

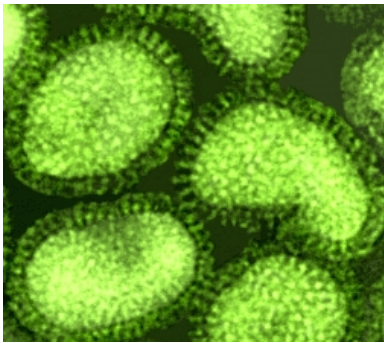
Virus and Bacterial Membrane Proteins

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Two viruses and seven bacteria have been chosen as examples to illustrate the structures of membrane proteins.

Influenza

All the pathogenic subtypes and mutations of the influenza virus so far identified have an abundance of particular prolyl peptides (**XPY**, where **P** is proline and one or both of the adjacent amino acids, **X** and **Y**, are hydrophilic amino acids – love water) in their hemagglutinin and neuraminidase surface membrane proteins. (see Figure 1: the hemagglutinin and neuraminidase proteins are the hair-like structures on the surfaces of the viruses) In addition, neuraminidase proteins - essential for host cell invasion and proliferation and in the former, for causing an immune response that damages the host's organs - in each variant of the influenza virus for which the structures are known.



For example, **the sequence of N1 neuraminidase** in the Japan/China H5N1 avian influenza (Mase, M., Eto, M., Tanimura, N., Imai, K., Tsukamoto, K., Horimoto, T., Kawaoka, Y., Yamaguchi, S. "Isolation of a genotypically unique H5N1 influenza virus from duck meat imported into Japan from China" **Virology**

Figure 1: influenza virus 339 (1), 101-109 (2005)) has multiple potential sites for ginger enzyme hydrolysis (**bolded**): these sites are adjacent to **P-3, 48, 93, 120, 154, 167, 169, 198, 246, 272, 283, 302, 326, 328, 340, 377, 410, 420, 431, and 458**, and the blue highlighted prolines are conserved over

the three H5N1, the H1N1 swine flu of 1918 and 2009, and H9N7 bird flu, with the pink highlighted prolines are not conserved in the H9N7 bird flu:

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1  MNPNQKITTI GSICMVGIV SLMLQIGNII SIWVSHSIQT GNQHQAEPCN QSIITYENNT
61  WVNQTYVNIS NTNFLTEKAV NLVTLAGNSS LCPISGWAVY SKDNGIRIGS KGDVFVIREP
121 FISCSHLECR TFFLTQGALL NDKHSNGTVK DRSPHRTLMS CPVGEAPSPYNSRFESVAWS
181 ASACHDGTSW LTIGISPDN GAVAVLKYDG IITDTIKSWR NNILRTQESE CACVNGSCFT
241 VMTDGPSNGQ ASYKIFRIEK GKVVKSAELNAPNYHYEECSCYPDAGEITCVCVRDNWHGSN
301 RPWVSFNQNL EYRIGYICSG VFGDNPRPNDGTGSCGPVSPKGAYGIKGFSFRYGNGVWIG
361 RTKSTNSRSG FEMIWDPNGW TGTDSNFSVK QDIVAITDWSGYSGSFVQHPELTGLDCIRP
421 CFWVELIRGR PKESTIWTSG SSISFCGVNS DTVGWSWPDG AELPFTIDK
```

Each letter in the sequence represents an amino acid. The important hydrophilic amino acids adjacent to prolines are: asparagine, **N**; glutamate, **E**; cysteine, **C**; serine, **S**; histidine, **H**; aspartate, **D**; arginine, **R**; lysine, **K**; tyrosine, **Y**.

Cleavage of the neuraminidase proteins by the ginger enzyme will prevent the virus invading the host cells and prevent proliferation of the virus in the host.

The identified epitopes for hemagglutinin are not linear peptides but are what is called "conformational" epitopes in which the participating amino acids are brought together via the three-dimensional structure of the protein. Seven of the H1N1 2009 (Swine flu) active prolyl peptides are included in the conformational epitopes and six of these are conserved prolines in H1 and H5. Numerous other non-proline amino acids in the epitopes are not conserved. (Deem, M.W., Pan, K. "The epitope regions of H1-subtype influenza A, with application to vaccine efficacy" **Protein Eng., Design & Selection**, 1-4 (2009, July 3))

EPITOPES (CONFORMATIONAL)

```
A:  RQLSSFERFPKSWPNHDKGTWGD
B:  VSCPHAGAFKDKGKE LVL GIHH
C:  DTVLENVVTH AFAMER  AGSSHTQPKNTLPFQNI
D:  AYIVDLLVKKGNSYPLSSSDQSLYQNADTYVFV SKFKPVDERNYY
E:  VNLEKHNLLGKCNIAG LGNPETFEATGLR
```

Underlined letters are three or more amino acid linear peptides.

Yellow highlighted letters are part of an active proline peptide group which is cleaved by the ginger enzyme.

The inhibitors currently on the market function differently to the ginger enzyme: they are specifically designed to 'plug' the active site of neuraminidase where the neuraminidase opens the virus' surface membrane so the virus can enter the host's cells. They rely on the amino acids that surround the active site (glutamate, arginine and aspartate) to stabilize the binding of the plug in the hole. (see Figures 2). The above N1 variant in H5N1 bird flu does not have arginine, R, in positions 92 and 371 and subsequently the binding and efficacy of the 'plug' could be significantly reduced.

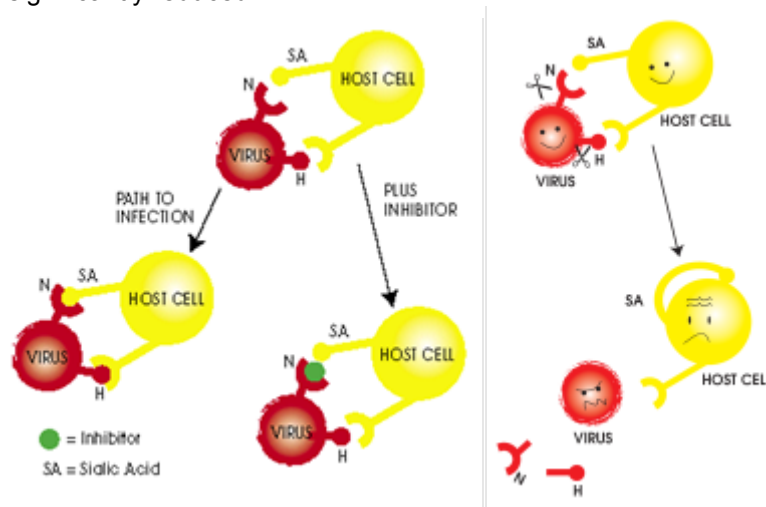


Figure 2: The influenza virus invades a host cell by initially binding to receptors on the host cell, one of which binds to the hemagglutinin (H) on the virus, and one with sialic acid (SA) which binds to the neuraminidase (N) protein on the virus. Current inhibitors such as Relenza are designed to mimic the sialic acid and to bind to the neuraminidase blocking the link to the host cell. The Biohawk ginger product acts (on the right) like a pair of scissors and specifically cuts off the hemagglutinin and neuraminidase proteins from the surface of the virus completely preventing the virus infecting the cell and replicating itself.

Importantly, although the various H5 structures show significant mutations, the potential sites for ginger enzyme hydrolysis are largely conserved and number at least 15. In the H5-hemagglutinins recently found in the Vietnam (Nguyen T.D., Hanh T.H., Puthavathana P., Long H.T., Buranathai C., Lim W., Webster R.G., Hoffmann E. "Lethality to Ferrets of H5N1 Influenza Viruses Isolated from Humans and Poultry in 2004" *J. Virol.* **79**, 2191–2198 (2005)) and Japan/China variants, and previously identified in Singapore H5N1 avian influenza (Ha, Y., Stevens, D.J., Skehel, J.J., Wiley, D.C. "Structure of Avian H5 Haemagglutinin Complexed with LSTA" *Proc. Nat. Acad. Sci. USA*, **98**, 11181 (2001)), the prolines are conserved. In addition there is an extra proline, P-233, in the Singapore H5N1, which has the hydrophilic arginine and lysine adjacent to it, but in the Vietnam and Japan/China variants, proline is replaced by serine. The hydrophilic amino acids adjacent to the prolines are also conserved, except for those adjacent to P-101 with asparagine (N-100) for the Vietnam and Singapore variants and serine (S-100) for the Japan/China protein, P-108 with the non-hydrophilic glycine for the Vietnam and Japan/China variants and the hydrophilic glutamate adjacent to the proline for the Singapore variant - giving an additional site for hydrolysis, P-134 with lysine for the Vietnam and Japan/China proteins and with arginine for the Singapore protein, and for P-337, which has glutamine following it for all three but with serine before the proline for the Vietnam and Japan/China variants and valine for the Singapore protein. The target prolines for ginger enzyme hydrolysis are: P-65, 81, 90, 101, 108 (Singapore only), 134, 174, 197, 210, 227, 233 (Singapore only) 251, 266, 297, 312, 319, 337, 506.

This gives the ginger enzyme excellent opportunity to hydrolyze the H5 at multiple sites. The conservation of the prolyl residues in the hemagglutinin structures is suggestive of these having a specific role in the function of the protein. Hydrolysis by Biohawk's ginger at these specific sites in the neuraminidase and hemagglutinin proteins, would cleave them from the virus inhibiting its ability to invade host cells and to proliferate. Further, hydrolysis of the hemagglutinin structure would prevent this viral protein from stimulating damaging cytokine fluxes. Independent studies have confirmed the ginger enzyme inhibits H5N1 bird flu (Selleck, P "Efficacy of Zingibain in inactivating H5N1 Avian Influenza Virus" Report July 2007- A/chicken/Vietnam/8/2004 **H5N1**)

Papilloma Virus

All capsid proteins of papilloma viruses are proline-rich with a high degree of conservation of the proline peptides. The known structures of the proteins associated with common warts and with anogenital papilloma infections are as follows. The proline peptides are highlighted:

PV L1A Sequences-warts common

CPV1	. MWRP <small>SD</small> NKLYVPPAPVSKVLTDDAYVTRTKIFYHASSRLLAVGHPYFPIRK.....ANKTIVPKVSGFQF	67
RHPV1R	MSMWRP <small>SD</small> SKVYLPP...PVSKVVSTDEYVSRYSIYHAGSSRLLAVGHPYVAVKK..GNNKVSVPKVSGLQY	68
HPV29	MALWRP <small>SD</small> NLVLYLPP.TPVSKVISTDD....YVTRTNIYYAGSSRLLTVGHPHYYSIKK...SNNKVAVPKVSGYQY	70
HPV2a	MALWRP <small>SD</small> NKLYLPP.TPVSKVI..STDVYVTRTNVYYHGGSSRLLTVGHPHYYSIKK...SNNKVAVPKVSGYQY	69
HPV27	MALWRP <small>SD</small> NKLYLPP.TPVSKVI..STDVYVTRTNVYYHGGSSRLLTVGHPHYYSIKK..GNNRLAVPKVSGYQY	70
HPV57	MAMWRP <small>SD</small> NKLYLPP.TPVSKVI..STDVYVTRTNVYYHGGSSRLLTVGHPHYYSIKK..SGNNKVSVPKVSGYQY	70
HPV26	MALWRTSDSKVYLPP.TPVSRVVTDE...YVTRTGIIYYAGSSRLLTVGHPYFSIPK...TGQKAEIPKVSAAYQY	69

CPV1	RVFKIVL...PDPNKFALPDTISIFDSTSRQLVWACI...GLEVGRGQPLGVGVCYGHPCLNKFDDVENSASYAVNPGQDNR	141
RHPV1R	RVFRVRLPDPNKFGLPDANFYDPNTQRLVWACLVGEVGRGQPLGVGTSGHPLLNKLDDEENGPKVAGGGQADNR	142
HPV29	RVFRVRLPDPNKFGLPDARIYNPEAERLVWACTGVEVGRGQPLGVGLSGHPLYNKLDDEENGPKVAGGGQADNR	144
HPV2a	RVFHVKLDPDNKFGLPDADLYDPDTQRLVWACLVGEVGRGQPLGVGVSGHPYNNRLDDEENGPKVAGGGQADNR	141
HPV27	RVFHVKLDPDNKFGLPDADLYDPDTQRLVWACLVGEVGRGQPLGVGVSGHPYNNRQDDTENAHTLDS..AEDGR	142
HPV57	RVFHVKLDPDNKFGLPDANLYDPDTQRLVWACLVGEVGRGQPLGVGISGHPYNNKQDDTENSHNPKDA..ADDGR	142
HPV26	RVFRVRLPDPNKFGLPDPQLYNPDTERLVWACLVGEVGRGQPLGIGLSGHPFLNKLDDEENGPKVAGGGQADNR	143

CPV1	VNVAMDYKQTQLCLVGCAPPVGEHWGKQKCGSVSVQDGDCCPPELVTSVIQDGDMDVDTGFGAMDFAELQSNKS215	214
RHPV1R	ECVSMYDQKQTLCLMGLCKPPVGEHWGKGNPC..TTGAAGDCPPALELVNSVIQDGDMDVDTGFGAMDFAELQSNKS	214
HPV29	ENISMDYKQTLCLMGLCKPPVGEHWGKGNPC..TTGAAGDCPPALELVNSVIQDGDMDVDTGFGAMDFAELQSNKS	218
HPV2a	ENISMDYKQTLCLMGLCKPPVGEHWGKGNPC..TTGAAGDCPPALELVNSVIQDGDMDVDTGFGAMDFAELQSNKS	214
HPV27	ENISMDYKQTLCLMGLCKPPVGEHWGKGNPC..TTGAAGDCPPALELVNSVIQDGDMDVDTGFGAMDFAELQSNKS	215
HPV57	ENISMDYKQTLCLMGLCKPPVGEHWGKGNPC..TTGAAGDCPPALELVNSVIQDGDMDVDTGFGAMDFAELQSNKS	215
HPV26	DNVSVNDKQTLCLMGLCKPPVGEHWGKGNPC..TTGAAGDCPPALELVNSVIQDGDMDVDTGFGAMDFAELQSNKS	217

CPV1	DVPLDICTSTCKYPDYLLQMAADPYGDRLLFFYLREKQMFARHFFNRAGTVGEQIPDELVFKGTT...SRATVSSN	286
RHPV1R	DVPLDICTSVCKYPDYLLKMSADPYGDSLFFYLRRQMFVRLHFNAGTMGDSVPDDLYIKGSG...SNVCLASH	285
HPV29	DVPLDICTSTCKYPDYLLQMAADPYGDSMFFLRREQLFARHFFNRAGTVGEQIPDELVFKGTT...SRATVSSN	289
HPV2a	DVPLDICTNTCKYPDYLLKMAAEPYGDSMFFLRREQLFARHFFNRAGTVGEQIPDELVFKGTT...SRATVSSN	284
HPV27	DVPLDICTNVCKYPDYLLKMAAEPYGDSMFFLRREQLFARHFFNRAGTVGEQIPDELVFKGTT...SRATVSSN	285
HPV57	DVPLDICTNICKYPDYLLKMAAEPYGDSMFFLRREQLFARHFFNRAGTVGEQIPDELVFKGTT...SRATVSSN	285
HPV26	DVPLDISQSTCKYPDYLLKMSADTYGNSMFFLRREQLFARHFFNRAGTVGEQIPDELVFKGTT...SRATVSSN	289

CPV1	IYFNTPSGSLVSSAQLFNKPYWHLKAQGHNNICWGNTLVFVTDTRSTNMTVCATSTSSP...SATYASE	357
RHPV1R	IYFNTPSGSMVTSDAQLFNKPYWHLKAQGHNNICWGNTLVFVTDTRSTNMTVCATSTSSP...SATYASE	356
HPV29	IYFNTPSGSMVTSDAQLFNKPYWHLKAQGHNNICWGNTLVFVTDTRSTNMTVCATSTSSP...SATYASE	360
HPV2a	IYFNTPSGSMVTSDAQLFNKPYWHLKAQGHNNICWGNTLVFVTDTRSTNMTVCATSTSSP...SATYASE	353
HPV27	IYFNTPSGSMVTSDAQLFNKPYWHLKAQGHNNICWGNTLVFVTDTRSTNMTVCATSTSSP...SATYASE	355
HPV57	IYFNTPSGSMVTSDAQLFNKPYWHLKAQGHNNICWGNTLVFVTDTRSTNMTVCATSTSSP...SATYASE	354
HPV26	IYFNTPSGSMVTSDAQLFNKPYWHLKAQGHNNICWGNTLVFVTDTRSTNMTVCATSTSSP...SATYASE	360

CPV1	YKQYMRHVEEFDLQFIFLQCLIKTAEIMAYIHTMNPVLEEWVFLGSPPPNGTLEDYRYVQSQAITCQK.P.	429
RHPV1R	FKEYLRHVEEFDLQFIFLQCLIKTAEIMAYIHTMNPVLEEWVFLGSPPPNGTLEDYRYVQSQAITCQK.P.	428
HPV29	IKEYLRHVEEFDLQFIFLQCLIKTAEIMAYIHTMNPVLEEWVFLGSPPPNGTLEDYRYVQSQAITCQK.P.	432
HPV2a	FKEYLRHMEEYDLQFIFLQCLIKTAEIMAYIHTMNPVLEEWVFLGSPPPNGTLEDYRYVQSQAITCQK.P.	425
HPV27	FKEYLRHMEEYDLQFIFLQCLIKTAEIMAYIHTMNPVLEEWVFLGSPPPNGTLEDYRYVQSQAITCQK.P.	427
HPV57	YKEYLRHMEEYDLQFIFLQCLIKTAEIMAYIHTMNPVLEEWVFLGSPPPNGTLEDYRYVQSQAITCQK.P.	426
HPV26	YKQFIRHGEEYDLQFIFLQCLIKTAEIMAYIHTMNPVLEEWVFLGSPPPNGTLEDYRYVQSQAITCQK.P.	432

CPV1	T.PDKEKQDPYAGLSFWEVNLKEKFSSELEQYPLGRKFLLQTVGQSTSLARAG...TKRAA...STST.ATP	493
RHPV1R	A.PPKKEKQDPYAGLSFWEVNLKEKFSADLDQFPLGRKFLLQAGMRAPTLRAP...KRTAS...STSS.SPR	493
HPV29	L.APTEKQDPYAKLNFWDVLDKDRFTLDSQFPLGRKFLLQAGARRRSVPSR...KRRT...TTTAPTPA	496
HPV2a	T.PPKTPTDPYANMTFWDVLDRESFMDLDQFPLGRKFLLQAGMRAPTLRAP...KRTAS...STSS.SPR	487
HPV27	T.PPKTPTDPYANMTFWDVLDRESFMDLDQFPLGRKFLLQAGMRAPTLRAP...KRTAS...STSS.SPR	479
HPV57	T.PPKTPTDPYATMTFWDVLDSEFSMDLDQFPLGRKFLLQAGMRAPTLRAP...KRTAS...STSS.SPR	487
HPV26	A.PPVKEDPFPQKFKFWDVLDKDRFTLDSQFPLGRKFLLQAGMRAPTLRAP...KRTAS...STSS.SPR	494

CPV1	TR.KKVKRK.....	501
RHPV1R	KR.KRTKR.....	500
HPV29	KR.KRSKK.....	503
HPV2a	KR.KRVRR.....	494
HPV27	AV.GRGH.....	485
HPV57	KR.KKVR.....	494
HPV26	KR.KKRLTK....	503
HPV52	KK.KVKR.....	503
HPV58	KR.KKVKK.....	498
HPV67	RK.KVKR.....	500

HPV L1A Sequences-Anogenital

HPV32	MSVWRP <small>SD</small> NKLYLPP.PPVSKVVSTDEYVQRTNIFYHASSRLLAVGHPYTIKK....TPNRTSIPKVSGLQY	69
HPV11R	.. MWRP <small>SD</small> STVYVPPPNPVSKVVATDAYVTRTNIFYHASSRLLAVGHPYYSIKK...YNKTVVPKVSGYQY	67
HPV6bR	.. MWRP <small>SD</small> STVYVPPPNPVSKVVATDAYVTRTNIFYHASSRLLAVGHPYYSIKR...ANKTVVPKVSGYQY	67
HPV18R	MALWRP <small>SD</small> NTVYLLPP.PSVARVVNTDD...YVTRTSIFYHAGSSRLLTVGHPYFVAVPAGGGNKQDIPKVSAAYQY	70
HPV16R	MSLWRP <small>SE</small> ATVYLLPP.VPVSKVVSTDEYVTRTNIFYHAGSARLLTVGHPYYSIKK...PNNNKILVVKVSGLQY	70
HPV31	MSLWRP <small>SE</small> ATVYLLPP.VPVSKVVSTDEYVTRTNIFYHAGSARLLTVGHPYYSIKK...PNNNKILVVKVSGLQY	71
HPV33	MSVWRP <small>SE</small> ATVYLLPP.VPVSKVVSTDEYVSRYSIYHAGSSRLLAVGHPYFSIKNPTNAKLLVVKVSGLQY	71

HPV32	RVFRVRLPDPNKFALPDTISYNPETQRMVWACVLEVGRGQPLGVGLSGHPLLNRLDDEENGPKVAGGGQADNR	143
HPV11R	RVFKVVLDPDNKFALPDSLSFDPTTQRLVWACT..GLEVGRGQPLGVGVSGHPLLNKYDDVENSAGGYPGQDNR	141
HPV6bR	RVFKVVLDPDNKFALPDSLSFDPTTQRLVWACT..GLEVGRGQPLGVGVSGHPLLNKYDDVENSAGGYPGQDNR	140
HPV18R	RVFRVRLPDPNKFGLPDTISYNPETQRLVWACVLEVGRGQPLGVGLSGHPLFNKLDDEENGPKVAGGGQADNR	144
HPV16R	RVFRVRLPDPNKFGLPDTISYNPETQRLVWACVLEVGRGQPLGVGLSGHPLLNKLDDEENGPKVAGGGQADNR	144
HPV31	RVFRVRLPDPNKFGLPDTISYNPETQRLVWACVLEVGRGQPLGVGLSGHPLLNKLDDEENGPKVAGGGQADNR	145
HPV33	RVFRVRLPDPNKFGLPDTISYNPETQRLVWACVLEVGRGQPLGVGLSGHPLLNKLDDEENGPKVAGGGQADNR	145

HPV32	ENVMDCQKQTLCLVGCAPPVGEHWGKGAACSA..QSNDCPPPELVNSVIQDGDMDVDTGFGAMDFALQTSKA	215
HPV11R	VNVGMDYKQTLCLMVGCAAPPVGEHWGKGTQCSNTSVQNGDCPPPELVNSVIQDGDMDVDTGFGAMDFALQTSKA	215
HPV6bR	VNVGMDYKQTLCLMVGCAAPPVGEHWGKGTQCSNTSVQNGDCPPPELVNSVIQDGDMDVDTGFGAMDFALQTSKA	214

HPV18R	DNVSDVYKQTQLCLGCA P AIGEHWAKGTACKS RPL SQGD CP PLELKNTVLEDGDMVDTGYGAMDFSTLQDTKC	218
HPV16R	ECISMDYKQTQLCLIGCK PP IGEHWGKGS SP CTNVA VNP GD CP PLELINTVIQDGDGMVDTGFGAMDFTTLQANKS	218
HPV31	ECISMDYKQTQLCLLIGCK PP IGEHWGKGS SP CNNAI TP GD CP PLELKNVSIQDGDGMVDTGFGAMDFTALQDTKS	219
HPV33	ECLSMYKQTQLCLLIGCK PP TGEHWGKGVACTN AA PAND CP PLELINTIIEDGDMVDTGFGCMDFKTLQANKS	218
HPV32	EV PL DIMNSISKY PD YLKMSAEAYGDNMFFFLRREQMFVRHLFN RAGTLGEPVPE DMYIKASNGASGRNNLASS	289
HPV11R	DV PL DICGTVCCKY PD YLQMAAD PY GDRLFFFLRKEQMFARHFFNRAGTV GEFV PDLLVKGGN...NRSSVASS	286
HPV6bR	DV PD ICGTTCCKY PD YLQMAAD PY GDRLFFFLRKEQMFARHFFNRAGEV GEFV PDLLIKGSG...NRTSVGSS	285
HPV18R	EV PL DICQSICKY PD YLQMSAD PY GDSMFFCLRREQLFARHFFNRAGTMGD TV PQSLYIKGTG...MRAS SP GSC	289
HPV16R	EV PL DICTSICKY PD YIKMVSE EPY GDSLFFFLRREQMFVRHLFN RAGTVGENV PDDLYIKGSG...STANLASS	289
HPV31	NV PL DICNSICKY PD YLKMA EPY GDTLFFFLRREQMFVRHFFNRSGTV GESV PTDLYIKGSG...STATLANS	290
HPV33	DV PD ICGTTCCKY PD YLKMTS EPY GDSLFFFLRREQMFVRHFFNRAGTL GEAV PDDLYIKGSG...TTASIQSS	289
HPV32	IYY PT PSGSMVTSDAQIFNK PY WLQQAQGHNNNGICWGNQVFLTVVD TTR STNMTVCATVTTED.....TYKSTN	358
HPV11R	IYVHT PS GSLVSSEAQLFNK PY WLQKAQGHNNNGICWGNHFLVTVVD TTR STNMTLCASVSKSA.....TYTNSD	355
HPV6bR	IYVNT PS GSLVSSEAQLFNK PY WLQKAQGHNNNGICWGNQVFLTVVD TTR STNMTLCASVTTSS.....TYTNSD	354
HPV18R	VY SP SPSGSIVTSDSQLFNK PY WLHKAQGHNNNGVCWHNQVFLTVVD TTR STNLTICASTQ SPV ...PGQYDATK	360
HPV16R	NY FPT PSGSMVTSDAQIFNK PY WLQQAQGHNNNGICWGNQVFLTVVD TTR STNMSLCAAISTSE.....TTYKNTN	359
HPV31	TY FPT PSGSMVTSDAQIFNK PY WMQRAQGHNNNGICWGNQVFLTVVD TTR STNMSVCAAANSND.....TTFKSSN	360
HPV33	AF FPT PSGSMVTSSESQFLFNK PY WLQQAQGHNNNGICWGNQVFLTVVD TTR STNMTLCTQVTSDS.....TYKNEN	358
HPV32	FKEYLRHAEEDYDQIFLQCKITLSVEVMSYIHTM NP DILDDWNVGV APPP SGTLEDSYRFVQSQAIRCOA.K	430
HPV11R	YKEYMRHVEEFDLQFIFQLCSITLSAEVMAYIHTM NP SVLEDWNFGL SPPP NGTLEDYRYVQSQAITCQK P	427
HPV6bR	YKEYMRHVEEYDLQFIFQLCSITLSAEVMAYIHTM NP SVLEDWNFGL SPPP NGTLEDYRYVQSQAITCQK P	426
HPV18R	FKQYSRHHVEEYDLQFIFQLCKITLADVMYIHSMNSSILEDWNFV GVPPP TTSLVDYRFVQSQVAITCQK.D	432
HPV16R	FKEYLRHGEEYDLQFIFQLCKITLADVMYIHSMNSTILEDWNFGL QPPP GGTLEDYRFVTSQAIACQK.H	431
HPV31	FKEYLRHGEEFDLQFIFQLCKITLSADIMTYIHSMN NP AILEDWNFGL TPPP SGLEDYRFVTSQAITCQK.T	432
HPV33	FKEYIRHVEEYDLQFVFLQCKVTLTAEVMTYIHAM NP DILEDWQFGL TPPP SASLQDQTYRFVTSQAITCQK.T	430
HPV32	V.TA PE KK DP FSDYSFWEVNLSSEKFSDDLQ FL GRKFLLQAGLRAR PK LTAV....KRTASS.SQKSS S .PAK	497
HPV11R	T. PE KEK QDP PKDMSFWEVNLSSEKFSSELQ FL GRKFLLQSGYRGRSARTG...IKR PAVSK PS .TAP	492
HPV6bR	T. PE KEK QDP PKNLSFWEVNLSSEKFSSELQ FL GRKFLLQSGYRGRSSIRTG...VKR PAVSKAS.AAP	491
HPV18R	A.APAENK DP YDKLFKWNVDLKEKFSADLDQ FL GRKFLLVQAGLR PK TI GRKRSAP..SATTSS.K PA	498
HPV16R	T. PP AP KEDP LKKYTFWEVNLKEKFSADLDQ FL GRKFLLQAGLKA KPK FTLG....KRKAT P .TTSSTS.TTA	498
HPV31	A. PQ K P KED P FKDYVFWEVNLSSEKFSADLDQ FL GRKFLLQAGYR ARPK FKAG...KRSAP..SASTT. TPA	497
HPV33	V. PK KEK EDP LKGYTFWEVNLKEKFSADLDQ FL GRKFLLQAGLKA KPK LKR....AAT TSTRT.SSA	492
HPV32	RR.KTRK.....	503
HPV11R	KR.KRTKTKK....	501
HPV6bR	KR.KRAKTKR....	500
HPV18R	KR.VRVRARK....	507
HPV16R	KR.KKRKL.....	505
HPV31	KR.KKTKK.....	504
HPV33	KR.KKVKK.....	499

The multiple highlighted proline peptides (and the other prolines which lack adjacent hydrophilic amino acids) render these proteins inert to normal hydrolysis. These intact proline-rich peptides in which the prolines are adjacent to hydrophilic amino acids are ideal epitopes for triggering a severe immune response and for binding to host cells. The Biohawk ginger with its high level of enzyme has the potential to cleave these peptides, resulting in breakdown of the viral capsid and structural proteins, prevention of host cell invasion and avoidance of immune system stimulation and the associated tissue damaging cytokine flux. The multiple exposed sites for the ginger enzyme cleavage on one side of the protein are coloured orange in the following model of the HVP16-L1 protein (Figure 3).

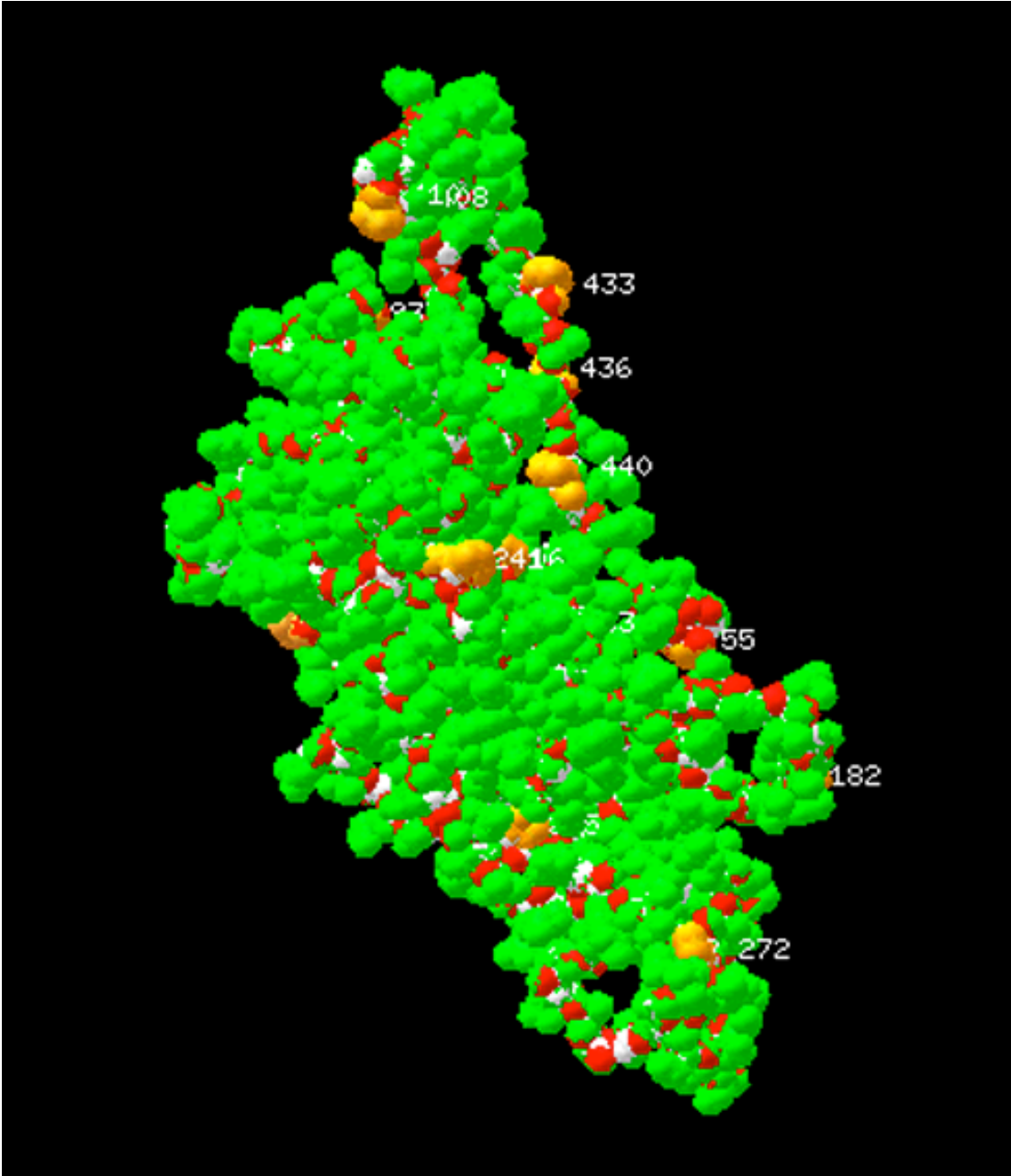


Figure 3. A model of the HPV16-L1 capsid protein responsible for generating a number of forms of squamous cell carcinomas. In the present picture green represents the surface groups and the backbone has been shown in red and white. The suitable prolines for ginger enzyme cleavage are shown in orange-yellow. The numbers represents the proline number in the HPV16-L1 protein.

Papilloma virus is of particular interest to Biohawk because it is associated with disease in humans and other species, for example equine sarcoid, bovine eye cancer, common warts, corns, some squamous cell carcinomas, and cervical cancer. Case studies have shown a very favourable response to treatment with the Biohawk creams (for example, Skin Rejuvenator). For example, the following common warts (Figure 4) were treated with a ginger cream once at 3 pm on one day and the next morning the growths had fallen off the hand.



Figure 4: Common wart treated with an active ginger cream

Bacteria

Some bacteria have a carbohydrate coating on the membrane mostly attached to the membrane proteins through the amino acids, serine (highlighted pink) and threonine (highlighted grey). This coating makes it more difficult for antibiotics to penetrate the membrane and makes it more difficult for the ginger enzyme to digest the proline-rich proteins. Biohawk's Pine Crush was developed to remove the carbohydrate coating. In other types of bacteria the proline-rich proteins project out of the membrane and although they may have serine and threonine, the carbohydrate coating does not prevent the ginger enzyme from digesting the proline-rich membrane proteins. Some examples of membrane proteins for common types of bacteria are given below.

Whooping Cough: *Bordetella pertussis*

Bordetella pertussis, the causative agent of whooping cough, is an aerobic coccobacillus capsule of the genus *Bordetella*. The acellular pertussis vaccine components, pertussis toxoid (PT), pertussis filamentous haemagglutinin (FHA) and pertactin (PRN) are extracted from phase I *Bordetella pertussis*, and are then purified and stabilised. The structures of FHA and PRN are given below.

FHA protein [*Bordetella pertussis*]

1 MNTNLYRLVF SHVRGMLVPV SEHCTVGNTEF CGRTRGQARS GARATSLVA PNALAWALML
61 ACTGLPLVTH AQLV PQGQT QVLQGGNKVP VVNIADPNSG GVSHNKFQQF NVANPGVVFN
121 NGLTDGVSRI GGALTKNP NL TRQASAILAE VTDTPSPRLA GTLEVYGKGA DLIANPNGI
181 SVNGLSTLNA SNLTLTTGRP SVNNGGRIGLD VQQGTVTIER GGVNATGLGY FDVVARLVKL
241 QGAVSSKQGGK PLADIADVAG ANRYDHATR ATPIAAGARG AAAGAYIDG TAAGAMYGKH
301 ITLVSSD SGL GVRQLGSLSSPSAITVSSQG EIALGDATVQ RGPLSLKGAG VV SAGKLASG
361 GGAVNVAGGG AVKIASASSV GNLA VQGGGK VQATLLNAGG TLLVSGRQAV QLGAASSRQA
421 LSVNAGGALK ADKLSATRRV DVDGKQAV AL GSASSNALS V RAGGALKAGK LSATGRLDVD
481 GKQAVTLG SV ASDGALSVSA GGNLRANELV SSAQLEVRGQ REVALDDASS ARGMTVVAAG
541 ALAARNLQSK GAIGVQGGEA VSVANANS DA ELRVRGRGQV DLHDL SAARG ADISGEGRVN
601 IGRARSDSDV KVS AHGALS I DSMTALGAIG VQAGG SVSAK DMR SRGAVTV SGGGAVNLGD
661 VQSDGQVRAT SAGAMTVRDV AAAADLALQA GDALQAGFLK SAGAMTVNGR DAVRLDGAHA
721 GGQLRVSSDG QAALGSLAAK GELTVS AARA ATVAELKSLD NISVTGGERV SVQSVNSASR
781 VAISAHGALD VGKVS AKSGI GLEGWGAVGA DSLGSDGAI S VSGRDAVRVD HARSLADISL 841
GAEGGATLGA VEAAGSIDVR GGS TVAANS L HANRDVRVSG KDAVRVTAAT SGGGLHVSSG
901 RQLDLGAVQA RGALALDGGG AVALQSAKAS GTLHVQGGEH LDLGTLAAVG AVDVNGTGDV
961 RVAKLVS DAG ADLQAGRSMT LGIVDTTGD L QARAQQKLEL GSVKSDGGLQ AAAGGALS LA
1021 AAEVAGALEL SGQGVTVDRA SASRARIDST GSVGIGALKA GAVEAASPRR ARRALRQDF
1081 TPGSVVVRAQ GNVTVGRGDP HQGVLAQGGI IMDAKGGTLL LRNDALTENG TVTISADSAV
1141 LEHSTIESKI QSVVLAAGD KGKPAVSVKV AKKLFNLGTL RAVNDNNETM SGRQIDVVDG
1201 RPQITDAVTG EARKDES VVS DAALVADGGP IVVEAGELVS HAGGIGNGRN KENGASVTVR
1261 TTGNLVNKG Y ISAGKQGVLE VGGALTNEFL VGS DGTQRIE AQRIENRGTF QSQAPAGTAG
1321 ALVVKAAEAI VHDGVMATKG EMQIAGKGGG SPTVTAGAKA TTSANKLSVD VASWDNAGSL
1381 DIKKGGAQVT VAGRYAEHGE VSIQGDYTVS ADAIALAAQV TQRGGAANLT SRHDTRF SNK
1441 IRLMGP LQVN AGPVSNTGN LKVREGVTVT AASFDNETGA EVMAKSATLT TSGAARNAGK
1501 MQVKEAATIV AASVSNPGTF TAGKDITVTS RGGFDNEGKM ESNKDIVIKT EQFSNGRVL
1561 AKHDLTVTAS GQADNRGSLK AGHDFTVQAQ RIDNSGTMAA GHDATLKAPH LRNTGQVVAG
1621 HDIHIINSAK LENTGRVDAR NDIALDVADF TNGTSLYAEH DATLTLAQQT QRDLVVDQDH
1681 ILPVAEGTLR VKAKSLTTEI ETGNPGLIA EVQENIDNKQ AIVVGKDLTL SSAHGNVANE
1741 ANALLWAAGE LTVKAQNITN KRAALIEAGG NARLTAVAL LNKLGRRIRAG EDMHLDAPRI
1801 ENTAKLSGEV QRKGVQDVG GEGHGRWSGIGYVNYWLRAGNGKKAGTIAAPWYGGDLTAEQ
1861 SLIEVGKDLY LNAGARKDEH RHLLNEGVIQ AGGHGHIGGD VDNRSVVRTV SAMEYFKT PL
1921 PVSALTALDNR AGLSPATWNF QSTYELLDYL LDQNRYEYIW GLYPYTEWS VNTLKNLDLG
1981 YQAKPAPTAP PMPKAP ELDL RGHTLESAEG RKIFGEYKKL QGEYKAKMA VQAVEAYGEA
2041 TRRVHDQLGQ RYGKALGGMD AETKEVDGII QEFAADLRTV YAKQADQATI DAETDKVAQR
2101 YKSQIDAVRL QAIQPGRVTL AKALS AALGA DWRALGHS QL MQRWKDFKAG KRGAEIAFY P
2161 KEQTVLAAGA GLT LSNGAIH NGENAAQNRG RPEGLKIGAH SATSVSSG SFD ALRDVGLEKR
2221 LDIDDALAAV LVNPHIFTRI GAAQTS LADG AAGPALARQA RQAPETDGMV DARGLGSSADA
2281 LASLASLDAA QGLEVS GRRN A QVADAGLAG PS AVAAPAVG AADVGV EPVT GDQVDQPVVA
2341 VGLEQPVATV RVAPPAVAL P RPLFETRIKF IDQSKFYGS R YFFEQIGY KP DRAARVAGDN
2401 YFDTTLVREQ VRRALGGYES RL PVRGVALV AKLMD SAGTV GKALGLKVG V APTAQQLKQA
2461 DRDFVWYVDT VIDGQKVLAP R LYLTEATRQ GITDQYAGGG ALIASGGDVT VNTDGHVSS
2521 VNGLIQGRSV KVDAGKGVV VADSKGAGGG IEADDEVDS GRDIGIEGK LRGKDVRLKA

2581 DTVKVA^T^SMR YDDKGR^LLAAR GDGALDAQGG QLHIEAK^RLE TAGATL^KGGK VKLDVDDV^KL
 2641 GGVYEAG^SSY ENK^SST^PLG^S LFAIL^SSTTE TNQ^SSAHANHY GTRIEAG^TLE GKMQNLEIEG
 2701 G^SVDA^AH^TDL^S VARDAR^FKA AADFAHAEHE KDVR^QL^SLG^A KVGAGGYEAG F^SLG^SES^GLE
 2761 AHAGRG^MTAG AEVKVG^YRAS HEQ^SSETEK^S YRNANLNFGG G^SVEAGNVLD IGGADIN^RNR
 2821 YGGA^AKNAG TEEAL^RMRAK KVE^STKYV^SE QT^SQSS^GW^SV EVA^STAS^ARS^SLLTAAT^RLG
 2881 D^SV^AQNV^EDG REIR^GELMAA QVAAEATQLV TADTAAVAL^S AGI^SADFD^SSS^HSR^STS^SQNTQ
 2941 YLGGNL^SIEA TEGDATLVGA KFGGGDQV^SL KAAK^SVNLMA AE^STFE^SSY^SE SHNFH^ASADA
 3001 NLGANAVQGA VGLGLTAGMG T^SHQITNETG KTYAGT^SVDA ANV^SIDAGKD LNL^SGS^RVRG
 3061 KHV^VLDVEGD INAT^SKQDER NYN^SSGGGWD ASAGVAIQ^NR TLVAP^VGS^AG FNFNTEHDN^S
 3121 RL^TNDGAAGV VASDGLTGHV KGDANLTGAT IADL^SGKGNL KVDGAVNAQN LKDYRDKDGG
 3181 ^SGG^LNVGI^SSS^TTLAP^TVGVA FGRVAGEDYQ AEQRATIDVG QTKD^PARLQV GGGVKGTLNQ
 3241 DAAQATV^VQR NKHWAGGG^SE F^SVAGK^SSLKK KNQV^RPVET^PTPDVVDG^PPS^RPTT^PASP^Q
 3301 PIRATVEV^SSS^PPP^VS^VATVE VV^PRP^KVETG SAASASAGGA QVV^PVT^PPK^VEVAKVEV^VPR
 3361 PKVETAQ^LPP^PRP^VVAEKVT TPAVQPQLAK VETVQ^PVKPE TTK^PL^PK^PL^PVAKVTKA^PPP
 3421 VVETAQ^LPP^PVK^PQKAT^PGP VAEVGKATVT TVQVQ^SAPP^KPAP^VAKQ^PAP^APK^PK^PK^PK^P
 3481 KAER^PK^PGKT T^PL^SGRHV^VQ QVQVLRQA S^DINNTK^SL^PGGKL^PK^PVTV KLTDENG^KP^Q
 3541 TY^TINRR^EDL MKLNGKVL^ST KTTLGLEQTF RLRVEDIGGK NYRVFYETNK

pertactin outer membrane protein [Bordetella pertussis]

1 MNM^SLS^RIVK AAP^LRR^TT^LA MALGALGAAP AAHADWNNQ^S IVKTGERQHG IHIQGS^DPGG
 61 VRTASG^TTIK V^SGRQAQ^GIL LE^NPAELQF R^NGS^VTSS^GQLS^DDDGIR^RFL GTVTVKAGK^L
 121 VADHATLANV GDTWDDDDGIA LYVAGEQAQA SIAD^STLQGA GGVQIERGAN VTVQR^SAIVD
 181 GGLHIGALQ^SLQPE^DLPP^SR VVLRD^TNVTA VPASGAPAAV^SVLGASEL^TL DGGHITGGRA
 241 AGVAAMQGA^VVHLQRATIRR GDAPAGGAV^PGGAV^PGGAV^PGGF^GPGG^FPG^VLDGWYGV^DV
 301 ^SSS^SVELAQ^SIVEAPE^LGAA IRVGRGARVT VSGG^SLS^APH GNVIE^TGGAR RFAPQAA^PLS
 361 I^TLQAGAHQ^G GKALLY^RVLP EP^VKLTLTGG ADAQGDIVAT EL^PSIP^GTSI GPLDVALAS^Q
 421 ARWTGATRAV^DLS^IDNATW VMTD^NSNVGA LRLASDGS^VD FQQA^EAGR^F KVLTVNTLAG
 481 ^SGLFRMN^VFA DLGL^SDKLVV MQDASGQHRL WVRN^SSG^SEPA SANTLLL^VQ^TPLGSAAT^FTL
 541 ANKDGKVDIG TYRYRLAANG NGQWSLVGAK APP^PK^PAP^QPG^QPP^QPP^QPP^QPP^QPEAP^AP^Q
 601 PAGREL^SAAA NAAVNTGGVG L^ASTLWYAE^SNALS^KR^LGEL RL^NPDAGGAW GRGFAQRQ^L
 661 DNRAGRRFDQ KVAGFELGAD HAVAVAGGRWHLGGLAGYTRGDRGFTGDGGGHTD^SVHVGG
 721 YATYIAD^SGF YLDATLRA^SR LENDFKVAG^SDGYAVK^GKYR THGVGAS^LLEA GRRFTHADGW
 781 FL^EPQAE^LAV FRAGGGAYRA ANGLRVRDEG GSSVLGRLGL EVGKRIELAG GRQV^QPYIKA
 841 ^SVLQEFDGAG TVHTNGIAHR TELRGTRAEL GLGMAAALGR GHS^LYAS^SYEY^SSKG^PK^LAMP^P
 901 TFHAGYRYS^W

Homology in sequences of above proteins

Pertactin: PKPAP- - - QPGPQPPQPPQPPQPEAPAPQP
 PKPAP- - - QP-P- P- - - P-P -P- A - -P- P
 FHA protein: PKPAPVAKQPAPAPKPKPK PKPKAERPKP

Staphylococcus aureus

Cap5P [Staphylococcus aureus]

1 MCLNFREDNV MKKIMVIFGT RPEAIKMAPL VKEIDHNGNF EANIVITAQH RDMLDSVLSI
 61 FDIQADHDLN IMQDQQTLAG LTANALAKLD SIINEEQPDM ILVHGDTT^TTVGSLAAFYH
 121 QIPVGHVEAG LRTHQKYS^PF PEELNRVMV^S NIAELNFAP^T VIAAKNLLFE NKDKERIFIT
 181 GNTVIDAL^ST TVQNDFV^STI INKHKGK^KVI LLTAHRRENI GEP^MHQIFKA VRDLADEYKD
 241 VVFIY^PPMHRN PKVRAIAEKY LSGRNRIELI EPLDAIEFHN FTNQS^SYLVL^TDSGGIQEEAP
 301 TFGK^PVLVLR NHTERPEGVE AGTSRVIGTD YDNIVRNVKQ LIEDEAYQR MSQANNPYGD
 361 GQAS^SR^RICEA IEYFGLR^TD KPDEFVPLRH K

Protein A signal fusion protein.

1 MKKKNIYSIR KLGVGIA^SVT LGTLLIS^SGGV TPAANAQHD EAVDNKFNKE QQNAFYEILH
 61 L^PN^LN^EEQRN AFIQ^SSLKDD^PSQSANLLAEA KKL^NDAQAPK VDNKFNKEQQ NAFYEILHL^P
 121 ^NL^NE^EEQRNAF IQ^SSLKDD^PSQSANLLAEA^KKL^NDAQAPK^VD AN^SSS^SVPG^DP LE^STCR^HAS^L
 181 ALAVVLQRRD WEN^PGVTQLN RLA^AHPP^FAS WRN^SSEEARTD RPSQQLRS^SLN^GEWRFRCNGW
 241 R

Biofilm-associated surface protein [Staphylococcus aureus]

1 MGNKQGFLPN KLNKYSIRKF TVGTASLLVG TTLFFGIGSE AQAALDITIT KEDVKSQDKG
61 EALDIKNIKESEKDVTTEDDNNAEVQNSAQTVDKSENSNDTAVESTNDSVKTDETKETS
121 NKSAAQDDDNIKEDSNTQEEESTNTSSQSSSEVPQTKKDTNETSETAIDEDASTKEQNNKDND
181 TAQDDDNIKE DNTQEEESTN TSSQSSSEVPQ TKKEQPKSS NSIKEPKQKQ EEEVAKKEKAI
241 TEIADKNKEL ELKNNKTDKN EESSELESNLSSENKKTVE SFLNSQLSDS ETKKIMENAN
301 IDYDKATDEE INTEILRASL IEMANNKKT ETLATPQRTM FRAMATPTAL RAAVNDQEEL
361 QKSLGYTDNY TFA SMLFDPG KLD SDDALNS NIIPFDLHYS MSGANSGNRY KIDLKLDPII
421 AEHVTKISAN PSGSNKPVEF VRNKDENGNL TDTWEVNFIR ANDGLFGGAE ILSQYTAKNG
481 KIELDDTVGN IISNAGNLSN NKLNHQVFVR DSRENKIVRT SSSGYFLTK ADDDLVNLN
541 NVSTENNAF KASSGSATYN ENVGEFGGIL IDQQIMKNGI FYSKTKANQ WAYNYQIDKD
601 LLPYIEGVEL HQDYDKGLNG FDKNYDAKNK VADLTIDEVG NGTITSDNLN KLIEFNALP
661 ETVGVRVVLK LNKSVNNILT KDAKYDSEGN LIRETTKQKE DFTFAGYLTDSK GALINNTL
721 GTSTLALQDY DKDGLLDREY RQLSLSDAEN EDTDGDGKND GDEVVNYKTSPLVKGPOAAD
781 ITTEDTVVSG SVPLKEGAAT QTAKVINAEG TTVGTATVNS DGTFSVSI PN SPEGTYTIAI
841 DSPNYDNDEV NTFEIVDNSK LPAPSPINPVD DNDQQIVVNG TSGSTVTVD SNNNVLTGVT
901 IPADDTSAAI NVDTPLEAGT VLTSTASKDG KTS DVSQDQIT VDATAPDAP TLDEVNTDAT
961 QVTGQAE PNSTV KLTTFPDGT TATGTADDQG NYTIDI PPSNV DLNGGEELQV TATDKDGNTS
1021 EPSSANVTD TAPDAPT VND VTS DATQVTG QAEPNSTVKL TFPDGT TATGTADDQGN YTI
1081 DIPSSNV DLNG GEELQVTATD KDGNTSE PPS ANVTDTTAPD APTVNDVTS DATQVTGQAE P
1141 NSTVKLTF PD GTTATGTADD QGN YTI DI PPS NVDLNGGEEL QVTATDKDGNTSE PPS ANVT
1201 DTTAPDAPT VNDVTS DATQV TGQAE PNSTV KLTTFPDGTTA TGTADDQGN YTI DI PPS NVDL
1261 NGGEELQVTA TDKDGNTSE PPS ANVTDTTA PDAPT VNDVT SDATQVTGQAE PNSTVKLTF
1321 PDGTTATGTA DDQGN YTI DI PPS NVDLNGGE ELQVTATDKD GNTSE PPS ANVTDTTAPDAP
1381 TVNDVTS DAT QVTGQAE PNSTV KLTTFPDGT TATGTADDQG NYTIDI PPS NVDLNGGEELQV
1441 TATDKDGNTS EPSSANVTD TAPDAPT VND VTS DATQVTG QAEPNSTVKLTF PDGTTATG
1501 TADDQGN YTI DI PPS NVDLNG GEELQVTATD KDGNTSE PPS ANVTDTTA PDAPT VNDVTS D
1561 ATQVTGQAE PNSTVKLTF PD GTTATGTADD QGN YTI DI PPS NGDLNGGEELQVTATDKDGN
1621 TSE PPS ANVT DTTA PDAPT VNDVTS DATQV TGQAE PNSTV KLTTFPDGTTATGTADDQGN Y
1681 TIDI PPS NVDL NGGEELQVTA TDKDGNTSE PPS ANVTDTTA PDAPT VNDVTS DATQVTGQAE
1741 EPNSTVKLTF PDGTTATGTA DDQGN YTI DI PPS NVDLNGGE ELQVTATDKDGNTSE P KLTN
1801 VDTTASDAP TVNDVTS DAS QVTGQAE PNSTV KLTTFPDGT TATGTADDQGN YTI DI PPS NV
1861 DLNGGEELQV TATDKDGNTS EPSSANVTD TAPDAPT VND VTS DATQVTGQAE PNSTVKL
1921 TFPDGT TATG TADDQGN YTI DI PPS NGDLNG GEKLQVTATD KDGNTSE PPS ANVTDTTA PD
1981 APTVNDVTS D ATQVTGQAE PNSTVKLTF PD GTTATGTADD QGN YTI DI PPS NVDLNGGEEL
2041 QVTATDKDGN TSE PPS ANVT DTTA PDAPT VNDVTS DATQV TGQAE PNSTVKLTF PDGTTA
2101 TGTADDQGN YTI DI PPS NVDL NGGEELQVTA TDKDGNTS ESTNTTIIDSDDNS DNGNNSGA
2161 GDTSDSDDNS GNGDNS GAGD NSDSDDNS DNGNNS GAGD NSDSDDNS DNE DNSSSNKDSIN
2221 QDSSNVNSNDS KHDKQNEL PE TGEKEVRNGT LFGTLFAGLG SLLLFTKRRR KENDKK

Streptococcus agalactiae

Protein immunoglobulin-a-binding beta antigen

1 AIKQQ IFDI DNAK EVID NLVHDAF SKM NA VAKFQKG LE N PE PD PKIPELPQA
61 PD PQAPD P HVPE SPK APE APRVPE SPK PD PHVPE SPK APE APRVPE SPK PD PHV
121 PESPK APE APRVPE SPK PD PHVPE SPK APE APRVPE SPK PD PHVPE SPK APE APRV
181 PESPK APE APRVPE SPK PE APKIPPEPK PDV PKLPDV P KLPDV PKLPD APKLPDGLNK
241 VGQAVF S D GN KV VVFD KP DADKLHL KEV KELA

Penicillin binding protein 1a

1 MI IKKE SVI KLLKYAFGII MGFIILAIVI GLLFAYYV S RSPKLT DQAL KSVNSSLVYD
61 GNNKLIADLG SEKRESVSAD S IPLNLVNAI T SIEDKRFFK HRGVDIYRIL GAAWHNLVSS
121 NTQGGSTLDQ QLIKLAYFST NKS DQ LK R K S QEVWLALQM ERKY TKEEIL TFYINKVYMG
181 NGNYGMR TTA KSYFGKDLKE L SIAQLALLA GI PQAP T QYD PYKNPES AQ T RRNTVLQQMY
241 QDKNIS KKEY DQAVA P VTD GLKELKQKST YPKYMDNYLK QVIS EVKQKT GKDIF TAGLK
301 VY T NIN T DAQ KQLYDIYNS D TYIAYPNNEL QIASTIMDAT NGKVIAQLGG RHQENISVFG
361 T NQSVL T DRD WGS T MKPISA YAPAIDS GYV NSTGQSLNDS VYYWPGTSTQ LYDWRQYMG
421 WMSMQTAIQG SRNVPARAL EAAGLDEAKS FLEKLGIIYP EMNYSNAISS NNSSSDAKYG
481 ASSSEKMAAAY SAFANGTTY KPQYVNKIEF SDG TND TYAA S GSRAMKE TTAYMMT DMLKT
541 VLTFG T G KA AIPGVAQAGK T G TSNY TEDE LAKIEA T TGI YNSAVGTMAP DENFVGYT SK
601 YTMAIW T GYK NR L T PLYG S Q LDIA TEVYRA MMS YL TGGYS ADW T MPEGLY RSGSYLYING
661 TTTTGT Y SSS VYKNYQNS G QSSQSSSS S SEKQKEDKN T ANDAN SSSPQ VE TPNNGNA T

721 **T**PNNS**N**Q**T**VP**G**GHGNGNGN NN**T**VP**N**GN

Pi-2a ancillary protein 2

1 MKKIRK**S**LGL LL**CC**FLGLVQ LAFF**S**VA**S**SVN AD**TP**N**Q**L**T**IT QIGL**Q**P**N**T**T**E EGI**S**Y**R**L**W**T**V**
61 **T**D**N**L**K**V**D**L**L****S** Q**M**T**D****S**EL**N**Q**K** Y**K****S**I**L**T**S**P**T**D T**N**G**Q**T**K**I**A**L**P** N**G**S**Y**F**G**R**A**Y**K** A**D**Q**S**V**S**T**I**V**P**
121 FYIEL**P**D**D**K**L** **S**N**Q**L**Q**I**N**P**K**R K**V**E**T**G**R**L**K**L**I** K**Y**T**K**E**G**K**I**K**K** **R**L**S**G**V**I**F**V**L**Y D**N**Q**N****Q**P**V**R**F**K
181 N**G**R**F**T**T**D**Q**D**G** I**T****S**L**V**T**D**D**K**G E**I**E**V**E**G**L**L**P**G** K**Y**I**F**R**E**V**K**A**L** T**G**Y**R**I**S**M**K**D**A** V**V**A**V**V**A**N**K**T**Q**
241 E**V**E**V**E**N**E**K**E**T** **P**P**T**N**P**K**S****Q** **P**L**F****Q**S**F**L**P**K T**G**M**I**G**G**G**L**T I**L**G**C**I**L**G**I**L F**I**L**R**K**T**K**N**S
301 **K**S**E**R**N**D**T**V

Streptococcus pneumoniae

PspA [Streptococcus pneumoniae]

1 MNKKKM**L**I**T****S** L**A**S**V**A**I**L**G**A**G** F**V**A**S**S**P**T**F**V**R** A**E**E**A**P**V**A**N**Q**S** K**A**E**K**D**Y**D**A**A**V** K**K****S**E**A**A**K**D**Y**
61 E**T**A**K**K**A**E**D**A Q**K**Y**D**E**D**Q**K** T**E**A**K**A**E**K**E**R**K** A**S**E**K**I**A**E**A**T**K** E**V**Q**Q**A**Y**L**A**Y**L** Q**A**S**N**E**S**Q**R**K**E**
121 A**D**K**K**I**K**E**A**T**Q** R**K**D**E**A**E**A**A**F**A** T**I**R**T**T**I**V**V****P**E**P**S**E**L**A**E**T**K**K**K A**E**E**A**T**K**E**A**E**V** A**K**K**K****S**E**E**A**A**K
181 E**V**E**V**E**K**N**K**I**L** E**Q**D**A**E**N**E**K**K**I** D**V**L**Q**N**K**V**A**D**L** E**K**G**I**A**P**Y**Q**N**E** V**A**E**L**N**K**E**I**A**R** L**Q****S**D**L**K**D**A**E**E
241 N**N**V**E**D**I**A**E**I**K**E**G** L**E**Q**A**I**T**N**K**K**A** E**L**A**T**T**Q**N**I**D K**T**Q**K**D**L**E**D**A**E** L**E**L**E**K**V**L**A**T**L** **D**P**E**G**K**T**Q**D**E**L
301 D**K**E**A**E**A**E**A**E**L**N K**V**E**A**L**Q**N**Q**V A**E**L**E**E**E**L**S**K**L** E**D**N**L**K**D**A**E**T**N** N**V**E**D**Y**I**K**E**G**L** E**E**A**I**A**T**K**K**A**E**
361 L**E**K**T**Q**K**E**L**D**A** A**L**N**E**L**G****P**D**G**D E**E**E**T****P**A**P**A**Q**P**E**K**P**A**E**E**P**E**N** **P**A**P**A**P**K**P**E**K**S**A**D**Q**Q**A**E**E**D**Y**A
421 R**R**S**E**E**E**Y**N**R**L**T**Q**Q**Q**P**P**K**A**E**K**P**A**P**A**Q**P**E**Q**P**A**P**A**P**K**I**G**W**K**Q**E**N**G**M**W**Y**F**Y**N**T**D**G**S**M**A**T**G**W**L**Q
481 N**N**G**S**W**Y**Y**L**N**S**N**G**A**M**A**T**G**W**L**Q**Y**N**G**S**W**Y**Y**L**N**A**N**G**A**M**A**T**G**W**L**Q**Y**N**G**S**W**Y**Y**L**N**A**N**G**A**M**A**T**G**W**L**Q**
541 Y**N**G**S**W**Y**Y**L**N**A**N**G**D**M**A**T**G**W**L**Q**Y**N**G**S**W**Y**Y**L**N**A**N**G**D**M**A**T**G**W**A**K**V**H**G**S**W**Y**Y**L**N**A**N**G****S**M**A**T**G**W**V**K 601
D**G**E**T**W**Y**Y**L**E**A** **S**G**S**M**K**A**N**Q**W**F Q**V****S**D**K**W**Y**Y**V**N G**L**G**S**L**S**V**N**T**T** V**D**G**Y**K**V**N**A**N**G** E**W**V

cbpA [Streptococcus pneumoniae GA19998] gram pos signal peptide

1 M**F**A**K****S****S**E**R**K**V** H**Y****S**I**R**K**F****S**I**G** V**A**S**V**V**V**A**S**L**F** L**G**G**V**V**H**A**E**E**V** R**R**G**N**N**L**T**V**T**S** **S**G**D**E**V**E**S**H**Y**Q
61 **S**I**L**E**K**V**R**K**S**L E**K**D**R**H**T**Q**N**V**D** L**I**K**L**Q**D**I**K**R T**Y**L**N**L**K**E**K**P**E**A**E**L**T****S**K**T**K**K** E**L**D**A**A**F**E**K**F**K**
121 **K**E**P**E**L**T**K**K**L**A E**A**E**K**K**A**K**D**Q**K** E**E**D**H**R**N**Y**P**T**N** T**Y**K**T**I**E**L**E**I**A** E**A**E**V**G**V**A**K**A**E** L**E**L**V**Q**A**Q**V**Q**I**
181 **P**Q**D**T**E**K**I**N**A**A K**A**K**V**E**A**A**K****S**N V**K**K**L**E**K**I**K**S**D** I**E**K**T**Y**L**Y**K**L**D** N**S**T**K**E**T****P**K**S**R V**R**R**N****S**P**Q**V**G**D
241 **S**R**E**L**K**E**T**I**D**K A**K**E**T**L**S**T**Y**M**V** T**R**L**T**K**L**D**P**S**V** F**W**F**A**D**L**L**M**D**A** K**K**V**V**E**E**Y**K**T**K** L**E**D**A****S**D**K**K**S**V
301 E**D**L**R**K**E**A**E**G**K** I**E****S**L**I**V**T**H**Q**N R**E**K**E**N**Q**P**A**P**Q** **P**G**G**Q**A**G**G****S**M**V** V**P**P**V**T**Q**T**P**P**S** T**S**Q**S**P**G**Q**K**A**T**
361 E**A**E**K**K**L**Q**D**L I**R**Q**F**Q**E**A**L**N**K** L**D**D**E**T**K**T**V****P**D G**A**K**L**T**G**E**A**G**K** A**Y**N**E**T**R**T**Y**A**K** E**V**V**D**K**S**K**L**L
421 **S**Q**T**A**V**T**M**D**E**L A**M**Q**L**T**K**L**N**D**A** M**S**K**L**K**E**A**K**A**K** L**V****P**E**V****K**P**Q**P**E** N**P**E**P**K**P**Q**P**E**G** E**K**P**S**V**P**D**I**N**Q**
481 E**K**E**K**A**K**L**A**I**A** T**Y**M**S**K**I**L**D**D**I** K**K**H**H**L**K**E**K**H H**Q**I**V**A**L**I**K**D**L** D**K**L**K**K**Q**A**L**S**E** I**D**N**V**N**T**K**V**E**I**
541 E**N**T**V**H**K**V**F**A**D** M**D**T**V**V**T**K**F**Q**K** G**L**I**Q**N**T****P**Q**V**P E**A**P**K**S**P**E**V**P**K** V**S**D**T****P**K**A**P**D**T **P**Q**V****P**E**A**P**K**S**P** 601
E**V**P**K**V**P**E**A**P**K** A**P**D**T****P**Q**V**P**E**A **P**K**S**P**E**V**P**K**V** **D**T**P**K**A**P**D**T**P**Q V**P**E**A**P**K**A**P**D**T** **P**Q**I**P**E**A**P**A**P**E
661 **T**P**A**P**A**P**E**A**P**K**T**G**W**K**Q**E**N**G**M**W**Y**F**Y**N**T**D**G****S**M**A**T**G**W**L**E**Y**N**G**S**W**Y**Y**L**N**A**N**G**A**M**A**T**G**W**L**E**Y**N**G**S**W**
721 Y**L**N**T**N**G**A**M**E**T**G**W**L**E**Y**N**G**S**W**Y**Y**L**N**T**N**G**A**M**E**T**G**W**L**E**Y**N**G**S**W**Y**Y**L**N**T**N**G**A**M**E T**G**W**L**E**Y**N**G**S**W**
781 Y**L**N**T**N**G**A**M**E**T**G**W**L**E**Y**N**G**S**W**Y**Y**L**N**T**N**G**A**M**E**T**G**W**L**E**Y**N**G**S**W**Y**Y**L**N**A**N**G****S**M**A** T**G**W**L**K**D**G**D**T**W**
841 Y**L**E**A**S**G**A**M**K E**S**Q**W**F**K**V**S**D**K** W**Y**Y**V**N**G**S**G**A**L** A**V**N**T**T**V**G**G**Y**R** V**N**A**N**G**K**W**V**N

Pseudomonas aeruginosa

ExoU [Pseudomonas aeruginosa PA103].

1 M**H**I**Q****S**L**G**A**T**A **S**S**L**N**Q**E**P**V**E**T **P**S**Q**A**A**H**K****S**A**S** L**R**Q**E**P**S**G**Q**G**L** G**V**A**L**K**S**T**P**G**I** L**S**G**K**L**P**E**S**V**S**
61 D**V**R**F****S**S**P**Q**G**Q G**E****S**R**T**L**T**D**S**A G**P**R**Q**I**L**R**Q**F E**N**G**V**T**E**L**Q**L**S** **R**P**P**L**T**S**L**V**L**S **G**G**G**A**K**G**A**A**Y**P
121 G**A**M**L**A**L**E**E**K**G** M**L**D**G**I**R**S**M**S**G** **S**S**A**G**G**I**T**A**A**L L**A**S**G**M**S**P**A**A**F** K**T**L**S**D**K**M**D**L**I** **S**L**L**D**S**S**N**K**L**
181 K**L**F**Q**H**I****S**S**E**I G**A**S**L**K**K**G**L**G**L**N K**I**G**G**F**S**E**L**L**L** N**V**L**P**R**I**D**S**R**A** E**P**L**E**R**L**L**R**D**E** T**R**K**A**V**L**G**Q**I**A**
241 **T**H**P**E**V**A**R**Q**P**T V**A**A**I**A**S**R**L**Q**S** **S**G**V**T**F**G**D**L**D** **R**L**S**A**Y**I**P**Q**I**K T**L**N**I**T**G**T**A**M**F** E**G**R**P**Q**L**V**V**F**N**
301 A**S**H**T****P**D**L**E**V**A Q**A**A**H**I**S**S**F**P G**V**F**Q**V**G****S**L**S**D **Q**P**Y**Q**A**G**V**E**W**T E**F**Q**D**G**G**V**M**I**N** V**P**V**P**E**M**I**D**K**N**
361 F**D**S**G**P**L**R**R**N**D** N**L**I**L**E**F**E**G**E**A** G**E**V**A**P**D**R**G**T**R** G**G**A**L**K**G**W**V**V**G** V**P**A**L**Q**A**R**E**M**L** Q**L**E**G**L**E**L**R**E
421 Q**T**V**V**V**P**L**K**S**E** R**G**D**F**S**G**M**L**G**G** T**L**N**F**T**M****P**D**E**I K**A**H**L**Q**E**R**L**Q**E** R**V**G**E**H**L**E**K**R**L** Q**A**S**E**R**H**T**F**A**S** 481
L**D**E**A**L**L**A**L**D**D** **S**M**L**T**S**V**A**Q**Q**N**P**E**I**T**D**G**A**V**A**F R**Q**K**A**R**D**A**F**T**E** L**T**V**A**I**V****S**A**N**G L**A**G**R**L**K**L**D**E**A**
541 M**R**S**A**L**Q**R**L**D**A** L**A**D**T****P**E**R**L**A**W L**A**E**L**N**H**A**D**N V**D**H**Q**Q**L**L**D**A**M** R**G**Q**T**V**Q****S**P**V**L A**A**L**A**E**A**Q**R**R 601
K**V**A**V**I**A**E**N**I**R** K**E**V**I**F**P**S**L**Y**R** **P**G**Q**P**D**S**N**V**A**L L**R**R**A**E**E**Q**L**R**H** A**T****S**P**A**E**I**N**Q**A L**N**D**I**V**D**N**Y****S**A
661 R**G**F**L**R**F**G**K**P**L** **S**S**T**T**V**E**M**A**K**A W**R**N**K**E**F**T

exotoxin A, partial [Pseudomonas aeruginosa].

1 ALLERNYP TG AEFLGDGGDV SFSTRGTQNW TVERLLQAHR QLEERGYV FV GYHGT FLEAA
61 QSIVFGG VRA RSQDLDAIWR GFYIAGDPAL AYGYAQDQ EP DARGRIRIGA LLRVYV PRSS
121 LPGFYRTGLT LAAP EAAGEV ERLIGHPLPL RLDAITG PEE EGGRL ETILG WPLAERTVVI
181 PSAIPTD PRN VGGDL DPSSI PDKEQAISAL PDYASQPGKP PREDLK

toxA gene product [Pseudomonas aeruginosa PAO1]

1 MHLTPH WIPL VASLG LLAGG SFASAAEEAF DLWNECAKAC VLDLKDGVRS SRMSVDP AIA
61 DTNGQGVLHY SMVLEGGNDA LKLAIDNALS ITS DGLTIRL EGGV EPNKPV RYSYTRQARG
121 SWSLNWLVP I GHEKPS NIKV FIHELNAGNQ LSHM SPIYTI EMGD ELLAKL ARDATFFVRA
181 HESNEM QPTL AISHAGV SVV MAQAQPRREK RWSEWASGKV LCLLDPLDGV YNYLAQQRCN
241 LDDTWEGKIY RVLAGNPAKH DLDIKPTVIS HRLHF PEGGS LAALTAHQAC HLPLETFRH
301 RQPRGWEQLE QCGYPVQRLV ALYLAARLSW NQVDQVIRNA LASP GSGGDL GEAIREQPEQ
361 ARLALTLAAA ESERFVRQGT GNDEAGAA SA DVVSLT CPVA AGE CAGPADS GDALLERNYP
421 TGAEFLGDGG DISFSTRGTQ NWTVERLLQA HRQLEERGYV FVG YHGT FLE AAQSIVFGGV 481
RARSQDLDAI WRGFYIAGDP ALAYGYAQDQ EPDARGRIRN GALLRVYVPR SSSLPGFYRTG
541 LTLAAPEAAG EVERLIGHPL PLRLDAITGP EE EGGRL ETI LGWPLAERTV VIPSAIPTDP
601 RNVGGDL DPS SIPDKEQAISAL PDYASQPG KPPREDLK

Escherichia coli

Surface protein [E. coli]

1 MTTPNPLAKT KGAGTTFW MY TGKGD AFANP LSDTDWLRLA MVKDLQPGEM TADAEDDTYL
61 DDEDADWKTT TQGQK SVGDT SATLAWRPGD SGQKKLVQLF DSGEVCAFRI KYPNGTVDVF
121RGWLS SSGKT IASKDVMTRT VKISGVGRPY LAEEGXETVG VTGLTVAPAS ASVKAGATTT
181LTFTVKPDGA SDKAISVHSS DPQTASV TLS GLVATVKGVK QGSVSVIVGMT SDGEFVAVAA
241VTVSAP

Plasmodium falciparum - malaria

PfMSA2

1 SIRR SMAESK SPTGTGASGS AGSGDGASGS AGSGDGASGS AGSGDGAVAS ARNGANP GAD
61 AEGSSSTPAT TTTTTTTTTT TTTNDAEAST STSSNPNHN NAETNPKGNG EVQEPNQANK
121 ETQNN SNVQQ DSQTKSNVPP TQDADTKSPT AQPEQAENSA PTAEQTESPE LQSAPEN

P proline **P** where ginger enzymes digest protein

T threonine that can bind to carbohydrate

S serine that can bind to carbohydrate