NOT DEAD	TOTAL	RISK
Vaccinated Antigen B	40%	60%	100%	40%
Not Vaccinated	57%	43%	100%	57%
TOTAL	97%	103%		

Relative Risk = 40/57 = .70 or 70%

Vaccine Efficacy = (57 - 40)/57 = 0.3 or 30%

[0161] The difference between vaccines and controls are significant ( $p < 0.05$ ) for two of the three vaccine groups, but the prevented fraction was 53%, 30%, and 44% for antigen 1, antigen 2, and the combined antigen, respectively. While the vaccine met the 53% PF level in one case, it could not be demonstrated consistently, and therefore is inadequately effective for a commercial fowl cholera vaccine. Based upon the banding profiles observed in example 4, we expected very good protection. Unfortunately, the protection observed was less than expected. These unexpected results suggest that bands visually appearing to overlap on a SDS-PAGE gel as shown in FIG. 1 are not enough to confer the desired protection level.

#### Example 10

#### Known Genome Analysis

[0162] Current known genomes were surveyed for potential metal regulated proteins. Several strains of *Pasteurella multocida* have full or partial genome sequences available in the public domain. The Universal Protein Resource (Uniprot) is a comprehensive resource for protein sequence and annotation data. At the time of the search, the full or partial sequence data of serotype 1 reference strain X-73 and the serotype 3 reference strain P-1059 were present in this database. For each of these organisms, we searched the database for all proteins having molecular weights between 50 kilodaltons and 150 kilodaltons that had any annotation reference to iron uptake. The identified proteins were organized into a table (FIG. 3), where those of different strains

having a similarity of greater than or equal to 95% were grouped together. Fourteen distinct proteins were identified among these strains.

[0163] The proteins were compared to expressed metal regulated proteins of the previous vaccine candidate (MS061130) as identified by MALDI-TOF analysis. There were eight proteins that appeared to be absent from the vaccine candidate, but were identified in additional genomes of other *Pasteurella multocida* strains derived from avian species. (See FIGS. 3 and 4).

[0164] These results demonstrated that our vaccine candidate (MS061130) was potentially missing 7 desirable proteins that were identified by genomic analysis of additional strains of *Pasteurella*. This same observation was seen in the analysis of the genome of the challenge strain that was used in example 9 which also had the seven protein subset that was not present in the MS061130 vaccine candidate. Thus, the vaccine strain described in Examples 6-9 was missing 7 proteins that were present in the challenge strain (FIG. 3). The Venn diagram in FIG. 4 illustrates the number of proteins present in relevant challenge strains P1059 and X73 that are not present in the vaccine strain MS061130. These observations led us to hypothesize that a combination of two or more strains with comprehensive siderophore receptor coverage of the seven protein subset would provide a higher degree of protective efficacy as well show cross protection independent of serotype.

#### Example 11

#### Comparison of Iron-Restricted Protein Banding Patterns in Wild and Type Strains of *Pasteurella Multocida*

[0165] In order to more fully understand metal regulated proteins in *Pasteurella*, thirty three type strains and clinical field strains of *Pasteurella multocida* were grown in iron-restricted medium and processed to purify membrane bound proteins in the manner of example 3. The resulting protein compositions were analyzed for metal regulated protein banding patterns on Sodium Dodecyl Sulfate-poly-acrylimide gel electrophoresis (SDS-PAGE) gels (Criterion TGX Stain-free gels, Bio-Rad laboratories, Hercules, CA). The individual lanes were compared and grouped using Phoretix 1D, a 1-dimensional gel analysis program (TotalLab, Newcastle upon Tyne, UK). A dendrogram was created with Phoretix 1D software showing banding pattern similarities as illustrated in FIG. 5. The analysis of the type strains and clinical field isolates revealed two predominant patterns of expressed metal regulated proteins; a four-band pattern as marked by the brackets [ ], and a three-band pattern as shown by braces { }. Other strains appear to have variations on these two patterns, missing one or two protein bands.

#### Example 12

#### Analysis of Proteins by MALDI-TOF

[0166] Thirteen isolates were chosen from example 11 as representative of the various band patterns observed in FIG. 5. Each protein band in the molecular weight range of 75,000 to 115,000 from these thirteen isolates of example 11 were excised from the SDS-PAGE gels for protein identification analysis via Matrix Assisted Laser Desorption Time of Flight mass spectrometry (MALDI-TOF MS) and peptide mass fingerprinting. The peptide masses were compared to a database (Uniprot, in this case) containing known protein

sequences of the genome. This was achieved by using computer programs that translate the known genome of the organism into proteins, then theoretically cut the proteins into peptides, and calculate the absolute masses of the peptides from each protein. They then compare the masses of the peptides of the unknown protein to the theoretical peptide masses of each protein encoded in the genome. The results are statistically analyzed to find the best match. The identities of these proteins were grouped in a greater than 93 percent similarity and the results are shown in the table in FIG. 6. A total of 38 bands from the 13 isolates were analyzed and were identified to be one of seven proteins as indicated by their molecular weight in FIG. 6. Two of the isolates (1121 and 1135, indicated by the ellipse in FIG. 6), when combined, expressed all seven protein bands. These two strains were chosen for a new combination vaccine because together they expressed all the proteins of the remaining eleven isolates in FIG. 6, and also were representative of the two major banding patterns identified in FIG. 5.

#### Example 13

##### *Pasteurella* Culture Seed Preparation

[0167] Master seed stocks of *Pasteurella multocida* strains 1121 and 1135 from example 12 and FIG. 6 were prepared. Strain 1121 was identified to be serotype 3x4 and strain 1135 was identified to be serotype 2x5. These isolates were isolated from turkeys that had died from avian Pasteurellosis. Typically, the liver or lung was swabbed and the resulting swab was streaked to 5% sheep blood agar plate and incubated at 37° C. for 24 hours. A Master Seed was prepared by inoculating the isolate into pBHI containing 6 g/l yeast extract and 12 µg/ml DP. The culture was then successively transferred into the above media with 18 µg/ml DP for five more passes to adapt the organisms to an iron-restricted environment and to maximize growth and expression of desired proteins. The cultures were concentrated by centrifugation for 30 minutes at 7,000 rpm (Beckman Coulter, Brea, CA) and resuspended in fresh pBHI containing 20% glycerol (as a cryoprotectant) but no DP. The resulting Master Seeds were aliquoted in 2.2 ml volumes into 3 ml cryovials and stored at or below-60° Celsius. The isolates were given the identification numbers PM1121 20140911 and PM1135 20140925 and established as Master Seeds MS1121 and MS1135, respectively. The Master Seeds were then expanded into working seeds that were used for the production of metal regulated proteins.

#### Example 14

##### Production of Metal Regulated Proteins

[0168] *Pasteurella multocida* can be grown under controlled fermentation conditions so as to express proteins, including proteins associated with the outer membrane. The bacteria can then be harvested and the proteins isolated, purified, and used as immunogens in a composition.

##### Fermentation:

[0169] A cryogenic vial of the working seed of strain 1135 (1 ml at 10<sup>9</sup> CFU/ml) of example 13 was used to inoculate 300 ml of 37° C. modified porcine brain heart infusion media (pBHI) (BD Difco) containing 6 g/l yeast extract

(BD). The culture was incubated at 35-38° C. while shaking at 30-400 rpm for 5 hours, at which point was transferred to 1500 ml of modified pBHI containing 22 µg/ml DP to restrict available iron. The culture was incubated at 37° C. for 2 hours and 38 minutes until it reached an optical density greater than or equal to 0.6. The entire culture was transferred to 15 liters of modified pBHI plus 22 µg/ml DP in an 18 liter flask. This culture was incubated for approximately 2 hours and 30 minutes and reached an optical density of 0.75 at 540 nm. The culture was then transferred to a 300 liter fermentor containing 270 liters of modified pBHI with 22 ug DP to maintain iron restriction, and 40 ml per liter of a 50% glucose solution as a carbon source. The culture was agitated at 35-38° C. with agitation controlled by dissolved oxygen content which was maintained at or above 30%. The culture grew to an optical density, as measured by a Beckman series 600 spectrophotometer, of approximately 5.4. At the end of fermentation, the culture pH was adjusted above 8.5 and the temperature adjusted at or below 15° C.

##### Harvest of Whole Cells:

[0170] The whole cell bacterial suspension was concentrated to approximately 1/10 volume by tangential flow filtration with a Pall Filtron Tangential Flow Maxisette-25 (Pall Filtron) equipped with two 30 ft<sup>2</sup> Alpha 300-K open channel filters, catalog No. AS300C5, (Pall Filtron) connected to a Waukesha Model U-60 feed pump (Waukesha Chemy-Burrell, Delevan, Wis.) set at about 44 Hz. The culture was further washed with Tris-buffered saline, and then concentrated to ~45 liters of whole cell suspension. A subsample of the cell suspension was dried and weighed to yield a final dried pellet weight. The final whole cell suspension was then frozen at -20°.

##### Whole Cell Disruption:

[0171] The whole cell suspension was thawed at 5-7° C. for 3 days. Individual containers of whole cell were pooled and mixed in a sterile tank equipped with a mixer set at 40%. The liquid cell suspension was disrupted by homogenization. Briefly, the suspended whole cell was passed twice through an Avestin homogenizer, model Emulsiflex C500B, (Avestin, Inc., Ottawa, Canada) at a pressure of 15,000-18,000 pounds per square inch (PSI) until a 1:100 dilution of homogenate reached 80 percent transparency (% T at 540 nm) on a Beckman series 600 Spectrophotometer (Beckman instruments, Brea, CA). A second process tank was connected to the homogenizer such that the fluid could be passed back and forth through the homogenizer for multiple passes while maintaining a closed system. Sodium lauroyl sarcosinate (Hamposyl L-30, Chattem Chemicals, Chattanooga, TN) was added at 3% volume/volume to the disrupted cell homogenate to solubilize the cell membrane and non-membrane proteins. Solubilization continued for approximately 72 hours at 2-7° C. Cell debris was removed by centrifugation utilizing a Sharples T1 continuous centrifuge set at 11 PSI.

##### Protein Harvest:

[0172] The protein suspension was washed with 800 liters of Tris-buffered water (TBW) (pH 7.4) containing 0.2% formalin by continuous wash tangential flow filtration (Maxisette 300 kDa filter, Pall corp, NY, NY). The protein was further washed with 100 liter of TBW containing

formalin and 10% ethanol. Following the ethanol wash, the protein suspension was further washed with 200 liters of TBW plus formalin (0.2%) to remove the ethanol. When the remaining TBW was washed from the system, the volume was reduced by continued filtration until the final antigen volume was approximately 20 liters. The final antigen lot 1135-A0001 was then inactivated with the 0.2% formalin with continuous mixing at 33° C. for 93 hours.

[0173] The working seed of strain 1121 was used to prepare antigen lot 1121-A0001 in the same manner as strain 1135.

#### Example 15

##### Preparation of Vaccine

[0174] The antigens prepared in Example 14 were used to prepare two 20 liter water in oil vaccines with 50% mineral oil as adjuvant and tween/span emulsifier as described in example 6. Briefly, 10,145 ml of Drakeol 6 mineral oil and 609 ml Span 85, as an emulsifier, were added to a sterile vessel equipped with a Silverson model 150UHSL mixer (Silverson Machines, East Longmeadow, MA). The mixer was set to 52 hz and the adjuvant phase recirculation was set at 2.5 liters per minute. The aqueous phase ingredients were added to a second vessel: Tween 85 (517 ml), 540 ml of antigen lot 1121-A0001, 1420 ml of antigen lot 1135-A0001 and 7010 ml of phosphate buffered saline, pH 7.4,—was slowly added over the course of 20 minutes. The vaccines were formulated to contain 100 µg total antigen (50 µg from each strain) per 0.25 ml dose, and 200 µg total antigen (100 µg of each strain), respectively. These two antigens, combined, represent the 7 proteins, of example 12 and FIG. 6, required for increased heterologous serotype protection.

#### Example 16

##### Vaccination and Challenge Model

[0175] The vaccines prepared in example 15 was used in a vaccination challenge model against type 1 challenge in SPF chickens. Briefly, 60 SPF leghorn chickens, both males and females, were obtained from Valo BioMedia (Adel, IA). The chickens were sorted by sex, and divided into three groups of 20 birds of mixed hens and roosters, each group with equal numbers of males and females. The birds were identified by numbered wing bands; each bird received identical numbers on the left and right wing to preserve identity. One group was vaccinated with 0.25 ml of the 200 µg vaccine of example 14. A second group was vaccinated with 0.25 ml of the 100 µg vaccine of example 14. A third group was vaccinated with a placebo containing the oil adjuvant in phosphate buffered saline (PBS) in place of the antigen. All birds were vaccinated subcutaneously in the back of the neck at 12 and 15 weeks of age, and all birds were challenged intramuscular in the breast at 17 weeks of age with approximately 6,000 cfu of the serotype 1 challenge strain X-73 prepared in the manner of example 8. The results of the study are shown in FIG. 7. It is clear that the level of protection is vastly improved with a preventive fraction of 69 and 75 percent and a p-value of 0.001 and ≤0.0004 for the 100 µg and 200 µg vaccinated groups, respectively. These results clearly show that the combination of two antigens, expressing all seven proteins of FIG. 6, confer increased prevented fraction against heterologous challenge in a chicken vaccination and challenge model.

#### Example 17

##### Zinc and Respiratory Pathogens

[0176] In our previous research (data not shown) we have shown that many gram-negative pathogens express a novel protein under zinc restriction. An abundance of these bacteria are respiratory pathogens. Some of these pathogens include *Moraxella catarrhalis*, *Haemophilus parasuis*, *Mannheimia haemolytica*, *Acinetobacter baumannii*, *Pasteurella multocida*, *Bordetella pertussis*, and *Actinobacillus pleuropneumoniae* (Stork et al. PLOS Pathogens, 6(7), e1000969 2010). It has been reported that this Zinc acquisition protein is a transmembrane beta-barrel protein now identified as ZnuD located in the outer cell wall that utilizes a single surface exposed loop containing two cysteines that when brought together by a disulfide linkage forms a cluster of histidines that can interact with a zinc ion (Stork et al., PLOS Pathogens, 6(7), e1000969. 2010). ZnuD thereby acts as a scavenger protein used for the acquisition of zinc sequestered by the host organism.

[0177] All these ZnuD homologs contain His- and Asp-rich regions that have been identified in the binding of zinc. This suggests that zinc acquisition mechanisms are important for bacteria residing in the respiratory tract. The current explanation is that unbound zinc concentration within the mucosal tract is too low to allow for simple diffusion across the bacterial membrane via porin proteins (Stork et al., PLOS Pathogens, 6(7), e1000969 2010). Thereby, proteins such as ZnuD are required to scavenge for zinc. This scavenging not only includes unbound zinc, but also the stripping of sequestered zinc from proteins such as calprotectin (Liu et al., Cell host & microbe, 11(3), pp 227-239. 2012). This theory has been supported by the detection of convalescent antibodies towards ZnuD from healthy carriers, which allows the assumption that ZnuD is expressed by bacteria that reside in the respiratory tract (Stork et al., PLOS Pathogens, 6(7), e1000969. 2010). The goal of this experiment was to identify, clone and evaluate the efficacy of a zinc acquisition protein of *Pasteurella multocida*. If a ZnuD homolog is expressed in *Pasteurella multocida*, it offers another possible target for vaccine development currently unexplored for controlling this respiratory pathogen.

#### Example 18

##### Genomic Sequencing of *Pasteurella Multocida* P-1059

[0178] The isolate of *Pasteurella multocida* used for this study was reference strain P-1059 of example 1. The genomic DNA was isolated from the *Pasteurella multocida* isolate using the Invitrogen by Life Technologies ChargeSwitch gDNA Mini Bacteria Kit (Life Technologies, Carlsbad, CA, product number: CS11301). Prior to extraction of genomic DNA, a fresh culture of the isolate was grown on Trypticase Soy Agar II with 5% Sheep Blood (Becton, Dickinson and Company, Franklin Lakes, NJ, product code: 221261) overnight at 37° C. The procedure followed the manufacturer protocol. The final yield was 33.7 µg of genomic DNA, which was stored at -20° C. until sequencing. The genomic DNA was submitted to ACGT, Inc. for sequencing (Wheeling, IL). Genbank reference sequences CP004752 and CP004753 were used in the assembly of the complete genomic sequence of the isolate.

## Example 19

## Possible Target Genes

**[0179]** After receiving the complete genomic sequence of the isolate, tblastn alignments were performed with the National Center for Biotechnology Information (NCBI) database to identify the possible genes of interest based on homology towards ZnuD (GenBank AAF62323.1). A blastx search was used to identify the proteins translated by the homologous genes found with the tblastn alignment. Pairwise sequence alignments of the identified ZnuD homologues and ZnuD zinc affinity region were performed using EMBOSS Matcher (Rice et al., Trends in genetics, 16(6), pp 276-277. 2000.).

## Example 20

## Expression of a Zinc Acquisition Protein

**[0180]** To identify the zinc acquisition protein of *Pasteurella multocida* we used the sequence of the zinc acquisition protein of *Mannheimia haemolytica* previously identified in our laboratory. Briefly, the expression of the target protein was evaluated by supplementing Bacto Brain Heart Infusion, Porcine culture medium (BHI) (Becton, Dickinson and Company, Franklin Lakes, NJ, product code: 256110) used for bacterial cell growth with TPEN (Tokyo Chemical Industry Co., Ltd., Portland, OR, product code: T1487). A titration of TPEN over a range of 0-100 µM during bacterial growth was used. The BHI media containing the various concentrations of TPEN were inoculated with a starter culture of the isolate in BHI containing no TPEN. The inoculum volume was 1 percent of the total volume of the final culture. The cultures were allowed to grow with vigorous agitation at 37° C. to an OD<sub>540</sub> of 1.0, unless bacteria were unable to grow at certain concentrations.

**[0181]** Once an OD of 1.0 was obtained, the bacteria were pelleted by centrifugation at 7,000×g for 10 minutes. An outer membrane preparation was performed on the pelleted bacteria to isolate the integral membrane proteins within the outer membrane following a procedure described by (Molloy et al., European Journal of Biochemistry, 267(10), pp 2871-2881. 2000) with modifications. The bacteria pellet was re-suspended in 30 ml of 60 mM Tris-HCl (Sigma-Aldrich, St. Louis, MO, product code: T3253) and 2.5 mM EDTA (EMD, Billerica, MA, product code: EX0539-3), pH 8.5. The bacteria were then lysed by sonication using a Sonifier S-450A analog ultrasonic processor with a ½" diameter Tipped Bio-Horn attached to a102-C converter (Branson Ultrasonics Corporation, Danbury, CT, product code: 101-063-198, 101-47-037, and 101-135-066 respectively) for 1 minute and 30 seconds on ice at a power setting of 9 and a duty cycle of 90. The sample was clarified by centrifugation at 39,000×g and 4° C. for 20 minutes to remove large cell debris. To remove the inner membrane, peptidoglycan, and proteins not found within the outer membrane, Hampsyl L-30 (Chattem Chemicals, Inc., Chattanooga, TN, product code: BD2099) was added to the supernatant at a final concentration of 1% and incubated for 16 hours at 4° C. while rocking end-over-end. The outer membrane was pelleted by centrifugation at 39,000×g and 4° C. for 2 hours. The outer membrane pellet was washed and re-suspended in 25 mM Tris-HCl buffered water, pH 7.2.

**[0182]** The integral membrane proteins were visualized by SDS-PAGE. Prior to the electrophoresis, a BCA (Thermo Fisher Scientific Inc., Rockford, IL, product code: 23225) was performed to quantify the protein within each sample as to allow for equivalent protein loads of 30 µg. Samples were brought up to equal volume prior to the addition of 3× loading buffer (New England BioLabs Inc., Ipswich, MA, product number: B7703S). The samples were compared on the gel to look for expression of the target protein.

## Example 21

## Protein Identification

**[0183]** The banding profile from the outer membrane preparation was analyzed by Matrix-assisted Laser Desorption/Ionization Mass Spectrometry (MALDI) to determine if any of the bands of interest match the genes identified by BLAST homology. The bands were cut out of the gel with a scalpel and sent to the Mayo Clinic Proteomics Core facility to be analyzed.

## Example 22

## Structure Modeling

**[0184]** Swiss Model was used to create a 3-D structural model of zinc acquisition protein of *Mannheimia haemolytica* based on homology sequencing (Arnold et al., Bioinformatics, (22), pp 195-201. 2006.). The PDB template used to create the protein model was 4epaA. The extracellular loop (ECL) sequence positions were identified using the 3-D structural model obtained from Swiss Model (Arnold et al., Bioinformatics, (22), pp 195-201. 2006.), Prediction of TransMembrane Beta-Barrel Proteins (Bagos et al., BMC Bioinformatics, 5(29). 2004.), Orientations of Proteins in Membranes (OMP) database and Positioning of Proteins in Membranes (PPM) server (Lomize et al., Nucleic Acids Res, 40 (Database issue): D370-6. 2012).

## Example 23

## Target Gene Identification

**[0185]** Once the proteins within the banding profile were identified, they were compared to the proteins of interest identified by BLAST homology. One protein within the banding profile matched a protein identified by the BLAST homology. The protein was identified as OMR family outer membrane iron receptor (GenBank: EDN73812.1) also referred to as the zinc acquisition protein in this paper. The target gene was located at nucleotides 1,160,319 . . . 1,162,691 within the genome. The signal peptide was identified using SignalP 4.1 (Petersen et al., Nature methods, 8(10), pp 785-786. 2011)). The nucleotide and amino acid sequences of the zinc acquisition protein identified in *Pasteurella multocida* is shown in FIG. 22.

## Example 24

## The Homology of the Zinc Acquisition Protein Across Respiratory Pathogens

**[0186]** Using the sequence of the acquisition protein of *Mannheimia haemolytica* (data not shown) the homology of zinc acquisition protein across respiratory pathogens, particularly the zinc affinity region, was determined with the

protein blast search tool established by NCBI. Ten different species of bacteria were found to contain a zinc acquisition protein homologue with greater than 49% identity. All species identified are respiratory pathogens. Aligning the homologous proteins to the zinc affinity region of ZAP using Clustal Omega highlights strikingly similar amino acid motifs. All the homologues contain two cysteines and multiple histidines and aspartic acids. The amino acid motifs are emphasized in FIG. 25 using GLAM2. The spatial arrangement of the cysteines, histidines, and aspartic acids remain constant across the homologues.

TABLE 2

Strain	Accession Number	% Identity	% Similarity
<i>Bibersteinia trehalosi</i>	AHG81836.1	98	97
<i>Actinobacillus pleuropneumoniae</i>	WP_005612269.1	93	95
<i>Mannheimia granulomatis</i>	WP_027074597.1	75	82
<i>Mannheimia variogena</i>	AHG73391.1	72	79
<i>Haemophilus parasuis</i>	WP_021114857.1	62	78
<i>Moraxella boevrei</i>	WP_026212957.1	53	69
<i>Psychrobacter phenylpyruvicus</i>	WP_028858792.1	54	69
<i>Pasteurella multocida</i>	WP_016534590.1	58	72
<i>Moraxella bovoculi</i>	KDN24548.1	52	67
<i>Conchiformibius steedae</i>	WP_027021676.1	49	64

The results show that multiple respiratory pathogens indeed show a degree of homology to the zinc acquisition protein of *Mannheimia Haemolytica* ranging from 49 to 98 percent identity.

#### Example 25

#### Expression and Identification of a Zinc Acquisition Protein

##### Cloning

**[0187]** The *Pasteurella multocida* zinc gene was PCR amplified from genomic DNA of strain P-1059 and inserted into plasmid vector pQE-T7-2 (Qiagen; Hilden, Germany) using Gibson assembly methods. The assembled plasmid was then transformed into NEB T7 Express Competent *E. coli* (New England Biolabs; Ipswich, MA; catalog #C2566H) for clonal selection and expression.

##### Expression

**[0188]** A starter culture of LB broth containing 50 µg/mL kanamycin was inoculated with a clone containing the recombinant plasmid. Culture was grown to an OD540 of 0.6 and then transferred to fresh LB broth. The new culture was allowed to grow again to an OD540 of 0.6. Expression was then induced with 1 mM IPTG and incubated an additional 16 hours. The cells were collected by centrifugation and stored at -80° C. until processing.

##### Inclusion Body Purification

**[0189]** Cells were thawed and resuspended in BugBuster Protein Extraction Reagent (Merck Millipore; Billerica, MA; catalog #70584-4). 10 µL of Lysonase Bioprocessing Reagent (Merck Millipore; Billerica, MA; catalog #71230) was added and the suspension was incubated for 1 hour at ambient temperature. The suspension was then sonicated (Branson Sonifier, model 102c; Branson Ultrasonics, Danbury, CT) for 30 seconds using a micro tip, output 6, at a 90% duty cycle. The lysate was centrifuged at 16000×g for 20 minutes to collect the insoluble material. The pellet was washed three times: first with BugBuster, second with 1/10x

BugBuster, and third with tris-buffered water. The inclusion body prep was solubilized in tris-buffered saline containing 8 M urea and 1 mM dithiothreitol at pH 8.0, yielding 10 mg recombinant protein. FIG. 24 shows the different stages of expression of the zinc acquisition protein run on a 10% Criterion stain free-free TGX Gel run at 300V (Bio-Rad, Hercules, CA) to include: the un-induced whole cell preparation, the induced whole cell preparation, the lysed cell supernatant, 1x bugbuster wash, 1/10x bugbuster wash, the solubilized inclusion bodies and the pellet post solubilization.

#### Example 26

#### Vaccine-Mediated Protection of a Novel Recombinant Zinc Protein of *Pasteurella Multocida* in a Chicken Sepsis Model

**[0190]** The purpose of the following experimental study was to evaluate the protective efficacy of a recombinant zinc protein (rZinc) against a virulent challenge of *Pasteurella multocida* in a chicken sepsis model. The efficacy of the rZinc protein of example 25 was compared to the efficacy of the *Pasteurella multocida* SRP extract composition of example 14 compared to non-vaccinated controls. Briefly, 30 SPF leghorn chickens, (males), were obtained from Valo BioMedia (Adel, IA). The chickens divided into three groups of 10 birds each. The birds were identified by colored leg bands. Treatment groups were designated as Group-A (Placebo), Group-B (rZinc protein) and Group C (*Pasteurella multocida* protein extract). The outcome parameter used to evaluate vaccine efficacy in this experiment was total mortality between vaccines compared to the non-vaccinated Placebo group. Food and water was supplied ad libitum to all chickens.

#### Example 27

#### Preparation of the Immunizing Compositions

**[0191]** The vaccines (SRP extract and rZinc) were prepared from the protein compositions of examples 14 and 25 by diluting the antigen into phosphate buffered saline (PBS) containing 8.0 g/l NaCl, 0.2 g/l KCl, 1.44 g/l Na<sub>2</sub>HPO<sub>4</sub> and 0.24 g/l KH<sub>2</sub>PO<sub>4</sub> pH 7.4. Each vaccine was prepared by emulsifying the following ingredients: 44.44% aqueous protein suspension in 0.1% formalized saline, 50% Drakeol 6 mineral oil (VOPAK USA, Inc, Kirkland, WA), 3.0% Span 85 and 2.56% Tween 85 (Ruger Chemicals, Hillside, NJ). The final bird dose for each vaccine composition (rZinc and SRP extract) was 250 µg and 400 µg administered in a volume of 0.5 ml and 0.25 ml respectively. The placebo was prepared by replacing the antigen with physiological saline and emulsifying the suspension using the above formulation in a dose volume of 0.5 ml. All birds were vaccinated subcutaneously at 14 and 17 weeks of age (21-day interval).

#### Example 28

#### Challenge and Results

**[0192]** At 19 weeks of age all birds were challenged intramuscularly in the breast with 5450 Colony Forming Units (CFU) of *Pasteurella Multocida* (serotype 1) strain X-73 prepared as described example 8. Birds were observed for mortality for 14 days following challenge. The results clearly demonstrate the efficacy of both the *Pasteurella*

*Multocida* SRP Extract and the rZinc vaccines having a percent survival of 100% and 50% respectively, in contrast to the non-vaccinated controls having no survivors (FIG. 23). The recombinant zinc protein in this study was evaluated at a single concentration of 250 µg, showing a 50% survival rate ( $p=0.0325$ ) against a virulent challenge.

[0193] The complete disclosure of all patents, patent applications, and publications, and electronically available material (including, for instance, nucleotide sequence submissions in, e.g., GenBank and RefSeq, and amino acid sequence submissions in, e.g., SwissProt, PIR, PRF, PDB, and translations from annotated coding regions in GenBank and RefSeq) cited herein are incorporated by reference in their entirety. Supplementary materials referenced in publications (such as supplementary tables, supplementary figures, supplementary materials and methods, and/or supplementary experimental data) are likewise incorporated by reference in their entirety. In the event that any inconsistency exists between the disclosure of the present application and the disclosure(s) of any document incorporated herein by reference, the disclosure of the present application shall govern. The foregoing detailed description and examples have been given for clarity of understanding only. No unnecessary limitations are to be understood therefrom. The

invention is not limited to the exact details shown and described, for variations obvious to one skilled in the art will be included within the invention defined by the claims. Unless otherwise indicated, all numbers expressing quantities of components, molecular weights, and so forth used in the specification and claims are to be understood as being modified in all instances by the term "about." Accordingly, unless otherwise indicated to the contrary, the numerical parameters set forth in the specification and claims are approximations that may vary depending upon the desired properties sought to be obtained by the present invention. At the very least, and not as an attempt to limit the doctrine of equivalents to the scope of the claims, each numerical parameter should at least be construed in light of the number of reported significant digits and by applying ordinary rounding techniques. Notwithstanding that the numerical ranges and parameters setting forth the broad scope of the invention are approximations, the numerical values set forth in the specific examples are reported as precisely as possible. All numerical values, however, inherently contain a range necessarily resulting from the standard deviation found in their respective testing measurements.

[0194] All headings are for the convenience of the reader and should not be used to limit the meaning of the text that follows the heading, unless so specified.

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#### SEQUENCE LISTING

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mol_type = genomic DNA
organism = Pasteurella multocida

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aaatttcaatt	accaacgtgg	tcgtttggat	ggcgatcgct	caatgaatgc	gattcaggct	2520
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tatatcacac	gtgtgagtga	aaaaaaagca	aagacacttta	taatatgttc	tataaagaaac	2640
agggatataa	agatagtgtc	gttctgttgg	gaagtgtatg	ctatacgct	gttgtatgcgg	2700
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accgttaataa	tttgacatgg	gaatctgtc	gttgcattaa	accatttgtt	acaagtaatt	2820
taattaatca	gaaaacaggc	cgaggattaa	atcggtttta	ctcaccagggt	cgtatattta	2880
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SEQ ID NO: 2            moltype = AA    length = 968

FEATURE                Location/Qualifiers  
source                1..968  
mol\_type = protein  
organism = Pasteurella multocida

SEQUENCE: 2

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FKELFEGYGN	PNNTRNSVEI	ETLKVKIAK	GADSVKVGSS	SLGGAVLFET	KDARDFLTEK	180
DWHIGYKAGY	STADNQGLNA	VTLAGRYQMF	DALIMHSKRH	CHELENYDYK	NGRDIQGKER	240
EKADPYTITK	ESTLVKFSFS	PTENHRFTVA	SDTYLQHSRG	HDLSYNLVAT	THIQLDEKES	300
RHANNDLTKR	NVSFTYENYY	VTPFWDTKLK	SYSQQRITTR	ARTEDYCDGN	EKCDSYKNPL	360
GLQLKEKGIV	DRNGDPVNLLK	LVDGKHQVUD	KAGKPFDVAS	GTNYAAFSGK	ELSPSSFWLD	420
CSIFDCSKPI	NTYKYRYTSS	EPTLQQITLN	KTMBINGKTF	ATYDGRGHYI	ILPNSKGYLP	480
LDYKERDLNT	TKQINLSDLT	KAFTLFEIEN	ELSYGGVYAK	TTKEMVNKAG	YYGRNPTWWA	540
ERTLQGSWKG	LRECKTSSSY	NGMLCPRHEP	LTSFLIPVEA	TTKSLYFADN	IKLHNMLMSVD	600
LGYRYDDIKY	QPEYIPGVTP	KIADDMVKGL	FIPLPEGEKV	TVGTMVFTRP	LTQAOQIRKNA	660
EENIAYIAQE	KRFKKHYSYL	GATFDPLNLF	RVQVKYSKGF	RAPTSDELHF	TFKHDFFTIL	720
PNPVLKPEEA	KNQEIALTVH	DNWGFVSTSV	FQTKYRHFI	LAYLGSRNLS	NSVGGQAAQR	780
DFQVYQNVNV	DNAVKVGLEI	NARLNLGYFW	HVLDFGNTSY	KFTYQRGRLD	GDRPMNAIQP	840
KASVFGGLYD	HKENKFGADL	YITRVSEKKA	KDTYNMFYKE	QGYKDASWR	RSDDYTLVDA	900
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KFSAEITF						968

SEQ ID NO: 3            moltype = DNA    length = 2373

FEATURE                Location/Qualifiers  
source                1..2373  
mol\_type = genomic DNA  
organism = Pasteurella multocida

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attgtcgtgt	atgcggatca	aaataatca	atgatgttca	caacagatgt	cactcaagat	180
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caacaatcac	gtcaaattaa	tccacaagg	gatcggtata	aacaaacaca	tacaagacgt	840
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ctccgcattt	accaacgagg	aaaagactac	aaaacgaat	gtttaacc	ccaaat	1860
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tggggctcac	gtttaaatct	tgatcgttt	gttccaataat	ttgatcttc	ttgggcta	2100
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gtatatcgta	gctatgatta	tggacttat	acacaatggg	ataccagtct	tcgctggcag	2220
ccaaacctcg	cagaaaaaca	ccgtccttat	attaaatttg	atgtacttaa	tgtgttaat	2280
aaaacacgta	aaggcgcggg	tccaaatgga	caagatctcg	gcatctacac	ccccggcgt	2340
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SEQ ID NO: 4            moltype = AA   length = 790  
 FEATURE                Location/Qualifiers  
 source                1..790  
                       mol\_type = protein  
                       organism = *Pasteurella multocida*  
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INNDMFGFSD	LFDGTMATPV	FANHSQGYFF	DANLLSSIVV	HDSNVASALG	GFAGGAIVAK	180
TKQYDGKDR	KFSYRTTDAS	WAKFKVEDKD	LERFKNAPE	GSVAEFQPKY	SKHFFNITAE	240
KGLSENLMGV	IGLSRRTSS	QOSRQINPQG	DRDKQTHTR	SDNALLNFL	TPNDKHFEL	300
GFRYSNYRER	KFFNTNIDSN	VFDYHRAYGV	TFWSWINALQS	GILTTTLAYD	NFDDTRKSAS	360
TYMKTTLTEE	CEEYETEGGMG	NSQLNQKNLH	TSLEYAMNP	NLGSIEHSVS	LGGIYQATKY	420
RFTRHSDAVG	ELYTPDWLNG	NTDKLIITQ	NIAKKGTVK	RYQNIALYVE	DLMTWKNLEF	480
RAGLRLRBRDD	YLKNTNLAPR	TVFRYKPFED	TAFPSVGWNRY	YGRSRFASMKL	SEGIFKLGDH	540
DTFRYKDLSQ	PKTPYSDELS	FGVEQYVANL	AFHLKYILRD	NKQIRIVLQEE	EVMLNGERKK	600
LRYYQRGKDY	KTNVLTFQIN	TQAPWELGBT	RWTSAVAPDW	LDSKAIDHGR	GYNGSTPVIL	660
DGKLMTYEQM	LKKVNAYKET	WGLRLNLDMP	VPIFDLSWAN	TIYVKPPTL	TERVSSNTPE	720
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EFWLEVGYEF						790

SEQ ID NO: 5            moltype = DNA   length = 2184  
 FEATURE                Location/Qualifiers  
 source                1..2184  
                       mol\_type = genomic DNA  
                       organism = *Pasteurella multocida*  
 SEQUENCE: 5  

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tttgcaccaa	cgcacaaat	taatgttac	catacggac	atgcacaatt	aggcgatatt	240
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SEQ ID NO: 6            moltype = AA   length = 727  
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source	1..964
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	organism = Pasteurella multocida
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9	
SEQ ID NO: 9 moltype = DNA length = 2547	
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	mol_type = genomic DNA
	organism = Pasteurella multocida
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SEQ ID NO: 10	moltype = AA length = 848
FEATURE	Location/Qualifiers
source	1..848
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KGLQSVPSYQ YAGSSRTY LDPDLISQVE IEKGPSLQMD ATGATGGVVR VETLRWQDII	180
PQGKNUVGVR KLGTMTNTVS PPPYYTRGGY QTGYISKCLIS NDTGLCQTQT YAPNARYSSH	240
GFDLNAYNYS LAFANKWQNA DLVLAYAKRK QGNYFVGVRHG QTPVIESIEF EEDSVEVKEP	300
RVHEEVEIIGS LTFKENRSTL YRPGEEARLNT SQDNTSYLAQ INVYNDVHL GLAYRHYSR	360
FGEIMSSLN FRAYGALQGE GTEVKVDSYH ANYSYNPTTP YVNLNVNAYF TDSDSNNFTP	420
FIEEYGYSL SRRHAFLVSE QKGLSIENTS IFOLNDKPFIS LKYGLAHSYR RIYQPRNAQA	480
RVRAKGYPED AIGPLYIRDG KRKEWSAFVA ANYPITSWLK ADIGLRYLQS TIYDIVRTE	540
RVNIGGALVP NPNGSGNIWU EKVDVVHKQ APVKNKGMSPI VMVLTFEPIN GVQIYTKYAE	600
ALRSPSLFQA TKGWSMSATA DNLEQLRPER AQNWEAGINL FYENLGGKD ILGFKLAYFN	660
NRIKDYLTRES YSPKDVTQI INIQOSAQFKG IELSAYYDMG KFYAKLAGTY YTAKTKFCILTA	720
EQAGKGEQCQ SGYVYRSNLN NAVPPRLNLH ATLGLTRLFEO KLDIGARYSY YSKRLPVLS	780
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SEQ ID NO: 11	moltype = DNA length = 2355
FEATURE	Location/Qualifiers
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	organism = Pasteurella multocida
SEQUENCE: 11	
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ggtatcgtat ttatgtgtgt ttctttgtt gatttcgtatca aaaaactact gtatgcgcgt	360
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ggcgaaaaat ttgtatgcctt attgtatgtt tcacaacggca cagggcacgca gtttaaaagt	660
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gatttatgtatccatccatcaatccatcaatccatcaatccatcaatccatcaatccatcaatccat	2160
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cgttccatccatccatcaatccatcaatccatcaatccatcaatccatcaatccatcaatccatcaatccat	2340
gaaatccatccatccatcaatccatcaatccatcaatccatcaatccatcaatccatcaatccatcaatccat	2355
SEQ ID NO: 12	moltype = AA length = 784
FEATURE	Location/Qualifiers
source	1..784
	mol_type = protein
	organism = Pasteurella multocida
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QNEMIRDTRD LVRYTTDVGI SDNGRFLKGF AMRGVEDNRV GISIDGVSLP DSEENSILYAR	120
YGNFNNSRSL IDPELIQTID IVRGSDFNA GSGSLGGGVN YNTLDPQHIV KTGNNSVALL	180
RGSYASKNRE WVRTLGIGYV GEKFDAALLMY SQRTGHEFKS RGSGPFRYS SSQHPDPVTQ	240
RFHNYLAKMN YQINDKQRIG LTLMQQTGGR YIDERSYTLM GSQWRADDQ NERLNANLYY	300

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IYAPSTGWLA	YSKFDLQYQK	TDLAAVNYKG	GRHFRTDAKE	LNEIYDRRMK	TVFTRGSVEL	360
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DHVRWNNTFS	SHLGLRYDHT	KLKPKELNAP	CSKACLEEGK	PKPTRFSTVS	TFAGFEAQLS	480
PSWMLGYNIS	TGYRVPTASE	MFFSFTNAYG	TWKSNPSSLK	EKSINHTLSL	KGNSEKGLLD	540
LTLYQTNYRH	PLFEQESLIQ	RTEMRYGRPY	TYQSQEQQMV	NLDKAKIYGV	ELKTHVNLDQ	600
MTAVIPOGFK	FYAALGYSKG	KLSNNASLLS	IOPLKIILGL	DYEATNGKWA	IFNRLTYLGE	660
KRASDAKVYE	IKRRCTEFVT	ETDPWTGQQI	TRCKKRELYP	DLSTYKHLNK	SAVFVDTFGY	720
YKITDDITFR	AGIYNLFNKK	YHTWDALARGI	NANSTLNSVD	REGKGLQRFY	APGRNYAASL	780
EIRF						784

SEQ ID NO: 13	moltype = DNA	length = 2229				
FEATURE	Location/Qualifiers					
source	1..2229					
	mol_type = genomic DNA					
	organism = Pasteurella multocida					
SEQUENCE: 13						
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caacaacaag	cagataatgc	tgctgactta	ataaaatattt	tgccctgggt	aaatatggcg	240
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tttatttgc	ctgtatgtt	aagaaaggat	acagtagaca	aaggaaata	ttctctccaa	420
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ttatgtgtt	ctgtaaagaaa	tgcgaacat	tatacaagac	ctgtatggaaa	taaaatttctt	660
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catttattaa	ctttatccag	tgtttatggc	attcataaa	ggtggaaacc	ttgggcagca	780
aaaagagatg	tgtatgtcgag	accaacagaa	acagaaatata	aacatctatgg	gattgtat	840
gctgtggaaac	gtttaacttgc	ttatggatgat	caaaaatgtt	aaagtatttc	attgaaatat	900
cgtatattac	ctggaaatata	taatgtggat	aattttatcg	ttcagctgag	ttatagtaaa	960
acagaacaga	atgatactcg	ccatggaaaa	gtcacttctt	cattcttagg	tacatttagga	1020
aataaaaatgtt	ggataacttta	ttcagatctt	acttttgcata	taagtaacac	aagtactcta	1080
aatatttggc	gtgtcgagca	tgaactacta	tttgggttac	agttggat	aaataaaaga	1140
aataccctta	tgtatctaa	atggggatgc	aagaaggcag	actataat	tggctat	1200
cagccttatt	atatgccttc	tggacggac	tatacacaag	cattttat	acaagatcaa	1260
ataaaaatggc	agaatttcc	cttttacagga	gggataagat	atgaccat	caataatata	1320
gggcagaaaa	atattagcgc	acgtatata	gatatctct	caggacat	tgatagccag	1380
aaaaaattata	atgggttgc	tattatata	ggtcttaat	atgtgtat	tcattat	1440
agtttattta	cgatatttt	taaaacttgc	cgagccccgt	ttattgtat	acgtat	1500
acacaatata	gtcaagcttc	tgtatctgc	acttctttaa	attttagaaa	agaaatgatt	1560
aatcaaccca	gagttgggtgg	aattttact	ctcaatcat	tatttcagg	aaatgtatgt	1620
tttcaattta	gaactactta	tttttacata	cgccggcaaga	atgaaatctt	caaaacgaga	1680
ggagtttaat	gogtagaaaa	tgcttttagat	gttgataata	gtgtttgtcc	taaaattt	1740
agtaattacc	gtatattacc	tggttatgtt	attcaaggag	cggaatttga	agcttattat	1800
caatcatcgat	attttttgg	tggactgaca	tattttat	taaaaggaaa	acgcgatact	1860
tcaccaagaa	atccatggag	taaaaatcat	acatggatgc	cagaaacatc	acctagaaaa	1920
gcacatcgata	cttttaggtt	taatattcc	gaatattat	ttacggcagg	ttggcgtgt	1980
gagtttgtga	gaaagcaaga	tagatccca	ctatctaatt	attcttaa	atcatattgg	2040
gcattaccc	cttcaaaagg	atatgcctt	catagtgtat	tcttccttg	gagtcctaca	2100
aagattaaag	gaatgaattt	caaagtact	gttgataat	tatthaaccc	accattatt	2160
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caattttaa						2229

SEQ ID NO: 14	moltype = AA	length = 742				
FEATURE	Location/Qualifiers					
source	1..742					
	mol_type = protein					
	organism = Pasteurella multocida					
SEQUENCE: 14						
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QQQADNAADL	INILPGVNMA	GGFRPSSGQTL	NINGMGAED	VRVQLDGATK	SFEKYQQGSI	120
FIEPELLRKV	TVDKGKNSPQ	YGNQGFACTV	KFETKDATDF	LKENQKIGGL	FKYGNNSNNN	180
QKTYSTALVL	YHKGWEPWAA	LGFSVNRASN	YTRPDKSKIL	FSKNNQKSGL	IKVNWQITPE	240
HILLTLSSVYG	IKHGWEWPAA	KRDVMSPRTE	TEIKHYGIDV	AWKRKLVYRD	QKDESYSLK	300
RYLPENNWKI	NLSVQLSYSK	TEQNDTRHEK	VTSSFLGTLG	NKSWITYSDL	TFDISNTSTL	360
NIGRAEHELL	FGLQNLKNKR	NTLMLYHKGGV	KKADYNVGYF	QPYYMPSPRQ	YTQAFYLQDQ	420
IKWQNLFITG	GIRDYDHINNI	GQKNLAPRYN	DISAGHDYSSQ	KNYNGWSYYL	GLKYDVNHYL	480
SLFTNFSKTW	RAPVIDEQYE	TQYSQASVSA	TSLNLEKEMI	NQTRVGGIIT	LNHLFQENDA	540
FQFRRTTYFYN	RGKNEIFKTR	GVNCVENALD	VDSVCPKII	SNYRNLPGYV	IQGAELEAYY	600
QSSYLFGGGLT	YSYVKGKRD	SPRNPKWSKTS	TWIAETSPrk	AIATLGFNIP	EYYFTAGWRA	660
EFVRKQDRSP	LSNDSKASYW	ALPSSKGYS	HSVFFSWSP	KIKGMNFKVT	VDNLNRPYY	720
PYLGELASGT	GRNVKFSLTQ	QF				742

SEQ ID NO: 15	moltype = AA	length = 968
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FEATURE	Location/Qualifiers
source	1..968 mol_type = protein organism = <i>Haemophilus influenzae</i>
SEQUENCE: 15	
VILMLMSQTN QPTNQPTNQN SNASEQLEQI NVLGSNNND NTPPKTAETV KTAQLKRQQ 60 VQDSRDLVRY ETGVTVVEAG RFGSSGYAIR GVDENRAVIT VGDLHQAETL SSQGFKELFE 120 GYGNFNNTRN SVEIETLKVIA KIAKGADSVK VGSGLGGAV LFETKDARDF LTEKDWHIGY 180 KAGYSTADNQ GLNAVTLAGR YQMFDALIMH SKRHGHELEN YDYKNRDIQ GKEREKADPY 240 TITKESTLVK FSFSPTENHFT FTVTADTYLQ HSRGHDFSYN LVKTTYINKV EEELRHTNDL 300 TKRKNSVFT ENYTVTFPWD TLKLQYSQR ITTRARTEDY CDGNEKCDSY KNPLGLQLKN 360 GQIVDRDGPV VNLKLINGR KVVDKNNKLF GLTDEDNNAA FDGKQLGLSG FWFDCTVFD 420 DKPVRTYKYK YSSSNPAVEN VELNKFMQVN GKRFTAYEDK IQSSEKRYVI LPNSKGYLPL 480 DYKERDLNTK TKQINLDSL AFTLFEIENE LSYGGVYAKT TKEMVNKAGY YGRNPTWAE 540 RTLGDQLSGN QHCNCNTNSY NGMLCPRHEP LTSFLIPVEA TTKSLYFSND IKLHNMLSVD 600 LGYRYDDIKY QPEYIPGVTP KIAADDMKGI FIPLPKGKEI KIGNYETTKP LTPEQIRKNA 660 EENIAYIAQE KRFKKHSYSL GATFDPLNFL RVQVKYSKGF RAPTSDELYF TFKHPDFTIL 720 PNPVLPKEEA KNQEIALTVH DNWGFVSTSV FQTKYRHFID LAYLGSRNLS NSVGGQAQAR 780 DFQVYQNVNV DNAVKVGLEI NARLNLGFW HVLDGFNTSY KFTYQRGRLD GDRPMNAIQP 840 KASVFGGLGYD HKENKFGADL YITRVSEKKA KDTYNMFYKE QGYKDSAVRW RSDDYTLVDA 900 VTYIKPVKNV TLQFGVYNLT DRKYLTWESA RSIKPFGTSN LINQQTGAGI NRFYSPGRNY 960 KFSAEITF 968	
SEQ ID NO: 16	moltype = AA length = 968
FEATURE	Location/Qualifiers
source	1..968 mol_type = protein organism = <i>Pasteurella multocida</i>
SEQUENCE: 16	
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SEQ ID NO: 17	moltype = AA length = 971
FEATURE	Location/Qualifiers
source	1..971 mol_type = protein organism = <i>Pasteurella multocida</i>
SEQUENCE: 17	
MRTTTIKFSA ITLALLSYCG VILADSHQEA TELDTITVSS QQDEMNIKEK KVGETVKTAS 60 QLKRQQVQDS RDLVRYETGV TVVEAGRFGS SGYAIRGVDE NRVAITVDGL HQAETLSSQG 120 FKELFEGYGN FNNTRNSVEI ETLKAVAKIAK GADSVKVGSS SLGGAVLFET KDARDFLTEK 180 DWHIGYKAGY STADNQGLNA VTLAGRYQMF DALIMHSKRH GHELEIYDYK NGRDIQGKER 240 EKADPYTITK ESTLVKFSFS PTENHRTVA SDTYLQHSRG HDLSYNLVAT TYIQLDEKES 300 RHANDTLTKR NVSFTYENYT VTPFWDTLKL SYSQQRITTR ARTEDYCDGN EKCDSYKNPL 360 GLQLKEKGIV DRNGDPVNLL VLDGKHQVVD KAGKPFWDAS GTNYAAFSGK QLGPSYFWLE 420 CTVFDCSKPV TTYKRYSTE TPVKBEDIQLN KTMEVNGKTF ATYDMGRERR YIILEPNSQGY 480 LPLDYKERDL NTKTQINLD LTKAFTLFEI ENELSYYGGVY AKTTNGVNVN AGYYGRNPTW 540 WAERTLGQSW NGTLRECKTS SSYNGMLCPR HEPLTSFLIP VEATTKSLYF ADNIKLHNML 600 SVDLGYRYDD IKYQPEYIPG VTPKIADDMV KGLFLIPLPEG EKVTGTVVF TKPLTPEQIR 660 KNAEENIAYI AQGKRPKKHS YSLGTTFDPL NFLRNVQKVH KGFRAPTSDE LYFTFKHPDF 720 TILPNPVLKP EEAKNQEIAL TVHDNWFVTS TSFQTKYRH FIDLAYLGSR NLSNSVGGQA 780 QARDFQVYQNVN VNVDNAVKVG LEINARLNLG YFWHVLDFGN TSYKFTYQRG RLDGDRPMNA 840 IQPKASVVEGL GYDHKENKFG ADLYIITRVS EKAKDTYNMF YKEQGYKDSA IRWRSDDYTL 900 VDAVGYIKPI KNLTQFGVY NLTERKYLTW ESARIKPFQ TSNLINQKGT AGINRFYSPG 960 RNFKFSAEITF 971	
SEQ ID NO: 18	moltype = AA length = 971
FEATURE	Location/Qualifiers
source	1..971 mol_type = protein organism = <i>Pasteurella multocida</i>

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SEQUENCE: 18

MRTTTIKFSA	ITLALLSYCG	AILADSHQEA	TELDTITVSS	QQDEMNIKEK	KVGETVKTAS	60
QLKRQQVQDS	RDLVRYETGV	TVVEAGRFGS	SGYAIRGVDE	NRAITVDGL	HQAETLSSQG	120
FKELFEGYGN	FNNTNRSVEI	ETLKAVAKIAK	GADSVKVGGG	SLGGAVLFET	KDARDFLTEK	180
DWHIGYKAGY	STADNQGLNA	VTLAGRYQMF	DALIMHSKRH	GHELENYDYK	NGRDIQKER	240
EKADPYITIK	ESTLVKFSFS	PTENHRFTVA	SDTYLQHSGRG	HDLSYNLVAT	THIQOLDEKES	300
RHANDLTKRK	NVSFTYENYT	VTPFWDTLKL	SYSQQRITTR	ARTEDYCDGN	EKCDSYKNPL	360
GLQLKEGKIV	DRNGDPVNLLQ	LVDGKHQVVD	KAGKPFDVTS	GTNYAASFSGH	QLGPSYFWLD	420
CTVFDCSKPV	TTYKYRSTE	TPVKEDIQLN	KTMEVNGKTF	ATYDMGRERR	YIILPNSQGY	480
LPLDYKERDL	NTKTKQINLL	LTKTCAFLFEEI	ENELSYGGVVY	AKTTKEMVN	AGYYGRNPTW	540
WAERTLGQS	NGTLRECKTS	SSYNGMLCPR	HEPLITSFLIP	VEATTKSLYF	ADNIKLNHML	600
SVDLGYRYDD	IKYQPEYIPG	VTPKIAADDMV	KGLFIPILPEG	EKVTVGTVVF	TKPLTPEQIR	660
KNAEENIAYI	AQEKRFKHS	YSLGATFDPL	NFLRVQVKY	KGFRAPTSDE	LYFTFKHPDF	720
TILPNPLVLPK	EEAKNQEYQ	TVHNDWGFVS	TSVFQTKYHR	FIDLAYLGSR	NLSNSVGGQA	780
QARDFQVYQ	VNVDAKVKKG	LEINARLNLG	YFWHVLQDFN	TSYKFTYQRG	RLDGDRPMNA	840
IOPKASVFG	GYDHKENKFG	ADLYITRVE	KKAKDTYMF	YKEQGYKDSA	VRWRSDDYTL	900
VDAVGYIKPI	KNLTLQFGVY	NLTDRKYLTW	ESARSIKPGF	TSNLINQKTG	AGINRFYSPG	960
RNFKLSAEIT	F					971

SEQ ID NO: 19                    moltype = AA    length = 792

FEATURE                         Location/Qualifiers  
source                         1..792  
mol\_type = protein  
organism = Pasteurella multocida

SEQUENCE: 19

MEKIDMESAK	NPLKKTTAL	LCCSTAFSLS	AKTDTNADKN	HFLTEIVVVA	DQNKSMSSTQ	60
SVTQDDMKKS	PVTNGNITDV	LRSPNPHVRYE	NSDQNGLQRG	EIKPENISIN	GADYQQTTF	120
VDNVNINNNDM	GFGSDFDGT	MATVPFANHS	QGYFFDANLL	SSIVVHDSNV	SASLGFFAGG	180
AVVAKTKQYD	GKDQLKFSYR	TTDASWAKFK	VEDKDLERFK	NAIPEGSVAE	FQPKYSKHFF	240
NITAEKGKSE	NLGMVIGLSR	RTTSSIQQSRQ	INPOGDRDKQ	THTRRSDNAL	LNFNLTTPNDK	300
HRFELGFRY	NYRERKFFNT	NIDSNTFVYH	RAYGVTFSWI	NALQSGILTT	TLAYDNFDDT	360
RKSASTSMKT	IIEDENDYTL	GGGMNSQLNQ	KNSHFSLEYA	MNSFDLSHIN	HSISLGSVFQ	420
HTQYRFHRES	DAEAEIINRI	DLENEKIEIK	SSNLAKKGTV	KTRYQNIALY	VEDLMTWKNL	480
EFFRAGLRLER	DDYKLKNTNL	PRTVFRYKPF	EDTAFSVGN	RYYGRSFASM	KLSEGIFKLD	540
GHDTFRYKPLSDE	INTQAPWELG	LSFGVQEYVA	NLAFHLKYIL	RDNKQRIVLQ	EEDVMLNGEK	600
KKLRYYQRGK	DYKTNVLTQ	PTRWTSAVAF	DWLDSKAIDH	GRGYNGSTPV	660	
ILDGKLMTYE	QMLKKVNAK	ETWGLRLNLD	MFVPRFDLW	ANTIYKPPT	TTERVSSNT	720
PEVYRSYDYG	TYTQWDTSLR	WQPTFAEKHR	PYIKLVDLNV	LNKTRKGAGP	NGQDLGIYTP	780
GREFWLEVGY	EF					792

SEQ ID NO: 20                    moltype = AA    length = 794

FEATURE                         Location/Qualifiers  
source                         1..794  
mol\_type = protein  
organism = Pasteurella multocida

SEQUENCE: 20

MESAKNPLKK	TTLALLCCST	AFSLSAKTD	DSDKNHFLTE	IVVYADQNK	MSSTQSVTQD	60
DMKSPVTNG	NITDYLRSNP	HVRYENSQDN	GLQRGEIKPE	NISINGADYQ	QTTFFVDNVN	120
INNDMGMGSD	LFDGTMATVP	FANHSQGYFF	DANLLSSIVV	HDSNVASALG	GFAGGA	180
TKQYDGKDRL	KFSYRTTDAS	WAKFKVEDKD	LERFKNAIPE	GSVAEFQPKY	SKHFFNITAE	240
KGLSENLMGV	IGLSRRTSS	QOSRQINPOQ	DRDKQTHTRR	SDNALLNFNL	TPNDKHRFEL	300
GFRYSNRYR	KFNTNIDSN	VFDYHRAYGV	TFSWINALQ	GILTTTLAYD	NFDDTRKSAS	360
TYMKTTLTDD	GEETYTEGGMG	NSQLNQKNLH	TSLEYAMNP	NLGSIEHSVS	LGGIYQATKY	420
RFTRHSDAVG	ELYTPNWL	NSNDIYDEL	LAQRNIACKG	TVKTRYQ	LYVEDLMTWK	480
NLEFRAGLRL	ERDDYKNTN	LAPRFTVFRYK	PFEDTAFS	VNRYYGRSFA	SMKLSEGIFK	540
LDGHDTFRYK	DLSQFKTPYS	DELSFGVQEY	VANLAFHLKY	ILHDNKQRIV	LQEEDEVMLN	600
EKKKLRYYQR	GKDYKTNVLT	FQISTQAPWE	FBGTRWTSAL	AFDWLDSKAI	DHGGRYNGST	660
PVILDGKLM	YEQMLKKVNA	YKETWGLRLN	LDMFVPKFDL	SWANTIVK	PTTLTERVSS	720
NTPEVYRSYD	YGTYTQWDT	LRWQPTFAEK	HRPYIKLDSL	NVLNKTRKG	GPNGQDLGIY	780
TPGREFWLEV	GYEF					794

SEQ ID NO: 21                    moltype = AA    length = 790

FEATURE                         Location/Qualifiers  
source                         1..790  
mol\_type = protein  
organism = Pasteurella multocida

SEQUENCE: 21

MESAKNPLKK	TTLALLCCST	AFSLSAKTD	NADKNHFLTE	IVVYADQNK	MSSTQSVTQE	60
DMKSPVTNG	NITDYLRSNP	HVRYENSQDN	GLQRGEIKPE	NISINGADYQ	QTTFFVDNVN	120
INNDMGMGSD	LFDGTMATVP	FANHSQGYFF	DANLLSSIVV	HDSNVASALG	GFAGGA	180
TKQYDGKDRL	KFSYRTTDAS	WAKFKVEDKD	LERFKNAIPE	GSVAEFQPKY	SKRFFNITAE	240
KGLSENLMGV	IGLSRRTSS	QOSRQINPOQ	DRDKQTHTRR	SDNALLNFNL	TPNDKHRFEL	300
GFRYSNRYR	KFNTNIDSN	VFDYHRAYGV	TFSWINALQ	GILTTTLAYD	NFDDTRKSAS	360
TYMKTTLEE	GEETYTEGGMG	NSQLNQKNLH	TSLEYAMNP	NLGSIEHSVS	LGGIYQATKY	420
RFTRHSDAVG	ELYTPDWL	NTDKLILTQR	NIAKKGT	RYQNIALYVE	DLMTWKNLEF	480
RAGLRLERDD	YLNKNTNLAPR	TVFRTYKPF	TAFPSVGNR	YGRSFAS	MKL SEGIFKLDGH	540

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SEQ ID NO: 22	moltype = AA length = 790
FEATURE	Location/Qualifiers
source	1..790
	mol_type = protein
	organism = <i>Pasteurella multocida</i>
SEQUENCE: 22	
MSAKNPLKK TTLALLCCST AFSLSAKTD NADKNHFLTE IVVYADQNKS MSSTQSVTQD	60
DMKSPVTNG NTIDYLRSNP HVRYENSDON GLQRGEIKPE NISINGADYQ QTTFVVNDVN	120
INNDMGGFSD LFDGTMATVP FANHSQGYFF DANLSSIVV HDSNVASALG GFAGGAVVAK	180
TQYDGDKRL KFSYRTTDAS WAKFKVEDKD LERFKNAIPE GSVAEFQPKY SKHFFNITAE	240
KGLSENLMGV IGLSRRTSSI QOSRQINPQG DRDKQTHTRR SDNALLNFLN TPNDKHREFL	300
GFFYSNRYER KFNTNIDSN VFDDYHRAYGV TFSWINALQS GILTTLAYD NFDDTRKSAS	360
TYMKTTLEE GEEYTEGGMG NSQLNQKNLN TSLEYAMNPF NLGSIEHSVS LGGIYQATKY	420
RFTRHSDAVG ELYTPDWLN TNDKLILTOR NIAKKGTWKT RYQNIALYVE DLMTWKNLDF	480
RAGLRLERDD YLKNTNLAPR TVFRYKPFED TAFSGWNR YGRSFASML SEGIFKLDGH	540
DTRFYKDLSQ FKTPYSDELS FGVEQYVANL AFHLKYILRD NKQRIVLQEE DVMLNGEKKK	600
LRYYQRGKDY KTNVLTFQIN TQAPWELGPT RWTSAVAFDW LDSKAIDHGR GYNQSTPVIL	660
DGKLMTYEQM LKKVNAVYKET WGLRLNLDMP VPRFDLSWAN TIYVKPPTL TEHVSSSTPE	720
VYRSYDYGTY TQWDTSLRWQ PTFAEKHRPY IKLDVLNVLN KTRKGAGPNG QDLGIYTPGR	780
EFWLEVGYEF	790
SEQ ID NO: 23	moltype = AA length = 730
FEATURE	Location/Qualifiers
source	1..730
	mol_type = protein
	organism = <i>Pasteurella multocida</i>
SEQUENCE: 23	
MSFKHKTAL FVAHACCTSV LAENAATTLE PIVVSDLSTSHT TLNLQDNKLE KESPKDLKAI	60
FATTPNINVII HTGHAQLGDI EIRGMGSSRE IFATGANRVT MELDGMDISP SFYFGHSSRH	120
GRQYFDPSDL KRVEVHKGPN SQGVAGHVRF QTKDPHDYLL PNQRTGAQLR AGYLGDSDAY	180
YVGITGAALL DEHSSALVSY TRRWFNEFNN KGGLDVTSQ RTKSNPSSGY SNAVNSKLRY	240
SPNDRHKFTL NLQHYDLKRT AYLEDLSGTT TTRRGTKTVH HNTNIQKNQR HAIAPSHDMQ	300
QTNHISIFDHL HWQIALQQTQ STSRNTGAVT STSGSPPPST PKFSQERSLD GFKTKTISLK	360
TEFNKSLQHQH VVHELHYGLK LQYSQMLR QTQSLNEQGS NRTSAAFFPT QQWQSKLHL	420
SDRISFGKSG LSLTPSIHLT QIRIKPKTEN VSKKNREQLF TYKDTAIGYG LRVDYALNEA	480
NLLSLNYQHA TRLPGYGENN AQSYGHWPWPK PNPHLQPETS DGIELSWSRA GAIGQQTTTL	540
FYNRYNDLIY DTTTACYADR TGQVQPCDL ANEKGRSYSY GIEFDGKLNL DTIGFAQGTY	600
LNAGFAYSKGK TANKQPOGR LDPLTGFGVGL GYQQPMWDWV IEGLKLFAAK KKTDL PANQ	660
GFEGGLPGYAV VDLTAYYNTV KQLYLGIGIY NVLDKKYARW AMARGDIKHG NYDKHTEAGR	720
HFGANIRYHF	730
SEQ ID NO: 24	moltype = AA length = 727
FEATURE	Location/Qualifiers
source	1..727
	mol_type = protein
	organism = <i>Pasteurella multocida</i>
SEQUENCE: 24	
MSFKHKTAL FVAHACCTSA LAENAATTLE PIVVSELSHT TLNLQDNKLE KESPKDLKAI	60
FTTTPNINVII HTGHAQLGDI EIRGMGSSRE IFATGANRVT MELDGMDISP SFYFGHSSRH	120
GRQYFDPSDL KRVEVHKGPN SQGVAGHVRF QTKDPHDYLL PNQRTGAQLR AGYLGDSDAY	180
YVGITGATLL DEHSSALVSY TRRWFNEFNN KGGLDVTSQ RTKSNPSSGY SNAVNSKLRY	240
SPNDRHKFTL NLQHYDLKRT AYLEDLSGTT TTRRGTKTVH HNTNIQKNQR HAIAPSHDMQ	300
QTAAFFDHHL WQIALQQTQ TSRRNTGTVTS TSAPPAPPSTP KFSQERSFDG FKTKTISLK	360
EFPNLSQHGV VVHELHYGLK QYSQMLRQ TQSLNEQGSN QQWQSKFHLS	420
DRISFGKSG SLTPSIHLT QIRIKPKTEN SKKNREQLFT YKDTAIGYGL RVDYALNEA	480
LLSLNYQHAT RLPGYGENNA QSYGHWPWPK NPHLQPETS GVELSWRSAG AIGQQTTLF	540
YNRYNDLIY DTTTACYADR GTQVPCDLANE KGRSYSYGIE FDGKLNLDTI GFAQGTYLNA	600
GFAYSKGKTA NKQPQGRLLDP LTGFVGLGYQ QPMDVWGLIEG KLKFAAKKKT KDL PANQ GFE	660
GLPGYAVVDSL TAYYNTVKQL YLGIGIYNL DKKYARWAMA RGDIKHGNYD KHTEAGRHF	720
ANIRYHF	727
SEQ ID NO: 25	moltype = AA length = 727
FEATURE	Location/Qualifiers
source	1..727
	mol_type = protein
	organism = <i>Pasteurella multocida</i>
SEQUENCE: 25	
MSFKHKTAL FVAHACCTSA LAENVATTLE PIVVSDLSTSHT TLNLQDNKLE KESPKDLKAI	60
FATTPNINVII HTGHAQLGDI EIRGMGSSRE IFATGANRVT MELDGMDISP SFYFGHSSRH	120
GRQYFDPSDL KRVEVHKGPN SQGVAGHVRF QTKDPHDYLL PNQRTGAQLR AGYLGDSDAY	180

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YVGITGATLL DEHSSALVSY TRRWLFNEFNN KGGLDVTGSQ RTKSNPSSGY SNAVNSKLRY	240
SPNDRHKFTL NLQHYDLKRT AYLEDLSGTT TTRRGTKTVH HNTNIQKNQR HAIAFSDMQ	300
QTAAFFDHHL WQIALQQTKS TSRNTGAVTS TSASPPPSTP KFSQERSFDG FKTKTISLKT	360
EFNKSLQHQH VHELYHGLKL QYSQMQLRQ TQSLNEQGSN TRTSAFFPTQ QQWQSKLHLS	420
DRISFGKSGL SLTPSIHLTQ IRIKPTEVN SKKNREQLFT YKD'TAIGYGL RVDYALNEAN	480
LILSINYOHAT RLPGYGENNA QSYGHWPAPK NPHLQPETSD GIELSWRSAG AIGQOTTTLF	540
YNRYNDLIYL DTTACYADRT GQVPCDLANE KGRSYSYGIE FDGKLNLDTI GFAQGTYLNA	600
GFAYSKGKTA NKQPQGRLEP LTGFVGLGYQ QPMDVWGIEG KLKFAAKKKT KDL PANQGFE	660
GLPGYAVV р TAYYNVTKQL YLGIGIYNVL DKKYARWAMA RGDIKHGNYD KHTEAGRHF	720
ANIRYHF	727

SEQ ID NO: 26	moltype = AA length = 727
FEATURE	Location/Qualifiers
source	1..727
	mol_type = protein
	organism = Pasteurella multocida
SEQUENCE: 26	
MSFKHKTLAL FVAHACCTSA LAENAATTLE PIVVSDLSHT TLNLDQNKE KESPKDLKAI	60
FATTPNINV1 HTGHAQLGDI EIRGMGSSRE IFATGANRVT MELEGMDISP SFYFGHSRH	120
GROYFDPSDL KRVEIHKGPN SQGVAGHVRP QTQDPRDYL PNQRTGAQLR AGYLGDSAY	180
YVGITGATLL DEHSSALMSY TRRWLFNEFNN KGGLDVTGSQ RTKSNPSSGY SNAVNSKLRY	240
SPNDRHKFTL NLQHYDLKRT AYLEDLSGTT TTRRGTKTVH HNTNIQKNQR HAIAFSDMQ	300
QTAAFFDHHL WQIALQQTKS TSRNTGAVTS TSASPPPSTP KFSQERSLDG FKTKTISLKT	360
EFNKSLQHQH VHELYHGLKL QYSQMQLRQ TQSLNEQGSN TRTSAFFPTQ QQWQSKLHLS	420
DRISFGKSGL SLTPSIHLTQ IRIKPTEVN SKKNREQLFT YKD'TAIGYGL RVDYALNEAN	480
LILSINYOHAT RLPGYGENNA QSYGHWPAPK NPHLQPETSD GIELSWRSAG AIGQOTTTLF	540
YNRYNDLIYL DTTACYADRT GQVPCDLANE KGRSYSYGIE FDGKLNLDTI GFAQGTYLNA	600
GFAYSKGKTA NKQPQGRLEP LTGFVGLGYQ QPMDVWGIEG KLKFAAKKKT KDL PANQGFE	660
GLPGYAVV р TAYYNVTKQL YLGIGIYNVL DKKYARWAMA RGDIKHGNYD KHTEAGRHF	720
ANIRYHF	727

SEQ ID NO: 27	moltype = AA length = 964
FEATURE	Location/Qualifiers
source	1..964
	mol_type = protein
	organism = Pasteurella multocida
SEQUENCE: 27	
MRTTIKFS A ITLALLSYCG VILADSHQEA TELDTITVSS QQDEMNKEK KVGETVKTAS	60
QLKRQQVQDS RDLVRYETGV TVVEAGRFGS SGYAIRGVDE NRVAITVDGL HQAETLSSQG	120
FKELFEGYGN FNNTTRNSVEI ETLVAKIAK GADSVKVGS SLGGAVLFET KDARDFLTEK	180
DWHIGYKAGY STADNQGLNA VTLAGRYQMF DALIMHSKRH GHELENYDYK NGRDIQGKER	240
EKADPYTITK ESTLVKFSFS PTENHRTVA SDTYIQRSRG HDRSYSLQPQ SNYFTYDEKE	300
SRHANDLTKR KNVSFTYENY SVTPFWDTLK LSYSQQKIRT RARTEDYCDG NEKCDSYKNP	360
LGLQLKDGKI VDPAQNQITL KGTGFNTIEV DKNGQNPFPST SGTNNAAFSN NIQLGPKEFW	420
LDCLSLFDCTQ PFTVNYQNG QTTPKQVELS EEITVNGKLY KTAKEERGVR NYLILPNSKG	480
YLPYDYLKEDR LDSNTKQINL DLTKTFTSFN IENELLYGAI YSRTEKKMVN KAGYDGRNPT	540
WWADRILGKS TNCNYNGLKC PRHEPLTSFL IPVEATTKSL YFSDN1KLHN MLSVDSLGYRY	600
DDIKYQPEYI PGVTPKIAAD MVKGIFIPLP KGEKVTTPW AETTKPLTQE QIRKNAEENI	660
AYIAQEKRKFK HSYSLGATF DPLNFLRVQV KYSKGFRAPT SDELYFTFKH PDFTILPNPV	720
LKPEEAKNQE IALTVDNNG FVTSVFQTKY YRFIDLAYL GSRNLSNSVG GQAQARDFQV	780
YQNVNVDNAK VKGLEINARL NLGYFWHVLDF GFNTSYKFTY QRGRLDGDRP MNAIQPKASV	840
FGLGYDHKENK FGADLYITR VSEKKAKDTY NMFYKEQGYKD SAIRWRSSDD YTLDAVGYI	900
KPIKNLTLQF CVYNLTDKLTWESARTSK PFGETSMLNQ KTGAGINRFV SPGRNFKFS	960
ETIF	964

SEQ ID NO: 28	moltype = AA length = 963
FEATURE	Location/Qualifiers
source	1..963
	mol_type = protein
	organism = Pasteurella multocida
SEQUENCE: 28	

MRTTIKFS A ITLALLSYCG AILADSHQEA TELDTITVSS QQDEMNKEK KVGETVKTAS	60
QLKRQQVQDS RDLVRYETGV TVVEAGRFGS SGYAIRGVDE NRVAITVDGL HQAETLSSQG	120
FKELFEGYGN FNNTTRNSVEI ETLVAKIAK GADSVKVGS SLGGAVLFET KDARDFLTEK	180
DWHIGYKAGY STADNQGLNA VTLAGRYQMF DALIMHSKRH GHELENYDYK NGRDIQGKER	240
EKADPYTITK ESTLVKFSFS PTENHRTVA SDTYIQRSRG HDRSYSLQPQ SNYFTYDEKE	300
SRHANDLTKR KNVSFTYENY SVTPFWDTLK LSYSQQKIRT RARTEDYCDG NEKCDSYKNP	360
LGLQLKDGKI VDPAQNQITL KGTGFNTIEV DKDGKPFPTT SGTNNAAFSN NLQLSPTGFW	420
LDCTIFDCTQ PFTVNYQNG KYEPREVMLS EEITIDGKLY KTAKEESGVR NYLILPNSKG	480
YLPYDYLKEDR LDSNTKQINL DLTKTFTSFN IENELLYGAI YSRTEKKMVN KAGYDGRNPT	540
WWADRILGKS TNCNYNGLKC PRHEPLTSFL IPVEATTKSL YFSDN1KLHN MLSVDSLGYRY	600
DDIKYQPEYI PGVTPKIAAD MVKGIFIPLP KGEKIKIENY ETTKPLTPEQ IRKNAEENIA	660
YIAQEKRKFK HSYSLGATF PLNFLRVQV YSKGFRAPTS DELYFTFKHP DFTILPNPV	720
KPEEAKNQE IALTVDNNG FVTSVFQTKY RHFIDLAYL SRNLSNSVG QQAQARDFQVY	780
YQNVNVDNAK VGLEINARL LGYFWHVLDF FNTSYKFTY QRGRLDGDRP NAIQPKASV	840
GLGYDHKENK FGADLYITR VSEKKAKDTY MFYKEQGYKD SAIRWRSSDD YTLDAVGYI	900

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PIKNLTLQFG VYNLTDRKYL TWESARSIKP FGTSNLINQK TGAGINRFYS PGRNFKFSAE ITF	960 963
SEQ ID NO: 29	moltype = AA length = 963
FEATURE	Location/Qualifiers
source	1..963
	mol_type = protein
	organism = Pasteurella multocida
SEQUENCE: 29	
MRTTTIKPSA ITLALLSYCG AILADSHQEA TELDTITVSS QDEEMNIKEK KVGETVKTAS QLKRQQVQDS RDLVRYETGV TVVEAGRFGS SGYAIRGVDE NRVAITVDGL HQAETLSSQG FKELFEGYGN FNNTNRNSVEI ETLKVAKIAK GADSVKVGSS SLGGAVLFET KDARDFLTEK DWHIGYKAGY STADNQGLNA VTLAGRYQMF DALIMHSKRH GHELENYDYK NGRDIQGKER EKADPYITK ESTLVKFSFS PTENHFRFTVA SDTYIQRSRG HDRSYSLQPQ SNYFTYDEKE SRHANDLTKR KNVSFTYENY SITPFWDTLK LSYSQQKIRT RARTEDYCDG NEKCDSYKNP LGQLQKDGKI VDPAGNQITL KGTGFNTIEV DKDGKPFPTT SGTNNAAFSI NLQLSPTEFW LDCTIFDCTK PFTVYNYKQN KYEPREVMLS EEITIDGKLK KTAKEESGVR NYLILPNNSKG YLPYDYKERD LDSNTKQINL DLTKTFSTFN IENELLYGAI YSRTEKKMVN KAGYDGRNPT WWADRILKRN TNCNYNGLKC PRHEPLTSFL IPVEATTKSL YFSDNIKLHN MLSVDLGYRY DDIKYQEPEI PGVTPKIADD MVKGIFIPLP KGEKIKIGNY ETTPKPLTPEQ IRKNAEENIA YIAQEKRFKK HSYSLGATFD PLNFLRVQVK YSKGFRAPTS DELYFTFKHP DFTILPNPV KPEEAKNQEI ALTVHNDWGF VSTSVFQTKY RHFIDLAYLG SRNLSNSVGG KAQARDFQVY QNVNVDAKVN KGVEINARLN LGYFWHVLDG FNTSYKFTYQ RGRLDGDRPM NAIQPKASVF GLGYDHKENK PGADLYITRV SEKKADTYN MFYKEQGYKD AIRWRSSDDY TLIDAVGYIK PIKNLTLQFG VYNLTDRKYL TWESARSIKP FGTSNLINQK TGAGINRFYS PGRNFKLSAE ITF	60 120 180 240 300 360 420 480 540 600 660 720 780 840 900 960 963
SEQ ID NO: 30	moltype = AA length = 971
FEATURE	Location/Qualifiers
source	1..971
	mol_type = protein
	organism = Pasteurella multocida
SEQUENCE: 30	
MRTTTIKPSA ITLALLSYCG VILADSHQEA TELDTITVSS QDEEMNIKEK KVGETVKTAS QLKRQQVQDS RDLVRYETGV TVVEAGRFGS SGYAIRGVDE NRVAITVDGL HQAETLSSQG FKELFEGYGN FNNTNRNSVEI ETLKVAKIAK GADSVKVGSS SLGGAVLFET KDARDFLTEK DWHIGYKAGY STADNQGLNA VTLAGRYQMF DALIMHSKRH GHELENYDYK NGRDIQGKER EKADPYITK ESTLVKFSFS PTENHFRFTVA SDTYLQHRSRG HDLSYNLVAT TYIQLDEKES RHANDLTKRN NUSFTYENY VTPVDFDTKL SYSQQRITTR ARTEDYCDGN EKCDSYKNP GLQLKEKGIV DRNGDPVNLQ LVDGKHQVVD KAGKPFWDTSF GTNYAAFSGK QLGPSYFWLE CTVFDCSKPV TTYKVYRSTE TPVKEDIQLN KTMEVNGKTF ATYDMGRERR YIILPNSQGY LPLDYKERDL NTKTQKINLD LTKAFTLFEI ENELSYYGGVY AKTTNGVVK AGYYGRNPTW WAERTLQGSW NGTLRECKTS SSYNGMLCPR HEPLTSFLIP VEATTKSLYF ADNIKLNHNM SVDLGYQYDD IKYQPEYIPG VTPKIAADDMV KGLFIPLPVE EKVTVGTVVF TKPLTPEQIR KNAEENIAYI AQGKRFKKHS YSLGTTFDPL NFRLRVQVKY KSFRAPTSDE LYFTFKHPDF TILPNPVLKPEEAKNQEIAL TVHDNNGFVS TSVFQTKYR FIDLAYLGSR NLSNSVGGQA QARDFQVYQN VNVDNAKVKG LEINARLNIG YFWHVLDFN TSYKFTYQRG RLDGDRPMNA IQPKASVFGK GYDHKENKPG ADLYITRVSE KKAADTYNMF YKEQGYKDSEA IRWRSSDDYTL VDAVGYIKPI KNLTQFGVY NLTERKYLTW ESARSIKPFG TSNLINQKTG AGINRFYSPG RNFKFSAEIT F	60 120 180 240 300 360 420 480 540 600 660 720 780 840 900 960 971
SEQ ID NO: 31	moltype = AA length = 848
FEATURE	Location/Qualifiers
source	1..848
	mol_type = protein
	organism = Pasteurella multocida
SEQUENCE: 31	
MQKSQPYPIH LGIFFLMLGLP TWAFSQANLE KSTINKLETI LVNESEEKNK FDENLIKTYL SSGSYSYLSQ SDISTFRGSS VGDPLSGVPG VIVGNKRNSG ALSVNLIRGIA NENRVPVWID KGLQSVPYSQ IEKGPQLSQM ATGATGGVVR VETLRWQDII PQGNKLGVRKL GLGTMTNTVS PPPYTYRGGY QTQYISKCLS NHTGLCQTQ YAPNARYSSH GFDLNAYNYS LAFANKWSNA DLVLAYAKRK QGNYFVGRHG QTPVIESIEF EEDSVEVKEP RVHEDVEIGS LTFKENRSTL YRPGEEARLNT SQDNTSYLAQ INVYNDVHRL GLAYRHYHSR FGEIMMSILN FRAYGALQGE GTEVKVDSYH ANYSYNPFTT YVNLNVNAYE TDSDSNFTP FIEEYGYSLS SRHAHFLVSK QKGLSIENT IFQLNDKPFY LKYGLAHSYE RIYQPRNAQA RVRAKGYPED AIGPLYIRDG KRKEWSAFVA ANYPITSWLK ADIGLYLQS TIYDYIVRTE RVNIGGALVP NPNGSGNIWV EKYKDVVHQ APVKNKGMP IVMFTFEPIN GVQIYTKYAE ALRSPSLFQA TKGWSMSATE DNLEQLRPER AKNWEAGINL FYENLGGKDN ILGFKLAYFN NRIKDYLTRS YSPKDKVTQT INIQSAQFKG IELSAYYDMG KFYAKLAGTY YTAKTFCLTA EQAGKGEQCN SGYVYRSNLN NAVPPRLNLH ATLGTRLFEO KLDIGARYSY YSKRLLPVLS AERFVNNTSSI EWAPYSLVLDL YANYVNSNNL KLTMTMDNFV NRYYLDINNM GLNTAPGRTL HHLGLEYRF	60 120 180 240 300 360 420 480 540 600 660 720 780 840 848
SEQ ID NO: 32	moltype = AA length = 848
FEATURE	Location/Qualifiers
source	1..848

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mol_type = protein
organism = Pasteurella multocida

SEQUENCE: 32
MQKQQPYPPIH LGIFLMLGLP TWAFSQANLE KSTINKLETTI LVNESEEKNK FDENLIKTYL 60
SSGSYSYLSQ SDISTFRGSS VGDFLSGVPG VIVGNKRNSG ALSVNIRGIA NENRVPVWID 120
KGLQSVPSYQ GYAGGSSTRY LDPDLISQVE IEKGPSLQMD ATGATGGVVR VETLRWQDII 180
PQGKNNWGVRL KLGTMTNTVS PPPYYTRGGY QTKYISKCLS NDTGLCQTQT YAPNARYSSH 240
GFDLNAYNYS LAFANKWQNA DLVLAYAKRK QGNYFVGRHG QTPVIESIEF EEDSVEVKEP 300
RVHEDVEIGS LTFKENRSTL YRPGEALNT SQDNTSYLAK INVYNDVHRL GLAYRHYSR 360
FGEIMSSILN PRAYGALQGE GTEVKVDSYH ANYSYNPTTP YVNLNVNAYF TDSDSSNFTP 420
FIEEYGYSLN SRHAHFLVSK QKGLSIENTS IFQLNDKPFT LKYGLAHSYE RIYQPRNAQA 480
RVRAKGYPED AIGPLYIRDG KRKEWSAFVA ANYPITSWLK ADIGLRYLQS TIYDYIVRTE 540
RVNIGGALVP NPNGSGNIWV EKYKDVVHKQ APVKNKGMSM IVMLTFEPIN GVQIYTKYAE 600
ALRSPSLFQA TKGWSMSATA DNLEQLRPER AKNWEAGINL FYENLGGKDN ILGFKLAYFN 660
NRIKDYLTRS YSPKDKVTQT INIQAQFKG IELSAYYDMG KFYTKLAGTY YTAKFCFLTA 720
EQAGKGEQCN SGYVYRSNLN NAVPPRLNLH ATLGLTRLFEO KLDIGARYSY YSKRLVPVLS 780
AERFVNNTSSI EWAPYSLVLDL YANYNVSNNL KLTMTMDNVF NRYYLDINNT GLNTAPGRTL 840
HLGLEYRF 848

SEQ ID NO: 33      moltype = AA length = 848
FEATURE           Location/Qualifiers
source            1..848
mol_type = protein
organism = Pasteurella multocida

SEQUENCE: 33
MQKQQPYPPIH LGIFLMLGLP TWAFSQANLE KSTINKLETTI LVNESEEKNK FDENLIKTYL 60
SSGSYSYLSQ SDISTFRGSS VGDFLSGVPG VIVGNKRNSG ALSVNIRGIA NENRVPVWID 120
KGLQSVPSYQ GYAGGSSTRY LDPDLISQVE IEKGPSLQMD ATGATGGVVR VETLRWQDII 180
PQGKNNWGVRL KLGTMTNTVS PPPYYTRGGY QTKYISKCLS NDTGLCQTQT YAPNARYSSH 240
GFDLNAYNYS LAFANKWQNA DLVLAYAKRK QGNYFVGRHG QTPVIESIEF EEDSVEVKEP 300
RVHEDVEIGS LTFKENRSTL YRPGEALNT SQDNTSYLAK INVYNDVHRL GLAYRHYSR 360
FGEIMSSILN PRAYGALQGE GTEVKVDSYH ANYSYNPTTP YVNLNVNAYF TDSDSSNFTP 420
FIEEYGYSLN SRHAHFLVSK QKGLSIENTS IFQLNDKPFT LKYGLAHSYE RIYQPRNAQA 480
RVRAKGYPED AIGPLYIRDG KRKEWSAFVA ANYPINSWLK ADIGLRYLQS TIYDYIVRTE 540
RVNIGGALVP NPNGSGNIWV EKYKDVVHKQ APVKNKGMSM IVMLTFEPIN GVQIYTKYAE 600
ALRSPSLFQA TKGWSMSATA DNLEQLRPER AKNWEAGINL FYENLGGKDN ILGFKLAYFN 660
NRIKDYLTRS YSPKDKVTQT INIQAQFKG IELSAYYDMG KFYTKLAGTY YTAKFCFLTA 720
EQAGKGEQCN SGYVYRSNLN NAVPPRLNLH ATLGLTRLFEO KLDIGARYSY YSKRLVPVLS 780
AERFVNNTSSI EWAPYSLVLDL YANYNVSNNL KLTMTMDNVF NRYYLDINNM GLNTAPGRTL 840
HLGLEYRF 848

SEQ ID NO: 34      moltype = AA length = 848
FEATURE           Location/Qualifiers
source            1..848
mol_type = protein
organism = Pasteurella multocida

SEQUENCE: 34
MQKQQPYPPIH LGIFLMLGLP TWAFSQANLE KSTINKLETTI LVNESEEKNK FDENLIKTYL 60
SSGSYSYLSQ SDISTFRGSS VGDFLSGVPG VIVGNKRNSG ALSVNIRGIA NENRVPVWID 120
KGLQSVPSYQ GYAGGSSTRY LDPDLISQVE IEKGPSLQMD ATGATGGVVR VETLRWQDII 180
PQGKNNWGVRL KLGTMTNTVS PPPYYTRGGY QTKYISKCLS NDTGLCQTQT YAPNARYSSH 240
GFDLNAYNYS LAFANKWQNA DLVLAYAKRK QGNYFVGRHG QTPVIESIEF EEDSVEVKEP 300
RVHEDVEIGS LTFKENRSTL YRPGEALNT SQDNTSYLAK INVYNDVHRL GLAYRHYSR 360
FGEIMSSILN PRAYGALQGE GTEVKVDSYH ANYSYNPTTP YVNLNVNAYF TDSDSSNFTP 420
FIEEYGYSLN SRHAHFLVSK QKGLSIENTS IFQLNDKPFT LKYGLAHSYE RIYQPRNAQA 480
RVRAKGYPED AIGPLYIRDG KRKEWSAFVA ANYPITSWLK ADIGLRYLQS TIYDYIVRTE 540
RVNIGGALVP NPNGSGNIWV EKYKDVVHKQ APVKNKGMSM IVMLTFEPIN GVQIYTKYAE 600
ALRSPSLFQA TKGWSMSATA DNLEQLRPER AKNWEAGINL FYENLGGKDN ILGFKLAYFN 660
NRIKDYLTRS YSPKDKVTQT INIQAQFKG IELSAYYDMG KFYTKLAGTY YTAKFCFLTA 720
EQAGKGEQCN SGYVYRSNLN NAVPPRLNLH ATLGLTRLFEO KLDIGARYSY YSKRLVPVLS 780
AERFVNNTSSI EWAPYSLVLDL YANYNVSNNL KLTMTMDNVF NRYYLDINNM GLNTAPGRTL 840
HLGLEYRF 848

SEQ ID NO: 35      moltype = AA length = 784
FEATURE           Location/Qualifiers
source            1..784
mol_type = protein
organism = Pasteurella multocida

SEQUENCE: 35
MKYPLSYKNI ARSIPFLSFI APPLYAQETT ELEQITVQES ATAEVNKTPS TVISKSATTI 60
QNKMIRDTRD LVRYTTDVGI SDNGRFLKGF AMRGVEDNVR GISIDGVSLP DSEENSLYAR 120
YGNFNNSRSLN IDPELIQTID IVRGSDSFNA GSGSLGGGVN YNTLDPQHIV KAGNSVGALL 180
RGSYASKNRE WVRTLGIGYV GEKFADLLMY SQTGHEPKS RGSGPPEFRYS SSQHDPVTQ 240
RFHNYLAKMN YQINDNQRIG LTLNQGTGGR YIDERSYLM GSQWREADDQ NERLNANLYY 300
IYAPSTGWLA YSKFDLDYQK TDLAAVNYKG GRHFTTDAKE LNEIYDRRMK TVFTRGSVEL 360
NAQPVHFYGE HTLTIKGYVS QRDFKNNINQD RIGIGTNYDT QYHYTIQYPI RTKQYGLSLK 420

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DHVRWNDTFS	SHLGLRYDHT	KLKPKELNAP	CSKACLEEGK	PKPTRFSTVS	TFAGLEAQLS	480
PSWMLGYNIS	TGYRVP TASE	MFFSFTNAYG	TWKS NPSLKP	EKSINHTLSL	KGNSEKGLLD	540
LTLYQTNYRH	PLFEQESLIQ	RTEIRYQPY	TYQSQEQQMV	NLDKAKIYGV	ELKTHVNLDQ	600
MIAVIPQGFK	FYAALGYSKG	KLSNNASLLS	IOPLKIIILGL	DYEATNGKWA	IFNRLTYLGE	660
KRASDAKVEE	I KRRCTEFVT	ETDPWSGEQI	THCKKRELYP	DLSTYKHLNK	SAFVFD TFGY	720
YKITDDITFR	AGIYNLFNKK	YHTWDALRG	IANSTLNSVD	REGKGLQRFY	APGRNYAASL	780
EIRF						784

SEQ ID NO: 36	moltype = AA	length = 784				
FEATURE	Location/Qualifiers					
source	1..784					
	mol_type = protein					
	organism = Pasteurella multocida					
SEQUENCE: 36						
MKYPLSYKNI	ARSIPFLSFI	AFPLYA QETT	ELEQITVQES	ATAEVNK TSP	TVISK SATTI	60
QNEMIRDTRD	LVRYTTDVGI	SDNGRFLKGF	AMRGVEDNRV	GISIDGVSLP	DSEENSLYAR	120
YGNFNNSRSL	IDPELIQTID	IVRGSDSFNA	GSGSLGGGVN	YNTLDPQHIV	KAGNSVGALL	180
RCGSYASKNRE	WVRTLGIGYV	GEKFDALLMY	SORTGHEFKS	RGSGP EFRYS	SSQHPDPVTQ	240
RFHNYLAKMN	YQINDNQRIG	LTLNGQTGGR	YIDERSYTLIM	GSQWREADDQ	NERLNANLYYY	300
IYAPSTGWLA	YSKF DLDYQK	TDLA AVNYKG	GRHFTTDAKE	LNEIYDRRMK	TVFTRGSVEL	360
NAQPVHFYGE	HTLT IKG YVS	QRDFKNINQD	RIGIGTNYDT	QYHYTIQYPI	RTKQYGLSLK	420
DHVRWNDTFS	SHLGLRYDHT	KLKPKELNAP	CSKACLEEGK	PKPTRFSTVS	TFAGLEAQLS	480
PSWMLGYNIS	TGYRVP TASE	MFFSFTNAYG	TWKS NPSLKP	EKSINHTLSL	KGNSEKGLLD	540
LTLYQTNYRH	PLFEQESLIQ	RTEIRYGRPY	TYQSQEQQMV	NLDKAKIYGV	ELKTHVNLDQ	600
MIAVIPQGFK	FYAALGYSKG	KLSNNASLLS	IOPLKIIILGL	DYEATNGKWA	IFNRLTYLGE	660
KRASDAKVEE	I KRRCTEFVT	ETDPWSGEQI	TRCKKRELYP	DLSTYKHLNK	SAFVFD TFGY	720
YKITDDITFR	AGIYNLFNKK	YHTWDALRG	IANSTLNSVD	REGKGLQRFY	APGRNYAASL	780
EIRF						784

SEQ ID NO: 37	moltype = AA	length = 784				
FEATURE	Location/Qualifiers					
source	1..784					
	mol_type = protein					
	organism = Pasteurella multocida					
SEQUENCE: 37						
MKYPLSYKNI	ARSIPFLSFI	AFPLYA QETT	ELEQITVQES	ATAEVNK TSP	TVISK SATTI	60
QNEMIRDTRD	LVRYTTDVGI	SDNGRFLKGF	AMRGVEDNRV	GISIDGVSLP	DSEENSLYAR	120
YGNFNNSRSL	IDPELIQTID	IVRGSDSFNA	GSGSLGGGVN	YNTLDPQHIV	KAGNSVGALL	180
RCGSYASKNRE	WVRTLGIGYV	GEKFDALLMY	SORTGHEFKS	RGSGP EFRYS	SSQHPDPVTQ	240
RFHNYLAKMN	YQINDNQRIG	LTLNGQTGGR	YIDERSYTLIM	GSQWREADDQ	NERLNANLYYY	300
IYAPSTGWLA	YSKF DLDYQK	TDLA AVNYKG	GRHFTTDAKE	LNEIYDRRMK	TVFTRGSVEL	360
NAQPVHFYGE	HTLT IKG YVS	QRDFKNINQD	RIGIGTNYDT	QYHYTIQYPI	RTKQYGLSLK	420
DHVRWNDTFS	SHLGLRYDHT	KLKPKELNAP	CSKACLEEGK	PKPTRFSTVS	TFAGLEAQLS	480
PSWMLGYNIS	TGYRVP TASE	MFFSFTNAYG	TWKS NPSLKP	EKSINHTLSL	KGNSEKGLLD	540
LTLYQTNYRH	PLFEQESLIQ	RTEIRYGRPY	TYQSQEQQMV	NLDKAKIYGV	ELKTHVNLDQ	600
MIAVIPQGFK	FYAALGYSKG	KLSNNASLLS	IOPLKIIILGL	DYEATNGKWA	IFNRLTYLGE	660
KRASDAKVEE	I KRRCTEFVT	ETDPWSGEQI	TRCKKRELYP	DLSTYKHLNK	SAFVFD TFGY	720
YKITDDITFR	AGIYNLFNKK	YHTWDALRG	IANSTLNSVD	REGKGLQRFY	APGRNYAASL	780
EIRF						784

SEQ ID NO: 38	moltype = AA	length = 784				
FEATURE	Location/Qualifiers					
source	1..784					
	mol_type = protein					
	organism = Pasteurella multocida					
SEQUENCE: 38						
MKYPLSYKNI	ARSIPFLSFI	AFPLYA QETT	ELEQITVQES	ATAEVNK TSP	TVISK SATTI	60
QNEMIRDTRD	LVRYTTDVGI	SDNGRFLKGF	AMRGVEDNRV	GISIDGVSLP	DSEENSLYAR	120
YGNFNNSRSL	IDPELIQTID	IVRGSDSFNA	GSGSLGGGVN	YNTLDPQHIV	KAGNSVGALL	180
RCGSYASKNRE	WVRTLGIGYV	GEKFDALLMY	SORTGHEFKS	RGSGP EFRYS	SSQHPDPVTQ	240
RFHNYLAKMN	YQINDNQRIG	LTLNGQTGGR	YIDERSYTLIM	GSQWREADDQ	NERLNANLYYY	300
IYAPSTGWLA	YSKF DLDYQK	TDLA AVNYKG	GRHFTTDAKE	LNEIYDRRMK	TVFTRGSVEL	360
NAQPVHFYGE	HTLT IKG YVS	QRDFKNINQD	RIGIGTNYDT	QYHYTIQYPI	RTKQYGLSLK	420
DHVRWNDTFS	SHLGLRYDHT	KLKPKELNAP	CSKACLEEGK	PKPTRFSTVS	TFAGLEAQLS	480
PSWMLGYNIS	TGYRVP TASE	MFFSFTNAYG	TWKS NPSLKP	EKSINHTLSL	KGNSEKGLLD	540
LTLYQTNYRH	PLFEQESLIQ	RTEIRYGRPY	TYQSQEQQMV	NLDKAKIYGV	ELKTHVNLDQ	600
MIAVIPQGFK	FYAALGYSKG	KLSNNASLLS	IOPLKIIILGL	DYEATNGKWA	IFNRLTYLGE	660
KRASDAKVEE	I KRRCTEFVT	ETDPWSGEQI	TRCKKRELYP	DLSTYKHLNK	SAFVFD TFGY	720
YKITDDITFR	AGIYNLFNKK	YHTWDALRG	IANSTLNSVD	REGKGLQRFY	APGRNYAASL	780
EIRF						784

SEQ ID NO: 39	moltype = AA	length = 745
FEATURE	Location/Qualifiers	
source	1..745	
	mol_type = protein	
	organism = Haemophilus influenzae	

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SEQUENCE: 39

MNIIINKRIF	LLVTFVGIGL	NVTAKQNSSN	SNREELLPII	VNTDEDSNKL	PGRSVLKQKN	60
IEQQQADNAA	NLINLPGVN	MAGGFRPGGQ	TLNINGMGA	EDVRVQLDGA	TKSFEKYQQG	120
SFIEPELLR	KTVVDKGNY	PQYGNGGFAG	TVKFETRDAR	DFLKENQKIG	GLLKVGNNSN	180
NNQKTYSTAL	VLQNEQKNID	LLLFGSVRNA	GDYKRPDNSK	ILFSKNNQKT	GLIKVNWQIT	240
PEHLLTSSLV	YGIHKGWEWP	AAKRDTVSRP	TETEIKRYGI	DVAWKRKLVY	RDQKDESYS	300
KYRLPENNPK	WINLSVQLSY	SKTEQNDTRH	EKVTSFLGT	LGNKSWITYS	DLTFDISNTS	360
TLNIGRAEHE	LLFGLQWLKN	TRNTLHYHK	KMNDKTYNYG	YFQPYMPMSG	RQYQAFYLQ	420
DQIKWKNIIF	STGARYDHIN	NIGQKNLAPQ	YNDISAGHNY	SQKNYNGWSY	YLGLKVDVNH	480
YLSLFTNFNR	TWRAPVIDEQ	YETOQSASV	PATSLNLEKE	MISQTRGGGT	VTLNNNLFQED	540
DTFQFRATYF	YHRGKNEIF	TRGVNCVGN	LDVDNKICPK	IISNYRNLPG	YVIQGAELEA	600
YYOSTYLFGE	LTYSVVKGKR	DTSPRNPWGK	TSTWIAEIP	RKATATLGFN	VPKYLTVGW	660
RAEFVRRQDR	SPSSRDPKAS	YYLSPLASRG	YSLHNLFLSW	TPEKIKGMNI	KITVDNLFNR	720
AYNPYLGELA	SGTGRNIKFS	LSQKF				745

SEQ ID NO: 40            moltype = AA    length = 742  
 FEATURE                Location/Qualifiers  
 source                1..742  
 mol\_type = protein  
 organism = *Pasteurella multocida*

SEQUENCE: 40

MDKNLMKGCV	FLLIVCGIQ	IGLASNPNNPP	DVDELLPII	NADEDNKLPG	RSVLQKQNI	60
QQQADNAADL	INILPGVN	GGFRPGGQTL	NINGMGAED	RVQLDGATK	SFEKYQQGSI	120
FIEPELLRKV	TVDKGNYSPO	YGNNGFAGTV	KFETKDADTF	LKENQKIGGL	FKYGNNSNNN	180
QKTYSTALVL	QNEQKNIDL	LFGSVRNASN	YTRPDKSIL	FSKNNQKSGL	IKVNWQITPE	240
HLLTLSSVYG	IHKGWEWPAA	KRDVMSRPT	TEIKRYGIVD	AWKRKLVYRD	QKDESYSLKY	300
RYLPENNWKI	NLSVQLSYSK	TEQNDTRHEK	VTSSFLGTL	NKSWITYSDL	TFDISNTS	360
NIGRAEHELL	FGLQWLKNKR	NTLHYHK	KKADYNNGYF	QPYYMPSGRQ	YTQAFYLQDQ	420
IWKQNFLFTG	GIRYDHINNI	GQKNLAPRYN	DISAGHDYSQ	KNYNGWSY	GLKYDVNHYL	480
SLFTNFSKTw	RAPVIDEQYE	TQYSQASVSA	TSLNLEKEMI	NQTRVGGIIT	LNHLFQENDA	540
FQRTTYYFYN	RGKNEIFKTR	GVNCEVALD	VDNSVCPKII	SNYRNLPGYV	IQGAELEAYY	600
QSSYLFGGLT	YSYVKGKRTD	SPRNPKWTS	TWIAETPPRK	ATATLGFNIP	EYYFTAGWRA	660
EFVRKQDRSP	LSNDSKASYW	ALPSSKGYS	HSVFFSWSP	KIKGMNFKV	VDNLNRPY	720
PYLGELASGT	GRNVKFSLTQ	QF				742

SEQ ID NO: 41            moltype = AA    length = 742  
 FEATURE                Location/Qualifiers  
 source                1..742  
 mol\_type = protein  
 organism = *Pasteurella multocida*

SEQUENCE: 41

MDKNLMKGCV	PLSIVCGIQ	IGLASNPNNPP	DVDELLPII	NADEDNKLPG	RSVLQKQNI	60
QQQADNAADL	INILPGVNMA	GGFRPGGQTL	NINGMGAED	RVQLDGATK	SFEKYQQGSI	120
FIEPELLRKV	TVDKGNYSPO	YGNNGFAGTV	KFETKDADTF	LKENQKIGGL	FKYGNNSNNN	180
QKTYSTALVL	QNEQKNIDL	LFGSVRNASN	YTRPDKSIL	FSKNNQKSGL	IKVNWQITPE	240
HLLTLSSVYG	IHKGWEWPAA	KRDVMSRPT	TEIKHYGIVD	AWKRKLVYRD	QKDESYSLKY	300
RYLPENNWKI	NLSVQLSYSK	TEQNDTRHEK	VTSSFLGTL	NKSWITYSDL	TFDISNTS	360
NIGRAEHELL	FGLQWLKNKR	NTLHYHK	KKADYNNGYF	QPYYMPSGRQ	YTQAFYLQDQ	420
IWKQNFLFTG	GIRYDHINNI	GQKNLAPRYN	DISAGHDYSQ	KNYNGWSY	GLKYDVNHYL	480
SLFTNFSKTw	RAPVIDEQYE	TQYSQASVSA	TSLNLEKEMI	NQTRVGGIIT	LNHLFQENDA	540
FQRTTYYFYN	RGKNEIFKTR	GVNCEVALD	VDNSVCPKII	SNYRNLPGYV	IQGAELEAYY	600
QSSYLFGGLT	YSYVKGKRTD	SPRNPKWTS	TWIAETPPRK	ATATLGFNIP	EYYFTAGWRA	660
EFVRKQDRSP	LSNDSKASYW	ALPSSKGYS	HSVFFSWSP	KIKGMNFKV	VDNLNRPY	720
PYLGELASGT	GRNVKFSLTQ	QF				742

SEQ ID NO: 42            moltype = AA    length = 742  
 FEATURE                Location/Qualifiers  
 source                1..742  
 mol\_type = protein  
 organism = *Pasteurella multocida*

SEQUENCE: 42

MDKNLMKGCV	PLSIVCGIQ	IGLASNPNNPP	DVDELLPII	NADEDNKLPG	RSVLQKQNI	60
QQQADNAADL	INILPGVNMA	GGFRPGGQTL	NINGMGAED	RVQLDGATK	SFEKYQQGSI	120
FIEPELLRKV	TVDKGNYSPO	YGNNGFAGTV	KFETKDADTF	LKENQKIGGL	FKYGNNSNNN	180
QKTYSTALVL	QNEQKNIDL	LFGSVRNASN	YTRPDKSIL	FSKNNQKSGL	IKVNWQITPE	240
HLLTLSSVYG	IHKGWEWPAA	KRDVMSRPT	TEIKHYGIVD	AWKRKLVYRD	QKDESYSLKY	300
RYLPENNWKI	NLSVQLSYSK	TEQNDTRHEK	VTSSFLGTL	NKSWITYSDL	TFDISNTS	360
NIGRAEHELL	FGLQWLKNKR	NTLHYHK	KKADYNNGYF	QPYYMPSGRQ	YTQAFYLQDQ	420
IWKQNFLFTG	GIRYDHINNI	GQKNLAPRYN	DISAGHDYSQ	KNYNGWSY	GLKYDVNHYL	480
SLFTNFSKTw	RAPVIDEQYE	TQYSRASVSA	TSLNLEKEMI	NQTRVGGIIT	LNHLFQENDA	540
FQRTTYYFYN	RGKNEIFKTR	GVNCEVALD	VDNSVCPKII	SNYRNLPGYV	IQGAELEAYY	600
QSSYLFGGLT	YSYVKGKRTD	SPRNPKWTS	TWIAETPPRK	ATATLGFNIP	EYYFTAGWRA	660
EFVRKQDRSP	LSNDSKASYW	ALPSSKGYS	HSVFFSWSP	KIKGMNFKV	VDNLNRPY	720
PYLGELASGT	GRNVKFSLTQ	QF				742

SEQ ID NO: 43            moltype = DNA    length = 2418

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SEQ ID NO: 46          moltype = AA  length = 790
FEATURE
source
1..790
mol_type = protein
organism = Bibersteinia trehalosi

SEQUENCE: 46
MFNKKLALLV ISAQFSPLVW ANNNDAVLD EVSVVGSTPS ISQGSEVTLV KVSDKIIAGK 60
EFKKRSATLG NALAAELGVH SNPFGGASK PIIRGQEGAR IRILQNGSDV IDMSNLSPDH 120
AVVADSLLAQ QVEILRGSSST LLYASSSPAG IVNVVDKRIP TEIPEKGYEV ELNSRFDTAA 180
KEKVGAAGAT FGIGKHIAVR AEGLTRHSDN YRVPGINLGE RLNYVPDTYN KSKVGTGLS 240
FVGEQGYIGA SYSKRDRNYG LPGHNNHKDFD CTGHIYGNKR DKYAYTLYP HLIGEENIGS 300
NPHFHCGTNH AEDGTHSHDN PFGHDHDHTH PGPWVDLESK RFDVKAELRQ PFKGIDKIKV 360
SYADADYYHD EKDAGVLATR YHKQLKKDQD YGKPVNIFKN RGKNARLEIY HAPLDGLTGV 420
WGQVYQTQKS SMHAPKDRREV KFPLVENTNK QMSLFGIEQY MWDNFALEFA GRVEKQKIEI 480
EYDRNEIKRL QDHYRISGGQ QVEPDLSYPN QNAYAYSGLT NWFFHPDYQL SFTASHNERF 540
PTPMELYHYG QHIAATNSFEY GNKDLKKEQS NNVELGLGYQ TERVGYKVNV YYNHFKNYIY 600
NEENLFRENQL FMRRYNQAKA RFYGIEEAS YRFNPKYQAT IFGDMVRGLW TNLPPLTVNS 660
DYSVFKDYL KDAKPGEDYL IYRADQNTPR TPPVRLGFRF NAEFTPNSWG DLELIRTFTQ 720
RRTSQLEYIT EGNTMLNIGV AYSNKWKDLD YKISLNGTNL LNQPVYIHTS YHQFVPMGR 780
NFILGMEMKF 790

SEQ ID NO: 47          moltype = AA  length = 782
FEATURE
source
1..782
mol_type = protein
organism = Actinobacillus pleuropneumoniae

SEQUENCE: 47
MLISAQFSPV VWWANNNDVAV LDEVS VVGST PSISQGSEV LLKVS DKIIIA GKEFKRSAT 60
LGNALAAELG VHSNPFGGG SKPIIRGQEG ARIRILQNGS DVIDMSNLSP DHAVVADSSL 120
AKQVEILRGS TVEILRGSST AGIVNNVVDKR IPTEIPEKGY EvelNSRFDT AAKEVKGALG 180
ATPGIGKHI A VRAEGLTRHS DNYRVPGINL GERLN YVDPD YNKS VGTGLG LSFVGEQGYI 240
GASYSKRRDN YGLPGHNHKF DFCIC HIYGN KQGKYAYT YPHLIGEENI GS NPHFHCGT 300
DHAEDGTHSH DNPF GHDDH THPGP WDLESK SKRFDVKAEL RQPF KGIDKI KV SYADADYY 360
HDEKDAGVLA TRYHKQLKKD QDYGKPVNIF KNRGK NARLEI YHAPLGGLT GVWGVQYQTQ 420
KSSMHPKDR EVKFP LVENT NKQMSLFGIE QYMW DNF ALE FAGR VKEQK EI EYDRNEIK 480
RQDHYRISG GKQV EPDLS YNQNAYAYSS TLNW FFHPDY QLSFTASHNE RFPT PMELY 540
HGQHIA TSF EYGNKDLKKE QSNNV ELL GL YQTERVGYKV NVYY NHFKNY IYNENL FREN 600
QLFMRRYNQAKA KARFY GIEAE ASYRFNPKYQ ATIFGDMVRG WLTNLPPLTV NSDYSVFKDY 660
LPKDAKPGED YLIYRADQNT PRTPVRLGF RFNAEFTP VN SGDLE LIRTF TQRRTS QLEY 720
ITEGNTMLNI GVAYSNKWD LDYKISLNGT NLLNQPVYIHTS TSYHQFV PQT GRNFILVVNV 780
KF 782

SEQ ID NO: 48          moltype = AA  length = 790
FEATURE
source
1..790
mol_type = protein
organism = Mannheimia granulomatis

SEQUENCE: 48
MTNKTILALL ISAHLSSLAL AQTKE DIAVL EEVSVI GN SD TPVVAQGSEV TILK VS DKII 60
DGKEF KRSATL TG NALAAEL GVHSN PFGGG ASKPIIRGQEG GARIKI LQNG AEVVD MSSL 120
PDH AVVADSL LASKV E VLRG SSTLL YTSSS PAGV VNVVDK RIPT SIPEKG YEVEL NSRFD 180
TVSKERL GVI GATVGLGQHV ALRVEGLN RH SD NYR VKS LQ LGESL NYV PD TYN RS RVG TM 240
GISFIG ERGY IGAAY SHRK D TYGLP GHN HK FDFCTG HIY VDRDK H AHTY LY PHL LT NEL 300
ISEN P FH C SDH GLD HN HS HDN PYG HKD HT HKGP VNDL RSKR L D LK M LK QP FSG IEK 360
LRLS HTDV NY YH DEKD AGV L LTFLK QL DKD YG KPV NI YK NR GKN SRL EF YH T P V GGL 420
TGVW GLO YQT QKSSM NAPK D RE VRP PLI EN N NK QL S L FGI EQ YMW DNF A V EL AGR IEK QK 480
IEIDYD LNEI KRL QDF YRIS GGAQ VEP DLS PYNK TAY AYS GTLN WFF HPN Y QLSFTASHN 540
ERHPT PMELY YHG QH LAT SS FEYGNK DLK K EQ SNN VEL GL GY QG EK FGY K TT VY YN HFKN 600
YIFNEN LYRE D QL FMR RY SQ AKAR FY GLE A EMSYRF NDK Y QATL FGD MVR GWL DLP PVK 660
VGGI HKHE LKP DAKP GED YL RADM NTPR TPPV RL GRL NAQF NEN WAG EA E F YRM FSQ 720
RRLS QLEY PT NGH SML NL GL SYSN KFK NAE YKISL NGT NL NQAVYIHTS YHQF VSP QGR 780
NFIL GLEM KF 790

SEQ ID NO: 49          moltype = AA  length = 790
FEATURE
source
1..790
mol_type = protein
organism = Mannheimia sp.

SEQUENCE: 49
MTNKTLLAVL ISAHLSTLAL AQT NED VAVL DEV SVI GNTA TPVIAQGSEV TLLK VS DKII 60
EGKEF KRSATL TG NALSAEL GVHSN PFGGG ASKPIIRGQEG GARIKI LQNG SD VV D MSSL 120
PDH AVVADSL LASQ V E VLRG SSTLL YTSSS PAGV VNVVDK RIPTI PEKG YEVEL NSRFD 180
TASKERL GVI GATVGLGQHV ALRVEGLN RH SD NYR VKS LQ LGER L NYV PD TYN RS RVG TV 240
GLSF VGERGY IGAAY NERTD TYGLP GHN HK FDEC IGH YIN EV RD KYAY T YK PH L DED L 300
ISHG P H FHC G TDHE MDAGHS HDN PYG HTD HT HKGP VNDL KSKR IDV KGE LLK PFR GLD K 360

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IRASYTDVNY	YHDEKDAGSV	STFVNKQLKK	DQNYGKPVNII	YKNRGKNSRL	EFYHSPIGGL	420
TGVWGVQYQT	QKSSMNAPRN	REVRPPLVEN	NNKQLSLFAI	EQYMWDFAL	ELAGRIEKQK	480
IEIDYDLEEI	RRLQKPFYSES	GGRQVEPDLS	PYDKTAYAYS	GTFNWFHFD	YQLSFTVSHN	540
ERHPTPMELY	YHGPFLATSS	FEYGNKDLKK	EQSNNVELGL	SYQGEKFQYK	TTVYYNNHFKN	600
YIFNENLYRE	NQLFMRRYSQ	AKARFYGLEA	EMSYRFNEQY	QATIFGDMVR	GWLTDLPPVK	660
VCGIHKKHLP	KDAKPGETYL	LYRADMNTPR	TPPLRLGLRV	NAQFNENWAG	EAEFYRMFSQ	720
RRLSQLEYPT	KGHSMNLNLGL	SYSNKFKNAE	YKISLNGTNL	LNQPIYIHTS	YHPFVSQPGR	780
NFLGVEVKF						790

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SEQ ID NO: 50	moltype = AA	length = 817				
FEATURE	Location/Qualifiers					
source	1..817					
	mol_type = protein					
	organism = Haemophilus parasuis					
SEQUENCE: 50						
MINNRTTEQQ	NRRTTEQQNN	RTTAFSLAFS	LLLCCLGINA	EQELELDEISV	MGKVPEGNSI	60
SFLKVSDAII	DGEKFKNRSA	TLGNALSSSEL	GVHSTPFGGG	ASAPIIRGQE	GVRVKILQNN	120
ADVVDMSNIS	PDHAITADTL	LANQVEILRG	ASTLLYASSS	PAGIVNIVDQ	RIPNKMPPKG	180
YEVTLSSRFD	TASKEVYAL	GTTIGIGKHL	ALRLEGGLDRQ	SQNYKVPQIK	LGETLNYVPD	240
TYHQSKVGTI	GLSFIGEKGY	LGASYNQRKD	RYGLPGHNNHK	FDTCTIAHYD	MRLQGKHSYT	300
NLYPHLMSDE	MVTENPHFHC	GTDYDLDPSH	SHDHPYGHDH	DHTHIGPWD	LHSKRIDIKG	360
EIKQPLPMLD	KIQLSYAQTD	YYHEDKACK	SGDTINPNR	DKSKDGFQPV	NIFKNQGKNA	420
RLEFFHTPIG	GLTGGMFGVQY	QTLQSSANTP	SNREVQWPPLV	DNRNQKQISLF	ALEYQAWDNF	480
AZIELGLRTEK	QNIHIDYDLA	KIQKQOKFNE	RTYGKQVDPD	LSDYDKEAIS	YTGAPNWWFH	540
PDYQLSFTAS	HNERLPTPME	LYYHGQHLAT	NSFEYGNKDL	KKEISNNFEL	GLGYHTEKLD	600
YKLSTYYNNF	DNYIYNETY	RSNNLFLMRRY	NQAKATFYGL	EGIIINYRFTP	DYQFSVFGDM	660
VKGKLKQLPD	IKGFLNDVYGE	PILPNDYDP	YDEPEDQYR	PYLKGEMIKQ	ADRVSPLRPP	720
IRLGARFNAQ	LTCNLGSVE	MVKVTFQNKV	SKLESSTKGY	QLLNASLNYR	RQIKGVEYTV	780
SLTGNNNLLNQ	AVYIHNNSYHP	YVPQMGRNF	LGLDLSF			817

SEQ ID NO: 51	moltype = AA	length = 825				
FEATURE	Location/Qualifiers					
source	1..825					
	mol_type = protein					
	organism = Moraxella boevrei					
SEQUENCE: 51						
MAVLASSSMS	VFANDSSPET	TLDTIVIEAT	SANKVNSLA	AESQKASDMV	ISKERLKSKS	60
ATLGNALAGE	SGIHSNPF	GASAPVIRGQ	EGVRVKILQ	GSAVADMSTV	SPDHVVAVDT	120
LLASRVEVNR	GASTLMHANA	SPAGVINVD	GRIPDKMPTG	ITGETMLRFN	SGSDEKLATA	180
GISAPLNDNF	VLRLEAMGRD	ANPYNVPAIN	FGEVNLNYLPD	SYNSTVTGTL	GLSYIGNQGH	240
IGLAYSERRD	KYGLVGHNHK	FDNCEGHAFN	TSRGLWGPER	RYLIPYPHLM	SDEDIMITSLH	300
FHCGTNYLDL	PSHSHEHVG	HKHDHTQKGP	WVDMTSKTS	LQGEINQPIP	SIDKIRLTAS	360
HSNYEHQEYD	DGKQIPDPSK	GKIVFKGNTS	YWDNQGLSSK	LSIYQSPTRD	LSFWGFDNQ	420
RNKTHALIPLS	PNEKAGNRR	LVKNTQKTHS	IFGLSEYKIG	NVKLHTALRQ	ERTRIPVEYN	480
MDEIKAQLAN	GIGTQBLPDL	TPYKSNATNS	AMGAIWIDNP	KLRLDTTFSH	NERTPTPMEL	540
YYHGKHLATN	SFLYGNKDLN	KEKSDNGEGL	LTFKGDKWRV	KGSIYANKFD	NYIHPENLYK	600
SGNLTMMRFT	QSKAKLRGAE	LEIGYQFNPN	LNVSLSFGDMV	RGKLYGFSP	TGGNLYEVVK	660
VIDPNCDLEK	DDPDYEDFCI	DIENKVVGD	TITRPDRTP	RLSPDRIGMR	INGEYGNFSP	720
SLEFIRVEDQ	NRTSNVAKA	YNSECTHHQY	GNERLCPIFI	QEDATKGYNL	LNVLQDHYNY	780
FKGLQYTVSL	NANNLLNEKV	YIHNSFLPFV	PQQGRNFSLA	LTTKF		825

SEQ ID NO: 52	moltype = AA	length = 869
FEATURE	Location/Qualifiers	
source	1..869	
	mol_type = protein	
	organism = Psychrobacter phenylpyruvicus	
SEQUENCE: 52		

MIKIKPLSYAV	SAIMLSQLSH	TPMAQAAVSS	KTVTDQDQK	QTPSQQASET	DSTPDSASNP	60
IAIFDTITVP	PSSTHVPSSL	AFDNAQKASD	VVIDNEKLRH	RSATLGNALA	DELGVHSNPF	120
GGGSSAPVVR	QGEGVVRKIL	QNGMDAIDMS	TLSPDHVVG	DTLLADKVEL	IRGASTLLYS	180
NASPAGVINV	VDGDRIPTEV	EGYTGEATLR	FNEENNDERVA	TAGITFGLTD	NVALRVEGLT	240
RKANEYEVPE	INLGDKLNYL	PDSQNKSNVG	TIGLSYVGER	GHIGVAYSER	EDKYGLVGHN	300
HKLDCGCYGHV	VYPQKNYKNC	PYLAAYPHLM	GDEDLAESFH	FHCDSDHMED	EPHSHDNPYG	360
HDHDHTQZGP	WVDMNSKSYY	LQGELLEPIP	AIEKVRLNVA	YHDYHHEEH	HGKTIPDPAK	420
GERFVKAQPS	WVDMNSKSYY	LEAYHTPTEH	LQGVWGIQSQ	SHKSSALIPS	KEEHQPNRRP	480
LVENKHRQFS	VFGVEQYKLN	DWLFEAGLRY	EHSKIPVTYN	LDEIEAQNKI	LGQLLKPEQ	540
DLTPYQESAF	SYALGAIWDM	TPEYRLLDMSY	SHNERLPTPM	ELYHGKLN	TNSFLYGNKD	600
LDKEESDNFE	LGIQFTGDKW	RYKASAYSNH	FDNYVHAENL	HKDGNLYMRR	MTQSKAKIQC	660
LEAEVGYEYL	PGHSATLFGD	YVRGKLYGFA	PVYGNEIKSQ	NEVIKYMPPE	ECGASPGDDV	720
YEEWCYPDY	EVIGIDKVER	PERNAPRMSP	LRLGLRLRNNE	YNDNWSTSLLD	FTRVFAQNK	780
STATVVNIPR	DESRLKVTVE	PEDSTSGYSL	LNVGVDYKNT	WDNAKHPV	TLSLRANNLL	840
DEIYIVHNSF	LPYVPQMG	FMLSLNVEF				869

SEQ ID NO: 53	moltype = AA	length = 805
FEATURE	Location/Qualifiers	
source	1..805	

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mol_type = protein
organism = Pasteurella multocida

SEQUENCE: 53
MISRGCKVNK FFAVLMCCCI PQVWANTEK KQIVFLDEIS VESKGAAFIRS DPLSLPKQN 60
DILVSKQKLK TGSSTLGNAL AGELSVHSNQ FGGSSAPVV RGQEGVRLKI LQNGSDVIDM 120
SQLSPDHAIG VDTLAEQVE IVRGASTLLY ANASPAVGIVN VVDKRIPTQL PQKGYEVDFN 180
TRYNTNSHEK LVTAALTGFL GKHTALRVEE LLRGSNYYHV PAFKLDKTLN YVPDTQNKTG 240
SGNYGVAFIG ERGYVGFAYN LRREKYGLPG HNHKLDSCAA HIWGGNVRND YYLELYPHLM 300
HDTDLVNTHF HCGSNHMDMG KHSHDHPYGH DHDHSIAGPL IDSYAKRYDI RAEVKQPMKA 360
IEKIKLSYE TRYKHDEKDG NIAVNLFKNN GYNLRVEIFH TPIAGLSGVI GAQYQTQTS 420
ANIPRIALCS NNASDPCHKK KQRDKPSKITK GDRKSVALINE NTQSQMSFFA IEQLRWQDFL 480
FEIGVRTEQ RIDETVHDRAW LFKVKRLEG CDPNSFFYSP SGCRQGSYPA PDFASYHDRA 540
TYSYGAISWN MTPDYTLSLT YSHNERHPTP MELYHGKHL ATVSFEHGNR NLKKEVSDNW 600
EVGLAYLGDK LSYKVNVYYN DFKNRIFNQT LNKSGNLSLN RYNQSKAKYY GVEGRIDYAL 660
TPELHMGLFG DYVRGKLYDL PPTYRVDHVA NSLEPVPPQD QDAPRVPPMR LGFRVNMEMT 720
ESLTSSLEYT VVYQQKKVAP LENQTAAYSL LNIRVDYSRQ IAGVNYQLFV QANNVLNRKV 780
YSHTSFLPFV PQMGRNVTLG LNIHF 805

SEQ ID NO: 54      moltype = AA length = 826
FEATURE          Location/Qualifiers
source           1..826
mol_type = protein
organism = Moraxella bovoculi

SEQUENCE: 54
MVNLYSRPTV LALSIALVSL SAHADDTATK TEVTLDLNA VVKGGMQTT S TLKKPSDMII 60
RDRDTLQRSA TLGNALAGEL GIHSNPFGGG ASKPIIRGQD GVRVKILQNG TDVIDMSALS 120
PDHVVAADTL LASRVELVRG AGTLLYGTAS QAGVVNVIDE RIPSRMPQGN IKEKIEGETL 180
LRYNTGSNEK VVTASLNMGV GQNVAVRVEG LTRRADDYEV PGFQSDVMLD YLPDSHNKST 240
VGTGVGSYIG DKHGIGVSYS RRQDKYGI PG HNHAYDNCIA HVLTPEASIS RYYLKAYPHL 300
IQNMDFSSSA HFHCCTDHAA DPGQSHEHPL GYEHDHTHPG PWIDMESERI DVRAQWEKPF 360
KGGLDKIALSF TSSDYYHEEN DHEVKLDKY TGRLQTTINNK PGYFGNQGKN VRVEFHHTPV 420
KDVSGLWGMQ WQKQESFAHL PHEREQGRY PLIANTNKQV SVFGLERWQA NDKWAEFEGT 480
RFEKQSIPIID YSEDKLDYR PKPECWDWGP SSGCLPAPNY EEPDLTTYKE KATSYSIGTT 540
WDFKPDYRLS ATYSHNERLP TPMELYHGK HLATHSFEFG NIGLDKEKSN NLDLGISFSG 600
DKWSYAVNAY HSRFKNYIYN ENVYREGNLF MRRHNQAKAD FYGLEGMVTVY NMDNHAISLF 660
GDLVRGKLKD LPNAYAKAYY CDGAYTTTKP EDTEENMQRQ NYNYFTDDFN YSYEPIIAQP 720
NMPTPRLPPA RLGLRWQGDL SANWSADAEBY MHVFGQNRIS KLESATASYD MLNLGLGYHN 780
HWGNVDYTLS LRANNVLDEK VYIHNSFLPF VPQMGRNFSL SANFKF 826

SEQ ID NO: 55      moltype = AA length = 850
FEATURE          Location/Qualifiers
source           1..850
mol_type = protein
organism = Conchiformibius steedae

SEQUENCE: 55
MFSRMLLPL LLGLGANAAY AEQNTSPA朴 ATEDNEGKTL PAMQGVGKRR DTAQPFSGNR 60
KASDMVISQE KLKSRSANLG EALAGELGIH GNPGGGASA PVIQGQEGVR VKILQGGSDV 120
VDMMSALSPDH AVAADTLLAQ QVEVLRGTST LAYAAASPAG VINITDKRIP DRLPAKGWEA 180
ETGKVRFDTAA KEKALTAGAT FGIGKHFAVR AEGLERKSDS YRVPGINLGE TLKYVPTDYN 240
RSHTGTLGVs WVGQNGHGLV SYSHRKDRY LGPGHNHMLDN CSGHVFVDVT ASAVERNYLL 300
PYPHLIGDED VNLSQFHCH TEHSSNAKHS HDNVYGHKHD HGEPPFWIDM RVRRYDVRGE 360
WRTQLPFLK IRLLTSAYTDY YHDEKNDGKV YISPDDPEGW RERKLKDAAA RKGKPDIIILK 420
NKGLNTRLEF PHRGFSVWNG MAGVQYQTQR SSARRVMPPM IGGERYINER NPLVDNTNKQ 480
LSLFALQQYR RGNWLAEGGV RWEKQRIPIR YDHDLAQYV KPGTQQPDLK PYSQKALSYS 540
GSSLWDFKPG YRLSLTASHN ERLPTPMELY YHKGKHLATNS FEYGNKDLKK ERSNNYEIGL 600
RYFGDKWDYK LSVYHNRFKN YIYNENLYRS GNLFIIRRNTQ AQGRFHGVEG EVSYRFKPTH 660
QVTLFGDMVR CRLFGLSPVY GDKIYREYEC VDEDGLEDTC FEVVGREKIE RPDRHAPRVP 720
PTRLGRLNLS QWGDNWTASL EYARWAQNR TAVSQFPRER DDEDEDEDE GNPKPRQQL 780
YAEPVLEDPT SGYHLLNAGI AYRKIGKAD YRSLDAFNL LNKKVYIHSN HLPYVPRPGR 840
NFVFGVNVSF 850

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- 1-27.** (canceled)
- 28.** A composition comprising:  
an isolated protein having at least 80% similarity to amino acids 26-805 of SEQ ID NO: 44; and  
an adjuvant.
- 29.** The composition of claim 1, further comprising:  
an isolated protein having at least 80% similarity to amino acids 25-968 of SEQ ID NO:2,  
an isolated protein having at least 80% similarity to amino acids 27-790 of SEQ ID NO:4,  
an isolated protein having at least 80% similarity to amino acids 23-727 of SEQ ID NO:6,  
an isolated protein having at least 80% similarity to amino acids 25-964 of SEQ ID NO:8,  
an isolated protein having at least 80% similarity to amino acids 26-848 of SEQ ID NO: 10,  
an isolated protein having at least 80% similarity to amino acids 27-784 of SEQ ID NO: 12,  
an isolated protein having at least 80% similarity to amino acids 25-742 of SEQ ID NO: 14, or a combination thereof.
- 30.** The composition of claim 28 further comprising: at least one isolated protein having a molecular weight of 249 kDa, 60 kDa, 42 kDa, 38 kDa, 27 kDa, 26 kDa, or 22 kDa, wherein the proteins are isolated from a *P. multocida*.
- 31.** The composition of claim 28 further comprising a pharmaceutically acceptable carrier.
- 32.** A composition comprising a bacterium engineered to express the protein of claim 28.
- 33.** The composition of claim 32, wherein the bacterium is gram negative.
- 34.** The composition of claim 33, wherein the gram negative bacterium is *E. coli*, *Salmonella* spp., or *Pasteurella* spp.
- 35.** The composition of claim 34, wherein the *Pasteurella* spp. is *P. multocida*.
- 36.** The composition of claim 32, wherein the bacterium is inactivated.
- 37.** The composition of claim 32, further comprising an adjuvant.
- 38.** A composition comprising a whole cell expressing the protein of claim 28 and an adjuvant.
- 39.** The composition of claim 38, wherein the bacterium is *P. multocida*.
- 40.** The composition of claim 38, wherein the bacterium is inactivated.
- 41.** A method comprising: administering to a subject an amount of the composition of claim 28 effective to induce the subject to produce antibody that specifically binds to the protein.
- 42.** A method for treating an infection in a subject, the method comprising: administering an effective amount of the composition of claim 28 to a subject.
- 43.** The method of claim 42, wherein the subject has an infection caused by a *P. multocida*.
- 44.** The method of claim 42, wherein the subject is a mammal.
- 45.** The method of claim 44, wherein the mammal is a bovine.
- 46.** The method of claim 42, wherein the subject is an avian.
- 47.** The method of claim 42, wherein the avian is a chicken or a turkey.
- 48.** The method of claim 42, wherein at least 0.01 micrograms and no greater than 3000 milligrams of protein is administered.
- 49.** A kit for detecting antibody that specifically binds a protein, comprising in separate containers:  
the isolated protein of the composition of claim 28; and  
a reagent that detects an antibody that specifically binds the protein.

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