GLYCOSYLATED HYDROXYTRYPTOPHAN IN A MUSSEL ADHESIVE PROTEIN FROM *PERNA VIRIDIS*

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The 3, 4-dihydroxyphenyl-L-alanine (Dopa) containing proteins of mussel byssus play a critical role in wet adhesion and have inspired versatile new synthetic strategies for adhesives and coatings. Apparently, however, not all mussel adhesive proteins are beholden to Dopa chemistry. The cDNA-deduced sequence of Pvfp-1, a highly aromatic and redox-active byssal coating protein in the green mussel *Perna viridis* suggests that Dopa may be replaced by a post-translational modification of tryptophan. The N-terminal tryptophan-rich domain of Pvfp-1 contains 42 decapeptide repeats with the consensus sequences ATPKPW1TAW2K and APPPAW1TAW2K. A small collagen domain (18 Gly-X-Y repeats) is also present. Tandem mass spectrometry of isolated tryptic decapeptides has detected both C2-hexosylated tryptophan (W1) and C2-hexosylated hydroxytryptophan (W2) of which the latter is redox active. The UV absorbance spectrum of W2 is consistent with 7-hydroxytryptophan, which represents an intriguing new theme for bioinspired opportunistic wet adhesion.

The amino acid, Dopa (3, 4-dihydroxyphenyl-L-alanine), occurs in many proteins of the mussel holdfast or byssus (1,2) and has recently been incorporated into mussel-inspired synthetic polymers with versatile adhesive consequences (3-7). One byssal protein in particular, mussel foot protein-1 (Mfp-1), has been investigated from over fifteen mussel byssus where it protectively coats compliant collagen-like proteins in the thread core (8-11). Mfp-1s typically contain 10 to 15 mole % Dopa in a highly conserved repeating peptide structure (8). In the blue mussel *Mytilus edulis* fp-1 (Mefp-1), for example, the consensus decapeptide AKPSYPPTYK is repeated over 70 times in tandem, and much of the tyrosine (Y) is converted to Dopa (Y*) (Fig. 1). In stark contrast to this, only trace levels of Dopa were detected in Pvfp-1 from the green mussel *Perna viridis* (Linnaeus 1758) (12), a notoriously invasive fouling species originally from the Indo-Pacific region (13). Since *P. viridis* fp-1 (Pvfp-1) and its homologue, Mefp-1, are both strongly aromatic, quinogenic, and composed of highly polar decapptide repeats (12), identifying the Dopa-mimetic substitutes in Pvfp-1 has been a matter of considerable interest.
Materials and Methods

Protein isolation from mussel feet

Green mussels (Perna viridis) were collected from Tampa harbor in Florida. The feet were severed from about 100 freshly shucked mussels and stored at –80 °C. Protein extraction from mussel feet was adapted from a previous report (12). Frozen mussel feet (in lots of 10 g wet weight) were thawed and depigmented by scraping with a scalpel. The depigmented feet were homogenized in a glass tissue grinder with 50 ml 5% acetic acid and two protease inhibitors (pepstatin A and leupeptin, both 30 mM). The homogenate was centrifuged (15,000 x g, 4 °C, 30 min) and the recovered supernatant (S1) was chilled in an ice bath. 70% perchloric acid (PCA) was added dropwise with stirring to a final PCA concentration of 1.4 v/v %. The mixture was further stirred and centrifuged (15,000g, 4°C) for 30 min. The supernatant (S2) was collected, dialyzed in 500 volumes of 5% acetic acid for 12 hr at 4°C, freeze-dried at –80 °C, and resuspended in 0.2 mL 5% acetic acid.

P. viridis foot protein-1 (Pvfp-1) was purified in 3 stages: gel filtration, reversed phase HPLC, and finally, again, gel filtration. Initial isolation was achieved by gel filtration chromatography on a Shodex KW-803 column (5µm, 8 x 300 mm). The column was equilibrated and eluted with 5% acetic acid at a flow rate of 0.2 ml/min. A maximum volume of 200 µl S2 was loaded onto the Shodex column per run. Eluted volume was monitored at 280 nm, and fractions under the Pvfp-1 peak were pooled (1-1.5 mL) and further resolved by C8 HPLC (Brownlee Aquapore RP-300, 7 µm, 4.6 x 250 mm) at a flow rate of 1 mL/min with an acetonitrile gradient described by reference 12. Collected fractions were assayed by redox cycling after acid urea gel electrophoresis (12) and amino acid analysis as described below to identify Pvfp-1 containing fractions. Pvfp-1 fractions were pooled and lyophilized at –80 °C. After resuspending in 200 µL 5% acetic acid, Pvfp-1 positive fractions were run once more on Shodex KW-803 (above conditions). Fractions eluting between 33 to 43 min were redox-active and had amino acid compositions consistent with previous studies (12). Purified Pvfp-1 fractions were recovered by freeze-drying. The N-terminus of purified Pvfp-1 was sequenced by automated Edman degradation on a Porton Instruments microsequencer (Porton Model 2090, Tarzana, CA) with a modified gradient program (14).

Mass Spectrometry: The mass of intact Pvfp-1 was determined by matrix-assisted laser desorption and ionization with time-of-flight (MALDI-TOF) mass spectrometry (Voyager DE, AB Biosystems, Foster City, CA) in the positive ion mode with delayed extraction. The MALDI matrix was prepared by dissolving sinapinic acid (10 mg/ml) in 30 vol % acetonitrile. Purified Pvfp-1 was dissolved in this matrix solution to give a final concentration between 1 and 10 pmole/µl. About 1 µl of sample was then spotted to the target plate and allowed to dry under vacuum (500 microns). Sample spots were irradiated at 337 nm using an N₂ laser (LSI, Inc., Cambridge, MA) with a pulse width of 8 ns and frequency of 5 Hz. Soft ionization generated singly, doubly, and triply protonated ions of the Pvfp-1 protein variants. The singly and doubly protonated ions of bovine serum albumin were used as molecular mass calibrants at m/z 66,430 and 33,215, respectively).
Cloning and sequencing Pvfp-1 cDNA - To obtain the partial cDNA sequence of Pvfp-1, total RNA was first isolated from the phenol gland at the tip of *P. viridis* foot tissue using an RNase Plant Mini Kit from Qiagen (Valencia, CA). A single freshly dissected foot was pulverized under liquid nitrogen using a mortar and pestle after which the Mini Kit protocol was adopted verbatim. Following that, the first strand cDNA was synthesized using Superscript II reverse transcriptase at 50 °C with an Adapter primer, 5’-GGCCACGCGTCGACTAGTACT (T)16-3’ (Invitrogen). The product of the reverse-transcribed reaction was used as template for subsequent PCRs.

Assuming repetition of the known trypsinized internal peptide sequence APPPAX₁TAX₂K (X denotes unknown amino acid residues) from a previous study (12), a degenerate oligo (sense 5’-AARGCNCCNCCNCCNGC-3’ which corresponds to the sequence KAPPPA) was designed and combined with an Abridged Universal Amplification Primer (antisense 5’-GGCCACGCGTCGACTAGTAC-3’, Invitrogen) to amplify the 3’ end of Pvfp-1 from RT templates. The cloned 3’ cDNA end provided the C-terminal sequence of Pvfp-1 along with some 3’-untranslated region.

Another degenerate oligo (sense 5’-GCNGTNATAYCACNNCCNTC-3’) was designed on the basis of N-terminal sequence of Pvfp-1 (AVYHPPS) and coupled with a gene specific reverse primer (antisense 5’-GGTTTGCCATACCCACCATTACACC-3’ corresponding to GYGGYK) cloned above to amplify the 5’ end cDNA of Pvfp-1.

Once the cDNA sequences encoding the mature Pvfp-1s were cloned, a GeneRacer kit (Invitrogen) was used to obtain the 5’ end untranslated region of the cDNA of Pvfp-1 from full-length transcripts by a 5’ RACE. PCR was performed with gene specific primer (as 5’-GCTTTCCATGCACTCCATGGCAGTGGATG-3’ (corresponding to HPPAWTAWK) coupled with a GeneRacer 5’ sense primer from Invitrogen.

Generally, PCR was carried out in 25 µl of 1x Buffer B (Fisher) and 5 pmole of each primer, 5 µmole of each dNTP, 1 µl first strand reaction, and 2.5 units Taq DNA polymerase (Fisher) for 35 cycles on a Robocycler (Stratagene). Each cycle consisted of 30 s at 94 °C, 30 s at 50 °C, and 1 min at 72 °C, with a final extension of 15 min. The PCR products were subjected to 1% agarose gel electrophoresis, purified and cloned into a PCR TA vector (TOPO TA Cloning Kit, Invitrogen) and transformed into competent “Top10” cells (Invitrogen) for amplification, purification and sequencing.

Byssal thread analysis - Fifteen to twenty threads were severed from the proximal end of each byssal stem, washed in 500 vols milli-Q water and blotted dry. Dry threads were weighed and hydrolyzed by one of two methods: 6 M HCl for all amino acids but Trp or 4 M NaOH for Trp (15). Acid hydrolysates were done with reusable hydrolysis vials (Pierce Chemical) at 110˚ C for 24 h and flash evaporated under high vacuum, whereas NaOH hydrolysates were titrated to pH 3 - 4 with glacial acetic acid. Purification of Trp-derived amino acids was achieved by C-18 HPLC (Brownlee Aquapore OD-300, 7 µm, 4.6 x 250 mm) over an acetonitrile gradient of 0-10%. The eluate was monitored continuously at 220 and 280 nm, and all those peaks absorbing at both wavelengths were collected for analysis by electrospray
ionization (ESI) and tandem mass spectrometry.

For routine amino acid analysis, the purified Pvfp-1s were hydrolyzed in one of three ways: 1) 4 M methanesulfonic acid with 0.5% 3-methyl-indole (16), 2) 6 M HCl with 5% phenol in vacuo at 110 °C for 24 hours or 3) 4 M NaOH in vacuo at 110 °C for 24 hours (15). The first method was dropped due to extremely low tryptophan derivative recovery. The HCl hydrolysate was flash evaporated at 50°C under vacuum and shaken to dryness with 0.5 mL of MilliQ-water followed by methanol, NaOH hydrolysates were titrated to pH 4. Amino acid analysis was performed according to conditions described earlier with a Beckman System 6300 Auto Analyzer (14) on which an authentic C²-(α-D-mannopyranosyl)-tryptophan (C-ManTrp) standard (17) eluted just after Met with a run time of 31.5 min.

Trp-containing peptides—Preparation of tryptic peptides from Pvfp-1, including purification by C18 HPLC (Brownlee OD 300, 4.6 x 260 mm) was done exactly as described by Ohkawa et al (12). Proteolysis was stopped by acidification to pH 4 with glacial acetic acid before HPLC. Three peak fractions corresponding to Ohkawa’s peptides d, e and j were selected for analysis by electrospray ionization (ESI) mass spectrometry followed by tandem mass spectrometry with collision induced decomposition (CID) using the PE Sciex QStar quadrupole/time-of-flight tandem mass spectrometer (Perkin-Elmer, CA) in the UCSB Mass Spectrometry Facility. Interpretation of fragments produced from the peptides by CID was assisted by comparison to an authentic C²-Man-Trp standard and by mock fragmentation of various model sequences using Protein Prospector, version 5.1.4 on ExPASy Tools.

UV-Vis spectra of indole derivatives. Chemical modifications based on 1-nitroso-2-naphthylation (18) and nitration (19) were performed as described. The former reacts only with the 5-hydroxy indole isomer, and the latter, which was developed for catechol detection, gave a bright yellow color for Pvfp-1 and its tryptic peptides. UV-visible spectra were obtained for 0.1 mM solutions of L-tryptophan (Sigma), C²-mannosyltryptophan (donated by S. Manabe), peptide e, and four hydroxyindoles [4-hydroxyindole (Acros Organics), 5-hydroxytryptophan (Sigma), 6-hydroxyindole (Oakwood Products, S. Columbia, SC), and 7-hydroxytryptophan (SynChem OHG, Felsberg, Germany)] using an HP model 8453 UV-Vis scanning spectrophotometer with UV Vis ChemStation (Rev 06.03). All solutions were buffered with 5% (v/v) acetic acid and scanned between 235 and 335 nm in masked 40 µL quartz cells (Starna, Atascadero, CA).

RESULTS

A previous investigation by Ohkawa et al. (12) reported the following attributes for Pvfp-1: 1) an apparent mass of 89 kDa based on SDS PAGE, 2) marked quinone-like redox cycling activity without Dopa, 3) significant carbohydrate content, particularly with respect to mannose, N-acetylglucosamine and fucose, and 4) an Edman derived primary sequence dominated by two closely related consensus repeats: AOOOX,TAX,K and APOKOX,TAX₂,K, in which O denotes trans-4-hydroxyproline. The X₁ and/or X₂ positions were consistent with the presence of an aromatic amino acid but
could not be reconciled with any known modification of Tyr or Dopa.

In the present investigation, purified Pvfp-1 was shown to consist of at least two variants (Fig. S1) with masses of 50 and 64 kDa observed by MALDI TOF mass spectrometry (Fig S2). The N-terminal sequence of the isolated protein was found to be (A)VY*HPPSX,T A,X1AOK (Fig. S1) where X1 and X2 represent Edman cycles devoid of detectable phenylthio hydantoin (PTH) derivatives. Y*, denotes Dopa and was the only Dopa detected in Pvfp-1.

To further explore the chemistry of X1 and X2, we first deduced the complete sequence of Pvfp-1 from its corresponding cDNA prepared by standard cloning procedures. Two Pvfp-1 variants, probably related by alternative splicing, were found with the shorter differing from the longer by exactly fifteen decapeptide repeats. In the longer variant 1 (calculated mass 61 kDa), [APPPAWTAWK] and [ATPKPWTAWK] consensus repeats occur 14 and 29 times, respectively (Fig. 2 and Fig. S3). Other intriguing features in Pvfp-1 are a distinct collagen-like sequence with eighteen tripeptide [Gly-X-Y] repeats, followed by a short interval (14 amino acids long), before returning to a W-rich C-terminal sequence. It must therefore be concluded that in Pvfp-1 some modification of tryptophan (W), not tyrosine, provides the basis for the unknown aromatic amino acids in the decapeptide repeats.

Since tryptophan is unstable to hydrolysis by HCl, purified Pvfp-1 and P. viridis byssal threads were hydrolyzed with 4M NaOH. After hydrolysis, component aromatic amino acids were chromatographically separated by C-18 HPLC (Fig. 3) and analyzed by mass spectrometry to help deduce their relationship to tryptophan. Two ions with masses of 367 and 529 Da eluting between 22-24 min were detected; the latter decomposed readily to 367 through a loss of 162 Da (hexose) (Fig. S4) suggesting a cleavage that is typical of O- and N-linked hexoses (20, 21). The 367 Da ion and its fragmentation, most notably the 120 Da loss to m/z 247, are identical with standard C2-[α-D-mannopyranosyl]-tryptophan (22, 23) (Fig. 4). The hexose linked to Trp in Pvfp-1 is likely to be mannose, but tandem MS by itself is unable to distinguish different hexoses. Since C2-ManTrp elutes just after Met in amino acid analysis, this method was used to estimate a C2-mannosylTrp content of 1-2 dry weight% in byssal threads and 3-4 dry weight % in Pvfp-1 following base hydrolysis (Table S1). Only trace levels of Trp could be detected in Pvfp-1, thus the Trp in byssus must come from other proteins.

In order to detect modified tryptophans more directly in the decapeptide sequences of Pvfp-1, we used tandem mass spectrometry with collision-induced decomposition of three peptides following trypsin digestion of Pvfp-1 (Fig. S5). Peptides d and e were selected since 75% of their sequence had already been established by Edman chemistry (12). The doubly charged parent ion (M+2H)2+ (m/z 953.8) for the tryptic decapeptide d readily decomposes to another ion (M+2H)2+ (m/z 771.25) after a neutral loss of 162 to m/z 873 (hexose) followed by another neutral loss of 204 to m/z 771 (N-acetyl hexose) (Fig. 5A). The oxonium ion of hexosyl-N-acetylhexose is evident at m/z 366. The 771.25 peak loses water to become m/z 762, which then undergoes the 120 Da (60 x 2) neutral loss typical of C2-
mannosylation; $y''_6$ or $OW_1TA\text{W}_2K (m/z 1144)$ does the same. The internal fragment, $OW_1TA (m/z 616.1)$, indicates a 366 Da mass for $X_1$ (Fig. 5B). Notably, the 511.1 ($y''_2$) and 290 ($y''_3^{2+}$) ions corresponding to $W_2K$ and $AW_2K$ are larger than the comparable $W_1$-containing fragments by 16 Da. C$^2$-ManTrp + 16 is consistent with C$^2$-hexosylated hydroxytryptophan ($W_2K$) and the quinoid counterpart of C$^2$-hexosylated hydroxytryptophan ($AW_2K$) (Fig. 5B).

A complete annotation of fragment masses is given in Fig. S6.

Mannose is plausible as the C$^2$-hexose given that $W-x_1-x_2-W$ is a well-accepted sequence signature for C$^2$-mannosylation of tryptophan in proteins (20, 24-26). The precise ring assignment of the hydroxyl group in hydroxytryptophan (OHTrp) is not possible from ESI ms/ms, but a phenyl ring placement is consistent with the loss of two hydrogens to form a doubly protonated quinoid ion at $m/z$ 290 and by the strong quinone-like redox cycling activity in Pvfp-1 and tryptic decapetides (12).

Peptide e ($m/z$ 757) is less glycosylated than d but shows some trends similar to those in the 771 ion during fragmentation (Fig. 6). The $y''_8$ ion (OPAW$_1TA\text{W}_2K$) at $m/z$ 1328 shows a neutral loss of 120 Da. The fragment ions implicate both $W_1$ and $W_2$ as C$^2$-ManOHTrp: the $y''_1$ ($W_1$K) and $y''_4$ ions ($\text{TAW}_2K$) are particularly suggestive of $W_2$ (Fig. 6; complete fragment annotation in Fig. S7).

The fragmentation of peptide j (Fig. 7 and Fig. S8) was undertaken to corroborate the odd collagen-like primary structure predicted by the cDNA-deduced Pvfp-1 sequence (Fig. 2). Peptide j is 14 residues long and its sequence corresponds to residues #496-509 with Gly at every third residue and three of the five prolines converted to trans-4-hydroxyproline.

Neither hydroxytryptophan nor its C-hexosylated forms were detected after MSA or NaOH hydrolysis of the byssus or Pvfp-1. We observed that whereas standard 5- or 7-hydroxytryptophan were stable to hydrolysis in MSA, standard C$^2$-ManTrp was not. OHTrp standards hydrolyzed in the presence of sugars were not detected by amino acid composition. C$^2$-ManOHTrp would thus be unlikely to survive MSA. OHTrp recovery following alkaline hydrolysis is prevented by the formation of quinonimine at alkaline pH (27).

Tandem mass spectrometry supports OHTrp presence but is unable to specify the position of the hydroxy group. To further clarify this important point, Pvfp-1 and peptide e were reacted with nitrosonaphthol, which is specific for the 5-hydroxy position on the indole (18); both were negative. In addition, the UV absorbance spectrum of peptide e at pH 3.5 was compared to a series of related indole derivatives (28-29). In contrast to L-tryptophan and C$^2$ Man-Trp, peptide e has a $\lambda_{max}$ 269 nm shifted to lower wavelength (Fig. 8 and Fig. S9), and an extinction coefficient of 18000 M$^{-1}$ cm$^{-1}$ at 269 nm that is consistent with the presence of two modified tryptophans enhanced by mannosylation (28). Of the four hydroxy-indole derivatives tested, only 7-hydroxytryptophan shared the same $\lambda_{max}$ values as peptide e. Indeed, the two spectra are nearly superimposable between 250-300 nm. Although future studies need to further characterize the tryptophan chemistry of peptide e by proton NMR and electrochemistry, the identification $W_2$ as C$^2$ hexosyl-7-hydroxy tryptophan in Pvfp-1 appears reasonable. A final test of hydroxy-indole
reactivity with Arnow’s reagent showed that only 7-hydroxy-tryptophan produced the same bright yellow product formed by Pvf-1 (Fig. S9).

**DISCUSSION**

Pvf-1 resembles other mussel coating proteins in its canonical, Pro- and Lys-rich repeats. It diverges significantly, however, in its reliance on modifications of tryptophan rather than tyrosine for its redox chemistry. Pf-1 consists of two closely related variants, the larger of which has predicted and observed masses of 61 kDa and 64 kDa, respectively. Both variants show a highly repetitive sequence with four distinct domains: a Trp-rich decapeptide repeat (consensus: APPPAWTAWK and ATPKPWTAWK) domain largely in the N-terminal two-thirds, a collagen domain (Gly-X-Y repeats), a short hinge (Res #450-459) and a predicted, short coiled-coil region before returning to W-rich repeats at the C-terminus (Fig. 9). The collagen domain is reminiscent of other collagen-like proteins such as the macrophage scavenger receptor (30), *Torpedo* acetylcholinesterase (31), and complement C1q (32) in which a small collagen domain directs trimer formation. The proposed trimeric structure remains speculative for Pf-1 as only the single chain mass has been determined with any accuracy by MALDI TOF.

Pvf-1 exhibits extensive post-translational decoration. Pro is targeted for hydroxylation throughout both decapeptide consensus repeats and in the collagen domains; Thr-2 in ATPKPWTAWK appears to be O-glycosylated with O-(N-acetyl)-hexosyl-hexose, which agrees with the high levels of N-acetyl-glucosamine reported earlier (12) but differs from the previous designation of Pro for position #2 in peptide e. Possibly, the PTH derivative of glycosylated Thr has the same elution time as PTH-Pro on C18 HPLC. Notably, Trp undergoes C-hexosylation (probably by mannose) and hydroxylation. Whereas W1 and W2 are always hexosylated, W2 seems favored for hydroxylation. Peptides d and e may be representative of other decapeptides in Pf-1 with respect to C-hexosylation and hydroxylation. At this stage, however, aside from ample evidence for 120 Da mass losses i.e. C-hexosyl-Trp in other peptides, collision-induced decomposition of many peptides remains too complex for confident interpretation. Pf-1 has only a single Dopa located near the N-terminus.

C-linked mannosylation of proteins is an unusual but not unprecedented modification of tryptophan. It was first reported in pancreatic ribonuclease and since then in over 47 other proteins including notably thrombospondin and complement proteins with the sequence motif W-x1-x2-W (20, 22-26). Although the function of this modification remains elusive, mannosylation is known to render the tryptophan more polar and solvent accessible (23). In complement proteins, C-ManTrp in the thrombospondin-like repeat domains has been proposed to have an adhesive function (33). The potential presence of up to eighty C-ManTrp residues in Pf-1, which functions as an adhesive and coating in byssus (9-11), begs the hypothesis that mannosylation makes tryptophan behave more like Dopa – sticky and prone to cross-link formation. This conjecture, however, is too simple since tryptophan needs to be hydroxylated before becoming redox active like Dopa. 5-Hydroxytryptophan in engineered proteins and as a free amino acid was
reported to undergo oxidation resulting in the formation of p-quinonimine and di-Trp cross-links (27, 34). Indeed, based on these studies, we expected to find C2 mannosylated 5-HOTrp in peptide e and Pvfp-1. UV spectra, however, support the rarer 7-OH rather than the 5-OH isomer and represent the first report of naturally occurring 7-hydroxytryptophan in proteins (35). 7-OHTrp seems a better mimic of Dopa than the other isomers. Upon oxidation, it forms an o-quinonimine (Fig. 10A), and the indolic N-H and phenolic OH groups are close enough to chelate metal ions (Fig. 10A), a common capability of mussel byssus (9, 36).

If 7-hydroxylation makes Trp more like Dopa, why bother with mannosylation? Trp mannosylation was observed to block cleavage in the vicinity of modified residues by exo- and endoproteases, hence renders proteins more resistant to degradation (28). In addition, Trp-rich proteins are quite hydrophobic thus prone to aggregation in solution (37). C-mannosylation of Trp residues appears to make them fully water accessible while preventing aggregation (38).

In summary, the green mussel *Perna viridis*, employs an intriguing alternative to Dopa in Pvfp-1, one of its byssal adhesive proteins. Although much bulkier, C-Man-7-OHTrp resembles Dopa in having attributes that contribute to both cohesive and adsorptive interactions necessary for adhesion (39). With respect to cohesion, the 2-electron oxidation of C-Man-7-OHTrp to an o-quinonimine (Fig. 10A) mimics Dopa-o-quinone, which is known to form covalent cross-links with other Dopa, cysteine, lysine, and histidine residues (Fig. 10B) (7, 40). The coordination of metal ions via the o-hydroxyl groups in Dopa also contributes to cohesion, particularly in the byssal cuticle of *M. galloprovincialis* (9). The ortho-placement of indolic-N and phenolic OH groups in C-Man-7-OHTrp seems a natural chelate metal ion binding but has yet to be investigated.

The stickiness of adhesive molecules is determined by their adsorptive tendencies. Dopa complexation of metal hydroxides on surfaces provides strong and reversible interactions (Fig. 10B) (41). Again, given the ortho-placement of electronegative elements such interactions also seem likely with C-Man-7-OHTrp. Future studies need to closely examine what adaptive advantages C-Man-7-OHTrp offers over Dopa.

**REFERENCES**

FOOTNOTES

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1* contributed equally.

The abbreviations used are: C2ManTrp, C2-[α-D-mannopyranosyl]-tryptophan; C2ManOHTrp, C2-[α-D-mannopyranosyl]-hydroxytryptophan; CID, collision induced decomposition; ESI, electrospray ionization; fp-1, foot protein 1; MALDI, matrix assisted laser desorption ionization; Mefp-1, Mytilus edulis foot protein 1; MSA, methansulfonic acid; OHTrp, hydroxytryptophan; PCR, ; PTH, phenylthiohydantoin; RACE, rapid amplification to cDNA ends; RT, reverse transcriptase; SDS PAGE, sodium dodecylsulfate polyacrylamide gel electrophoresis; TANDEM MS or MS MS, tandem mass spectrometry
FIGURE LEGENDS

Fig. 1. The common blue mussel *Mytilus edulis* and green mussel *Perna viridis* are shown with attached byssal threads. Distinct consensus decapeptide repeat sequences are associated with the repeat domains in fp-1 foot proteins of the 2 species. The amino acid Dopa (Y*, right) is prominent in mefp-1 repeats but absent from Pvfp-1. O denotes trans-4-hydroxyprolines. Residues denoted as X₁ and X₂ are shown by this study to be derived from tryptophan.

Fig. 2. Complete protein sequence of Pvfp-1 variant 1 deduced from cDNA. Signal peptide is *italicized* and the mature N-terminus indicated by *arrows*. Three directly sequenced peptide sequences are underlined: N-terminus (solid), decapeptide repeats (peptides e and d, dashed) and collagen (peptide j dotted). The single dopa (Y) residue is circled. Variant 2 sequence is in the supporting data. GenBank accession numbers for variants 1 and 2 are AAY46226.1 and AAY46227.1.

Fig. 3. C-18 HPLC of C²-hexosyl Trp from NaOH hydrolyzed *P. viridis* byssal threads (*top*); standard L-Trp (*middle*); standard C²-mannosylTrp (*bottom*). Fractions at 22-25 min were sampled by ESI mass spectrometry and collision induced decomposition. Standard 7-hydroxytryptophan, which does not survive NaOH hydrolysis, eluted at 17 min.

Fig. 4. Mass spectrometric analysis of the m/z 367 peak obtained from C18 HPLC of NaOH hydrolyzed *P. viridis* threads (above) and purified Pvfp-1 (below). The loss of 120 Da is a signature for C-hexosylation.

Fig. 5. Tryptic peptide d (parent ions [M+2H]²⁺ m/z 953 and 771) following collision induced decomposition. O denotes trans-4-hydroxyproline; W₁ C²-hexosylTrp and W₂ C²-hexosylhydroxyTrp. See supporting data for complete fragment annotation.

Fig. 6. Tryptic peptide e (parent ion [M+2H]²⁺ m/z 757) following collision induced decomposition. O denotes trans-4-hydroxyproline; W₁ and W₂ both are consistent with C²-hexosylhydroxyTrp. See supporting data for complete fragment annotation.

Fig. 7. Tryptic peptide j (parent ion [M+2H]²⁺ m/z 692) after collision induced decomposition. O denotes trans-4-hydroxyproline. See supporting data for complete fragment annotation.

Fig. 8. Ultraviolet absorbance spectra of Pvfp-1 derived peptide e and model hydroxyindoles in 5% acetic acid with extinction coefficients at selected wavelengths. A. 4-hydroxyindole, B. 5-hydroxytryptophan, C. 6-hydroxyindole, D. 7-hydroxytryptophan. The spectrum of peptide e (broken line) is superimposed with all other spectra.

Fig. 9. Model of trimeric Pvfp-1 based on the trimerization imposed by formation of the collagen domain. Structure of the long N-terminal and short C-terminal repeat domains is not predictable at present. The coiled coil region was predicted by Coils (window 14) in ExPAsy Tools.

Fig. 10. C²-hexosyl-7-OHTrp reactivity (A) has parallels with peptidyl-Dopa (B). In the redox pathway, C²-hexosyl-OHTrp loses 2 hydrogens to become an o-quinonimine (*bottom*); Dopa loses 2 hydrogens (2H⁺ + 2e⁻) to become an o-quinone that reacts with cysteine, histidine, lysine and other Dopa residues to form cross-links (B). In the metal chelate pathway, both the indolic nitrogen and phenolic oxygen of C²-hexosyl-7-OHTrp should contribute electrons for a bidentate complexation (A, *top right*); Dopa forms stable bidentate complexes with metal ions and metal hydroxides (Mⁿ⁺[OH]ₙ⁻)(B, *center*).
The preferred metal (M) bound to Dopa in *Mytilus* byssus is Fe$^{III}$; in Pvfp-1 of *Perna viridis* byssus it is unknown. Covalent cross-links and bis-bidentate metal complexes contribute to adhesive cohesion whereas mono-bidentate metal hydroxide complexes contribute to adhesive adsorption.
Figure 1. The common blue mussel *Mytilus edulis* and green mussel *Perna viridis* are shown with attached byssal threads. Distinct consensus decapeptide repeat sequences are associated with the repeat domains in fp-1 foot proteins of the 2 species. The amino acid Dopa (Y*, right) is prominent in mefp-1 repeats but absent from Pvfp-1 decapeptide repeats. O denotes trans-4-hydroxyprolines. Residues denoted as $X_1$ and $X_2$ are shown by this study to be derived from tryptophan.
Figure 2. Complete protein sequence of Pvfp-1 variant 1 deduced from cDNA. Signal peptide is italicized and the mature N-terminus indicated by arrows. Three directly sequenced peptide sequences are underlined: N-terminus (solid), decapeptide repeats (peptides e and d, dashed) and collagen (peptide j dotted). The single dopa (Y) residue is circled. Variant 2 sequence is in the supporting data. GenBank accession numbers for variants 1 and 2 are AAY46226.1 and AAY46227.1.
Figure 3. C-18 HPLC of C²-hexosyl Trp from NaOH hydrolyzed *P. viridis* byssal threads (*top*); standard L-Trp (*middle*); standard C²-mannosylTrp (*bottom*). Fractions at 22-25 min were sampled by ESI mass spectrometry and collision induced decomposition. Standard 7-hydroxytryptophan, which does not survive NaOH hydrolysis, eluted at 17 min.
Figure 4. Mass spectrometric analysis of the $m/z$ 367 peak obtained from C18 HPLC of NaOH hydrolyzed *P. viridis* threads (above) and purified Pvfp-1 (below). The loss of 120 Da is a signature for C-hexosylation.
Figure 5. Tryptic peptide $d$ (parent ions [M+2H]$^{2+}$ m/z 953 and 771) following collision induced decomposition. O denotes trans-4-hydroxyproline; W$_1$ C$^2$-hexosylTrp and W$_2$ C$^2$-hexosylhydroxyTrp. See supporting data for complete fragment annotation.
Figure 6. Tryptic peptide e (parent ion [M+2H]^{2+} m/z 757) following collision induced decomposition. O denotes trans-4-hydroxyproline; W₁ and W₂ both are consistent with C²-hexosylhydroxyTrp. See supporting data for complete fragment annotation.
Figure 7. Tryptic peptide \( j \) (parent ion \([M+2H]^{2+}\) \( m/z \) 692) after collision induced decomposition. O denotes trans-4-hydroxyproline. See supporting data for complete fragment annotation.
Figure 8. Ultraviolet absorbance spectra of Pvfp-1 derived peptide e and model hydroxyindoles in 5% acetic acid with extinction coefficients at selected wavelengths. A. 4-hydroxyindole, B. 5-hydroxytryptophan, C. 6-hydroxyindole, D. 7-hydroxytryptphan. The spectrum of peptide e (broken line) is superimposed with all other spectra.
Figure 9. Model of trimeric Pvfp-1 based on the trimerization imposed by formation of the collagen domain. Structure of the long N-terminal and short C-terminal repeat domains is not predictable at present. The coiled coil region was predicted by Coils (window 14) in ExPAsy Tools.
Figure 10. C$_2$-hexosyl-7-OHTrp reactivity (A) has parallels with peptidyl-Dopa (B). In the redox pathway, C$_2$-hexosyl-OHTrp loses 2 hydrogens to become an o-quinonimine (*bottom*); Dopa loses 2 hydrogens (2H$^+$ + 2e$^-$) to become an o-quinone that reacts with cysteine, histidine, lysine and other Dopa residues to form cross-links (B). In the metal chelate pathway, both the indolic nitrogen and phenolic oxygen of C$_2$-hexosyl-7-OHTrp should contribute electrons for a bidentate complexation (*top right*); Dopa forms stable bidentate complexes with metal ions and metal hydroxides (M$^{n+}$[OH]$_n$). The preferred metal (M) bound to Dopa in *Mytilus* byssus is Fe$^{III}$; in Pvfp-1 of *Perna viridis* byssus it is unknown. Covalent cross-links and bis-bidentate metal complexes contribute to adhesive cohesion whereas mono-bidentate metal hydroxide complexes contribute to adhesive adsorption.
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