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Peptides derived from HIV-1, HIV-2, Ebola virus, SARS coronavirus and coronavirus 229E exhibit high affinity binding to the formyl peptide receptor

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Abstract

Peptides derived from the membrane proximal region of fusion proteins of human immunodeficiency viruses 1 and 2, Coronavirus 229 E, severe acute respiratory syndrome coronavirus and Ebola virus were all potent antagonists of the formyl peptide receptor expressed in Chinese hamster ovary cells. Binding of viral peptides was affected by the naturally occurring polymorphisms at residues 190 and 192, which are located at second extracellular loop–transmembrane helix 5 interface. Substitution of R190 with W190 enhanced the affinity for a severe acute respiratory syndrome coronavirus peptide 6 fold but reduced the affinity for N-formyl-Nle–Leu–Phe by 2.5 fold. A 12 mer peptide derived from coronavirus 229E (ETYIKPWWVWL) was the most potent antagonist of the formyl peptide receptor W190 with a $K_i$ of 230 nM. Fluorescently labeled ETYIKPWWVWL was effectively internalized by all three variants with EC50 of $\sim$25 nM. An HKU-1 coronavirus peptide, MYVKWPWYWVL, was a potent antagonist but N-formyl-MYVKWPWYWVL was a potent agonist. ETYIKPWWVWL did not stimulate GTPγS binding but inhibited the stimulation by formyl-NleLeuPhe. It also blocked β arrestin translocation and receptor downregulation induced by formyl-NleLeuPhe. This indicates that formyl peptide receptor may be important in viral infections and that variations in its sequence among individuals may affect their likelihood of viral and bacterial infections.

Keywords: Virus; Signal Transduction; G protein coupled receptor; Polymorphism

1. Introduction

Neutrophils play an essential role in innate immunity. In addition to activating phagocytosis and secreting superoxides and hydrolytic enzymes, the neutrophil must be able to chemotax, or migrate toward the source of a chemoattractant. Upon activation by a chemoattractant, G-protein coupled receptors on the membrane of the neutrophil change conformation and bind to G proteins, activating a signal transduction pathway that results in intracellular actin polymerization, the formation of pseudopodia, reorientation toward a chemoattractant gradient, and adhesion to and infiltration through the blood vessel endothelium [1,2].

The formyl peptide receptor (FPR) is a chemoattractant G protein-coupled receptor found on the surface of phagocytes. It is thought to play an important role in allowing phagocytic cells to recognize the presence of bacteria, which are believed to be the source of formyl peptides [3]. FPR is activated by other ligands in addition to fMLF. FPR binds peptides derived from the GP-41 envelope protein of HIV-1 [4,5]. FPR activation is inhibited by a polypeptide secreted by S. aureus [6,7], an undecapeptide (Cyclosporin H) derived from the fungus Tolypocladium inflatum [8], and peptides derived from murine leukemia virus and human T cell leukemia virus [9]. A peptide derived from herpes simplex
virus type 2 elicited chemotaxis and superoxide production in neutrophils in a process that appeared to involve FPR [10]. Thus FPR appears to interact with peptides derived from a variety of sources.

Lentiviruses are associated with immunological impairment in their respective hosts and both human immunodeficiency and feline immunodeficiency viral infections increase the likelihood of secondary bacterial infections [11]. Recently, Kubes et al. [12] demonstrated that feline neutrophils exhibited a marked (>90%) reduction of neutrophil chemotaxis toward fMLF following infection with FIV. It is possible that viral infections may release factors which impair chemotaxis in vivo, which would be consistent with the observations of virally derived peptides on FPR.

Polymorphisms in the FPR appear to be very common. Sahagun-Ruiz et al. [13] did an extensive haplotype investigation of FPR and reported finding at least 23 haplotypes for FPR. No polymorphisms were found in the closely related receptor FPRL1 [13]. One possible reason for the existence of polymorphisms is that polymorphisms might prevent pathogens from binding to receptors and using them as an entry point into cells. Several G protein-coupled receptors have been used by pathogens as a point of entry into cells and the use of CCR5 and CXCR4 as co-receptors for HIV-1 and CXCR4 as a co-receptor for FIV are notable examples [14–16]. The CCR5 receptor in African green monkeys [17] has been shown to be polymorphic and these polymorphisms provide some protection against SIV infection.

Numerous studies have indicated that patients with aggressive periodontitis exhibit a ~2 fold reduction in chemotaxis toward fMLF [18–23]. Zhang et al. [24] found that aggressive periodontitis patients of African American decent were more likely (P = 0.002) to have at least one allele as the R190/K192 polymorphism when compared to African American controls (R/W190 and N/K192 are located at the extracellular loop 2–TMH5 interface). They also found that aggressive periodontitis patients of African American decent were much less likely (P = 0.003) to have one of their alleles as the W190/N192 polymorphism. R190/K192 was not associated with an increased risk of aggressive periodontitis among either Brazilians or Turks nor was W190/N192 associated with reduced incidence of aggressive periodontal disease in these two populations [24]. It is unclear at this point why FPR W190/R190 and N192/K192 are only associated with altered risk of aggressive periodontitis in African Americans but the effects of these polymorphisms on receptor function clearly needs to be determined.

We previously observed that a single amino acid mutation, D106N in TMH3, markedly enhanced the ability of FPR to bind a peptide derived from HIV-1 gp41 but also markedly reduced its affinity for fMLF [25]. Thus, determination of the affinity of these two very different ligands provides a way to observe changes in ligand specificity associated with changes in FPR. In order to determine whether polymorphisms W190 or K192 might similarly affect ligand binding and possibly explain why K192 is associated with an increased risk of periodontal disease and W190 is associated with reduced risk of disease, we expressed R190/N192, W190/N192, and R190/N192 FPR variants in CHO S cells and determined their affinity for numerous ligands, including several formyl peptides and peptides derived from HIV-1 gp-41 and analogous peptides derived from HIV-2, SARS coronavirus, 229E, coronavirus OC43, and Ebola virus.

2. Materials and methods

2.1. Cell transfection

Geneticin- and ampicillin-resistant and FPR-expressing DNA plasmid PBGSA was prepared as previously described [26]. Mutagenesis was performed according to the Quickchange Mutagenesis Kit instructions using pfu DNA polymerase essentially as described previously [27]. The plasmids were cut with BspEI and Xbol and ligated back into the original FPR-expressing DNA plasmid PBGSA. This step was necessary to obtain good expression levels. DNA was purified, sequenced (Davis Sequencing) and transfected into CHO S cells (Invitrogen) using Lipofectamine 2000, cloned, and maintained in Gibco CHO-S-SFMII media containing 100 μg/ml G418. Numerous clones expressing R190/N192, R190/ K192, or W190/N192 were evaluated for their expression by FACScan analysis using 10 nM formyl-Nle–Leu–Phe–Nle–Tyr–Lys–FITC. A clone was chosen for each sequence that exhibited nearly identical levels of surface expression, less than ~20% difference between the three variants.

2.2. Peptide synthesis

Peptides were obtained from either Genscript or Macromolecular Resources and purified by reverse phase HPLC. Peptides were synthesized with unmodified N and C terminals. Mass spectral data was provided with each peptide and the major mass peak matched the expected mass of the peptide in all cases. Formyl-Nle–Leu–Phe was obtained from Sigma and used without further purification.

2.3. Fluorescent labeling of peptides and anti β-arrestin antibody

Peptides were reacted with Alexa Fluor 488 n-hydroxysuccinimidelester (Molecular Probes) at a 1.5 to 1 mole ratio (peptide to fluorophore) in DMSO/1% triethylamine at 30 °C for ~4 h and purified by reversed phase HPLC. For formyl-Nle–Leu–Phe–Nle–Tyr–Lys the only site of labeling is the Lys. For ETYIKWPWWVL either the Lys could be labeled or the N-terminus could be labeled. However, the intrinsic reactivity of the ε amino of Lys is generally ~100 fold greater than the N-terminal amino group so it is likely that almost all labeling occurs at the lys when the peptide is used in excess to the fluorophore. Anti β-arrestin antibody (BD Transduction Laboratories) at 250 μg/ml in PBS was incubated with 5 mM 6-(fluorescein-5-carboxamido)hexanoic acid, succinimidyl ester (Molecular Probes) for 1 h at 37 °C and the labeled antibody was purified by HPLC gel filtration on a Sigmachrome 100 column. The antibody contained approximately 1.5 moles of fluorescein/mole of antibody.

2.4. Determination of GTPγS binding

Membranes from CHO S cells expressing FPR were prepared as follows: The cells were centrifuged at 100 g for 5 min. The cell pellets were resuspended in 10 mM phosphate, pH 7.4, 10 mM KCl, 100 mM NaCl, 0.1 μM GDP and 5 μg/ml leupeptin. The cells were sonicated for 30 s at 20% power and stored as aliquots of approximately 1.5 moles of fluorescein/mole of antibody.

2.5. FACScan analysis of ligand binding to FPR variants expressed in CHO S cells

CHO S cells (100 μl in Gibco CHO-S-SFMII) were added to 3 mM KCl, 10 mM NaCl, 10 mM sodium phosphate, pH 7.4, 1 mM Mg++, 1 mM Ca++.
containing 5% fetal bovine serum and incubated at 4 °C for 60 min with varying concentrations of fluorescent peptide. The mean fluorescence of the cells was determined, and the mean fluorescence observed in non-expressing CHO S cells (non-specific binding) was subtracted. The data were then analyzed by nonlinear least squares analysis and the Kᵢ and Bₘₐₓ determined. The affinity of peptides was determined using increasing concentrations of peptide in the presence of constant formyl-Nle–Leu–Phe–Nle–Tyr–Lys–FITC. The Kᵢ was then determined by nonlinear least squares analysis using the known Kᵢ for formyl-Nle–Leu–Phe–Nle–Tyr–Lys–FITC for each FPR variant and the observed EC₅₀ of each peptide. Formyl-Nle–Leu–Phe–Nle–Tyr–Lys–FITC concentrations of 0.3 nM, were used for all three FPR variants, and the Kᵢ was determined from the equation Kᵢ = EC₅₀/(1 + (ligand/Kᵢ)) using the graphics/statistics program Graphpad Prizm.

2.6. Translocation of arrestin

CHO S cells were incubated in Gibco CHO-S-SFMII at 37 °C for 1 h with 10 μM Nle–Leu–Phe, 10 μM ETYIKWPWWVWL, 10 μM each of both peptides or no peptides. DMBSO was the same concentration in all cases. The cells were washed with 10 ml 10 mM phosphate, pH 7.4, 3 mM KCl, 100 mM NaCl. Cells were resuspended in 10 mM phosphate, pH 7.4, 3 mM KCl, 100 mM NaCl, 0.02% saponin, 60 μg/ml leupeptin, protein phosphatase inhibitors (Sigma) 1/100, and 8 μM of fluorescein labeled anti arrestin. The cells were incubated for 15 h at 4 °C, diluted 1 to 3 with 10 mM phosphate, pH 7.4, 3 mM KCl, 100 mM NaCl, and then analyzed by FACScan. Background was determined by adding unlaeled anti-arrestin in 10 fold molar excess to the fluorescein labeled anti arrestin.

2.7. Downregulation of surface FPR

CHO S cells expressing FPR were incubated in Gibco CHO-S-SFMII at 37 °C for 1 h with 10 μM Nle–Leu–Phe, 10 μM ETYIKWPWWVWL, 10 μM each of both peptides or no peptides. DMBSO was the same concentration in all cases. The cells were washed 2× with 10 ml 10 mM phosphate, pH 7.4, 3 mM KCl, 100 mM NaCl at 4 °C. The cells were resuspended in 100 mM NaCl, 3 mM KCl, 10 mM sodium phosphate, pH 7.4, 1 mM Mg++, 1 mM Ca++, 5% fetal bovine serum and 50 nM formyl-Nle–Leu–Phe–Nle–Tyr–Lys–FITC and incubated for 1 h at 4 °C and analyzed by FACScan as described above. Background observed with non-expressing CHO S cells was subtracted.

2.8. Internalization of ETYIK-(Alexa Fluor)WPWWVWL

CHO S cells expressing FPR were incubated in Gibco CHO-S-SFMII at 37 °C for 1 h with varying concentrations ETYIK-(Alexa Fluor)WPWWVWL. The cells were washed 2× with 10 ml 10 mM phosphate, pH 7.4, 3 mM KCl, 100 mM NaCl at 4 °C. The cells were resuspended in 100 mM NaCl, 3 mM KCl, 10 mM sodium phosphate, pH 7.4, 1 mM Mg++, 1 mM Ca++, 5% fetal bovine serum and 10 μM formyl-Nle–Leu–Phe and incubated for 1 h at 4 °C to displace any ligand bound to surface receptor. Internalized ligand was determined by FACScan as described above. Background observed with non-expressing CHO S cells (<20% observed with expressing cells) was subtracted.

2.9. Statistical analysis

Experimental results are expressed as the mean ± standard error of the mean. The significance was evaluated by unpaired t test (2 tailed) using the computer program Graphpad Prizm.

3. Results

3.1. Choice of viral peptides for analysis

Peptides derived from HIV-1 GP-41 have previously been shown to interact with FPR [4] and we previously observed that a gp41 36 mer (T20/DP178) inhibited formyl-Nle–Leu–Phe–Nle–Tyr–Lys–FITC binding to FPR [25]. Since gp41 is a fusion glycoprotein which contains two heptad repeat regions, we chose viral glycoproteins with this property. T20/DP178 is derived from the second heptad repeat region of HIV-1 gp41 but also includes residues C-terminal to the end of the heptad 2 repeat region [28]. Sainz et al. called these residues in HIV-1 and SARS coronavirus the “aromatic rich region” and they noted a similar region in Ebola virus [29]. This region has also been called the membrane proximal region. We therefore chose to synthesize peptides from HIV-2 and several human coronaviruses (SARS, OC43, and 229E) that corresponded to T20/DP178 in HIV-1. An alignment of sequences is shown in Table 1, which also includes the newly discovered coronaviruses HKU-1 and NL-63 [30–32] and the aromatic rich region from the Ebola spike protein.

The alignments show that all these viral sequences contain aromatic rich C-termini which are very rich in tryptophan. The 12 C-terminal residues of the coronaviruses are highly conserved. The 12 C-terminal residues of HIV-1 T20/DP178 are also highly conserved among known HIV-1 sequences and this 12 amino acid region overlaps the epitopes of 2 broadly protective anti-HIV antibodies 2F5 and 4E10 [33,34].12 mer peptides corresponding to C-termini of the sequences for HIV-1, HIV-2, SARS coronavirus, and coronavirus 229E were synthesized (residues are underlined in Table 1) as well as a 12 residue peptide corresponding to the aromatic rich region from the Ebola spike protein. We also synthesized smaller peptides of coronavirus 229E to localize residues which might be important in binding (ETYIKWP, WPWWVWL, and WWVWL).

3.2. Ligand specificity and affinity determination of R190/N192, W190/N192, and R190/K192 variants

We transformed CHO S cells with FPR variants R190/N192, W190/N192, and R190/K192. A schematic figure showing the topology of FPR is shown in Fig. 1. The location of R/W 190 and N/K 192 at the extracellular loop 2–TMH5 interface is indicated in the figure. Residues that have been shown in previous crosslinking and mutagenesis studies to be important in ligand binding are V83 (TMH2), R84 (TMH2), K85 (TMH2) indicated in the figure. Residues that have been shown in previous crosslinking and mutagenesis studies to be important in ligand binding are V83 (TMH2), R84 (TMH2), K85 (TMH2) [35]; L78A (TMH2), D106N (TMH3), L109A (TMH3), T157A...
R201A (TMH5), I204Y (TMH5), R205A (TMH2), W254A (TMH6), Y257A (TMH6) and F291A (TMH7) [26]. D106 and R201 are the sites of formyl group binding and R205 interacts with the C-terminus of fMLF [25]. The beginning of TMH5 was assigned to residue 191 and the end to residue 228 by hydrophobicity analysis according to Kyte and Doolittle [36]. We analyzed CHO S cells expressing R190/N192, W190/N192, and R190/K192 FPR for their ability to bind formyl-Nle–Leu–Phe–Nle–Tyr–Lys–FITC. We observed that the polymorphisms in FPR result in significantly altered formyl-Nle–Leu–Phe–Nle–Tyr–Lys–FITC binding affinity (Fig. 2) with R190/K192 having the highest affinity (\(K_d = 0.71 \pm 0.10\) nM), R190/N192 intermediate (\(K_d = 1.06 \pm 0.09\) nM), and W190/N192 having the lowest affinity (\(K_d = 1.68 \pm 0.13\) nM). We also synthesized formyl-Nle–Leu–Phe and determined the \(K_d\)s: R190/K192 (\(K_d = 0.76 \pm 0.05\) nM), R190/N192 (\(K_d = 1.20 \pm 0.14\) nM), and W190/N192 (\(K_d = 2.0 \pm 0.10\) nM), the results being essentially identical as that observed with formyl-Nle–Leu–Phe–Nle–Tyr–Lys–FITC.

We then determined the affinities of a broad range of peptides for these three polymorphisms based on their ability to compete with formyl-Nle–Leu–Phe–Nle–Tyr–Lys–FITC (Table 2). We observed that R190/K192 exhibited the highest affinity for formyl-Nle–Leu–Phe (chosen over fMLF in order to prevent possible oxidation of methionine) and MMWLL, a potent non-formylated chemoattract for FPR, [25,37]. Both formyl-Nle–Leu–Phe and MMWLL bound with \(\sim 3\) fold higher affinities to R190/K192 than to W190/N192. W190/N192 had the highest affinity for most virally derived peptides, especially the coronavirus peptides. W190/N192 bound the SARS coronavirus peptides with \(\sim 4–6\) fold higher affinity than either R190/N192 or R190/K192. This indicates that changes in these two positions affected ligand specificity, but most of the change in specificity was associated with the R to W change at residue 190. Most of the binding affinity of the coronavirus 229E peptides appears to reside in the 5 C-terminal residues since WWVWL exhibited only a modest reduction in affinity compared to ETYIKWPWWVWL. ETYIKWP was completely devoid of activity. WPWWVWL had similar affinity to WWVWL. ETYIKWPWWVWL bound to all three FPR variants with 15 to 30 fold higher affinity than the

Fig. 1. Transmembrane model of FPR. Residues in black are residues that have been shown in previous crosslinking and mutagenesis studies to be important in ligand binding [25,26,35]. Residues in gray are sites of known polymorphisms [13]. **R or W, *N or K.
N192 was determined as a function of ligand concentration. The greatest binding at saturation, but the peptides was specific for FPR, we labeled ETYIKWPWWVWL for W190/N192. WYVWL enhanced its affinity 6 fold for both R190/N192 and W190 can contact AlexaFluor during the excited state. This indicates that W190 quenches the fluorescence of formyl-Nle-Leu—Phe—Nle—Tyr—Lys—Alexa Fluor when the peptide is bound to W190/N192 FPR so that W190 can come in contact with the Alexa Fluor of ETYIK-(Alexa Fluor)WPWWVWL when bound to FPR. After correcting for the quenching of formyl-Nle—Leu—Phe—Nle—Tyr—Lys—Alexa Fluor on W190/N192, we estimate that ETYIK-(Alexa Fluor)WPWWVWL is internalized at about 20–30% of the rate of formyl-Nle—Leu—Phe—Nle—Tyr—Lys—Alexa Fluor in R190/N192, R190/K192, and W190/N192, based on their relative internalization rates by FACScan. This level of internalization was too low to accurately quantitate by direct fluorescence readings so the degree of quenching of Alexa Fluor- ETYIKWPWWVWL could only be estimated by FACScan analysis.

We also tested several peptides including formyl-Nle—Leu—Phe, ETKYIKWPWWVWL, MYVKWPWYVWL and formyl-MYVKWPWYVWL for their ability to stimulate GTPγS binding. Both 300 nM formyl-Nle—Leu—Phe and 300 nM formyl-MYVKWPWYVWL stimulated GTPγS binding in all three FPR variants, but neither ETYIKWPWWVWL nor MYVKWPWYVWL had any effect on the three FPR variants (Fig. 5). This indicated that the formyl group on formyl- MYVKWPWYVWL was essential for agonist activity but not for binding. 2 μM MYVKWPWYVWL inhibited 300 nM formyl-MYVKWPWYVWL stimulation of GTPγS binding by W190/N192 FPR by 60% but did not have a significant effect on formyl-MYVKWPWYVWL stimulation of GTPγS binding in either R190/N192 FPR or R190/K192 FPR. This is consistent with the data in Table 2 where formylation of MYVKWPWYVWL enhanced the affinity for R190/N192 and R190/K192 by 6 fold but only 2 fold for W190/N192. We then tested ETKYIKWPWWVWL for its ability to inhibit the stimulation of those required for receptor saturation. Since the peptides behaved as antagonists, we were somewhat surprised that they were so readily internalized. This probably reflects turnover of FPR on the cell surface, since treatment of the cells with ETKYIKWPWWVWL did not result in net receptor down regulation (see Fig. 9) below. Nonetheless, it indicates that if a virus interacts with FPR, it could be effectively internalized even if it is unable to activate the receptor.

We also quantitated formyl-Nle—Leu—Phe—Nle—Tyr—Lys—Alexa Fluor internalization in R190/N192, R190/K192, and W190/N192, by fluorescence spectroscopy in the absence or presence of 0.1% SDS to correct for any quantum yield change in Alexa Fluor fluorescence which might occur upon binding to FPR. Interestingly, formyl-Nle—Leu—Phe—Nle—Tyr—Lys—Alexa Fluor fluorescence internalized by W190/N192 FPR was enhanced in the presence of 0.1% SDS by 95±5%, but 0.1% SDS did not affect the fluorescence of formyl-Nle—Leu—Phe—Nle—Tyr—Lys—Alexa Fluor internalized by either R190/N192 FPR or R190/K192 FPR. This indicates that W190 quenches the fluorescence of formyl-Nle—Leu—Phe—Nle—Tyr—Lys—Alexa Fluor by ∼50% when it is bound to the receptor and thus must be sufficiently near the Alexa Fluor moiety of formyl-Nle—Leu—Phe—Nle—Tyr—Lys—Alexa Fluor when the peptide is bound to W190/N192 FPR so that W190 can contact AlexaFluor during the excited state [38]. This indicates that W190 resides in or very close to the ligand binding pocket (∼1 to 1.5 nm average distance [38]). Since Alexa Fluor- ETKYIKWPWWVWL exhibited twice the fluorescence of formyl-Nle—Leu—Phe—Nle—Tyr—Lys—Alexa Fluor when bound to W190/N192 (the expected value if it was not quenched), it is unlikely that W190 can come in contact with the Alexa Fluor of ETYIK-(Alexa Fluor)WPWWVWL when bound to FPR.

In order to determine whether the binding of coronavirus peptides was specific for FPR, we labeled ETKYIKWPWWVWL with Alexa Fluor 488, purified it by HPLC, and compared the binding to CHO S cells expressing R190/N192, W190/N192, R190/K192 or cells not expressing FPR. All three FPR variants exhibited binding of Alexa Fluor 488 labeled peptide, compared with non-expressing cells (Fig. 3). W190/N192 exhibited the greatest binding at saturation, but the $K_s$ for R190/N192, R190/K192, and W190/N192 were similar: 121±18 nM, 115±10 nM, and 99±10 nM, respectively. When ETKYIK-(Alexa Fluor)WPWWVWL and formyl-Nle—Leu—Phe—Nle—Tyr—Lys—Alexa Fluor binding to R190/N192, R190/K192, and W190/N192 were determined at saturating concentrations of either ligand, Alexa Fluor- ETKYIKWPWWVWL exhibited twice the fluorescence of formyl-Nle—Leu—Phe—Nle—Tyr—Lys—Alexa Fluor when bound to W190/N192, but only about 60% the fluorescence of formyl-Nle—Leu—Phe—Nle—Tyr—Lys—Alexa Fluor bound to either R190/N192 or R190/K192 (data not shown, but see below for an explanation).

We also investigated the ability of ETKYIK-(Alexa Fluor)WPWWVWL to be internalized at 37°C. The results are shown in Fig. 4. The EC$_{50}$s for internalization were ∼4 fold lower than the $K_s$ for binding (R190/N192, R190/K192, and W190/N192: 29±5 nM, 27±8 nM, and 20±3 nM, respectively). This indicates that either the affinity of ETKYIK-(Alexa Fluor)WPWWVWL for FPR is greater at 37°C used for internalization than at 4°C used in the binding experiments to prevent internalization or that saturation of the internalization process occurs at concentrations well below HIV-2 12 mer (KLN$\text{N}$SWDVFGNW$\text{F}$) indicating a preference of coronavirus peptides over HIV-2 peptides. The reverse peptide, LWVWWPWK$\text{I}$YTE, exhibited very little inhibition at concentrations up to 40,000 nM. Formylation of MYVKWPWYVWL enhanced its affinity 6 fold for both R190/N192 and R190/K192 but had only a very modest 2 fold affect on affinity for W190/N192.

![Fig. 2. Determination of affinity of ligand binding. Binding of formyl-Nle—Leu—Phe—Nle—Tyr—Lys—Alexa Fluor to FPR variants R190/N192, R190/K192, or W190/N192 was determined as a function of ligand concentration.](image-url)
of GTPγS by 300 nM formyl-Nle–Leu–Phe (Fig. 6). ETYIKWPWWVWL was more effective in inhibiting the stimulation of GTPγS binding in W190/N192 (EC50=300±120 nM; (P=0.003 vs. R190/K192; P=0.005 vs. R190/N192) than either R190/N192 (EC50=1600±400 nM) or R190/K192 (EC50=1600±440 nM). These results are consistent with the results in Table 2, which show that W190/N192 has a similar affinity for formyl-Nle–Leu–Phe and ETYIKWPWWVWL but that R190/N192 and R190/K192 have 6–8 fold higher affinities for formyl-Nle–Leu–Phe than ETYIKWPWWVWL. We also tested ETYIKWP and WWVWL for their ability to inhibit the stimulation of GTPγS binding by 300 nM formyl-Nle–Leu–Phe in W190/N192 FPR. The results are shown in Fig. 7. WWVWL inhibited formyl-Nle–Leu–Phe stimulation of GTPγS binding with a slightly higher EC50 (EC50=1300 nM) than ETYIKWPWWVWL (EC50=300±120 nM). ETYIKWP had no effect. Again this is consistent with the results shown in Table 2. We also tested the HIV-1 36 mer and the Ebola virus 12 mer for their ability to inhibit the stimulation of GTPγS binding by 300 nM formyl-Nle–Leu–Phe in W190/N192 FPR. The results are shown in Fig. 8. Formyl-Nle–Leu–Phe caused β-arrestin translocation in CHO S cells expressing R190/N192, R190/K192, and W190/N192 but not in CHO S cells not expressing FPR. ETYIKWPWWVWL did not cause translocation in CHO S cells expressing R190/N192, W190/N192, or R190/K192 FPR. ETYIKWPWWVWL inhibited translocation in response to formyl-Nle–Leu–Phe by 18±4% in R190/N192 FPR expressing cells, 20±9% in R190/K192 FPR expressing cells, and 77±10% in W190/N192 FPR expressing cells (P<0.006; W190/N192 vs. either R190/N192 or R190/K192). Again ETYIKWPWWVWL was more effective against W190/N192 than with R190/N192 or R190/K192, but it exhibited only antagonist activity in this assay.

Finally, we tested R190/N192, R190/K192, and W190/N192 expressing CHO S cells to undergo surface receptor downregulation in response to 10 μM formyl-Nle–Leu–Phe, 10 μM ETYIKWPWWVWL, or 10 μM each of both peptides (Fig. 9). Formyl-Nle–Leu–Phe caused β-arrestin dependent signaling pathways [39–42]. Therefore, we also tested R190/N192, W190/N192, and R190/K192 expressing CHO S cells for the ability to undergo β-arrestin translocation in response to 10 μM formyl-Nle–Leu–Phe, 10 μM ETYIKWPWWVWL, or 10 μM each of both peptides (Fig. 8). Formyl-Nle–Leu–Phe caused β-arrestin translocation in CHO S cells expressing R190/N192, R190/K192, and W190/N192 but not in CHO S cells not expressing FPR. ETYIKWPWWVWL did not cause translocation in CHO S cells expressing R190/N192, W190/N192, or R190/K192 FPR. ETYIKWPWWVWL inhibited translocation in response to formyl-Nle–Leu–Phe by 18±4% in R190/N192 FPR expressing cells, 20±9% in R190/K192 FPR expressing cells, and 77±10% in W190/N192 FPR expressing cells (P<0.006; W190/N192 vs. either R190/N192 or R190/K192). Again ETYIKWPWWVWL was more effective against W190/N192 than with R190/N192 or R190/K192, but it exhibited only antagonist activity in this assay.
ETYIKWPWWVWL did not cause surface receptor down-regulation in CHO S cells expressing R190/N192, R190/K192, or W190/N192 FPR. ETYIKWPWWVWL inhibited receptor down-regulation in response to formyl-Nle-Leu-Phe by 16±2% in R190/N192 FPR expressing cells, 20±2% in R190/K192 FPR expressing cells, and 64±6% in W190/N192 FPR expressing cells (P<0.001 for W190/N192 vs. either R190/N192 or R190/K192). As with the previous experiments, ETYIKWPWWVWL was more effective against W190/N192 than R190/N192 or R190/K192. The effects on downregulation closely correlated with receptor affinity. 

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Fig. 3. Determination of affinity of Alexa Fluor labeled ETYIKWPWWVWL. Binding of Alexa Fluor labeled ETYIKWPWWVWL to W190/N192 FPR, R190/K192 FPR, and R190/N192 FPR and to CHO-S cells not expressing FPR are shown as a function of concentration. K_d for W190/N192, R190/K192, and R190/N192 were 99±10 nM, 115±10 nM, and 121±18 nM, respectively.

Fig. 4. ETYIKWPWWVWL labeled with Alexa Fluor 488 is internalized in CHO cells expressing FPR. CHO cells expressing FPR W190/N192, FPR R190/N192 or FPR R190/K192 were incubated with indicated concentrations of Alexa Fluor labeled peptide for 37 °C for 1 h and the amount of internalized ligand was determined. The very low internalization of CHO cells not expressing FPR has been subtracted.

Fig. 5. Stimulation of GTPγS binding by formyl-Nle-Leu-Phe, formyl-MYVKWPWYVWL, MYVKWPWYVWL or ETYIKWPWWVWL in cells expressing W190/N192, R190/K192, or R190/N192 FPR.

Fig. 6. Inhibition of stimulation of GTPγS binding induced by formyl-Nle-Leu-Phe by ETYIKWPWWVWL in FPR variants R190/N192, R190/K192, or W190/N192. Inhibition of FPR activation by viral peptides. Inhibition of stimulation of GTPγS binding induced by formyl-Nle-Leu-Phe by ETYIKWPWWVWL, ETYIKWP, or WWVWL in W190/N192 FPR.
with β-arrestin translocation, and this was not unexpected given association between β-arrestin translocation and down-regulation [43].

4. Discussion

R190/K192 exhibited the highest affinity for formyl-Nle–Leu–Phe, formyl-Nle–Leu–Phe–Nle–Tyr–Lys–Alexa Fluor, and formyl-MYVKWPWWVWL, which indicates that it is not defective in binding formyl peptides. African American patients expressing the R190/K192 polymorphism were associated with increased risk of aggressive periodontitis whereas those expressing the W190/N192 polymorphism were associated with reduced risk [24]. Our data clearly show no evidence that R190/K192 is defective and we find that W190/N192 exhibited a 2.6 fold lower affinity for formyl peptides than R190/K192. Zhang et al. also reported that R190/K192 was not associated with an increased risk of aggressive periodontitis in either Brazilians or Turks nor was W190/N192 associated with reduced incidence of aggressive periodontal disease in these two populations [24] so it was unclear why this association was only true for African Americans. A recent study by Maney et al. [44] was unable to confirm the association of aggressive periodontitis with R190/K192 in African Americans, but did find an association with a silent polymorphism I116I. Our data finding no defect in R190/K192 are consistent with the study of Maney et al. [44] and also with the observation that R190/K192 was not associated with increased risk for aggressive periodontitis in either Turks or Brazilians.

In Fig. 1, we show the relative positions of R/W 190 and N/K 192 at the extracellular loop 2-TMH5 interface to other residues in FPR which we previously identified as important in ligand binding [25,26,35]. We previously determined that D106 and R201 likely form an ion pair between TMH3 and TMH5 and that this ion pair imparts formyl group selectivity to ligand binding since mutation of either residue changes the preference of formylated to non-formylated peptide from 4000/1 to ∼1/3 and 1/1, respectively, for D106N and R201A [25].

Fig. 7. Inhibition of FPR activation by viral peptides. Inhibition of stimulation of GTPγS binding induced by formyl-Nle–Leu–Phe by ETYIKWPWWVWL, ETYIKWP, or WWVWL in W190/N192 FPR.

In Fig. 8, we show the relative positions of R/W 190 and N/K 192 in the presence of various FPR ligands. Relative arrestin translocation was determined in non-expressing CHO cells expressing R190/N192 FPR, R190/K192 FPR expressing cells, or W190/N192 FPR expressing cells in response to 10 μM formyl-Nle–Leu–Phe, 10 μM ETYIKWPWWVWL, or 10 μM formyl-Nle–Leu–Phe + 10 μM ETYIKWPWWVWL. 10 μM ETYIKWPWWVWL inhibited translocation in response to 10 μM formyl-Nle–Leu–Phe by 18±4% in R190/N192 FPR expressing cells, 20±4% in R190/K192 FPR expressing cells, and 77±10% in W190/N192 expressing cells (P<0.006 for W190/N192 FPR vs. either R190/N192 or R190/K192).
better than R190/N192 or R190/K192, it is likely that W190 interacts with some as yet unidentified residues in these peptides. One explanation of why R190/K192 binds formyl peptides better than W190/N192 is that the former contains two additional positive charges. Formylation removes the positive charge from the N-terminus of peptides. If R190 is near the entry site of the ligand, the ligands with free N-termini and peptides with net positive charges are likely to be attracted toward R190 but not W190. W190 would be expected to exhibit a relative preference for peptides with free N-termini and peptides with net positive charge over R190/K192.

Su et al. [4] originally observed that DP178/T-20, a peptide derived from the C-terminal HR and membrane proximal region of gp41 of human immunodeficiency virus type 1 (HIV-1), elicited chemotaxis in neutrophils and monocytes. They determined that this chemotaxis was a result of its ability to bind to and activate FPR. Su et al. [4] also observed that shorter versions of DP-178 acted as antagonists of FPR. We subsequently showed that DP178 binds to FPR expressed in CHO S cells and observed that two FPR mutants (D106N and R201A), which bind fMLF poorly, exhibited an enhanced ability (30 fold and 3 fold, respectively) to bind DP178/T20 [25]. Here we observed that R190/N192, W190/N192, and R190/N192 FPR variants all exhibited similar affinities for DP178/T20 and that DP178/T20 behaved as an antagonist based upon its inability to stimulate GTPγS binding and its ability to inhibit the stimulation of GTPγS by formyl-Nle–Leu–Phe.

HR regions appear to be a common motif in many viral fusion proteins [45]. They exist as sequential helical domains; one n terminal HR region (HR1) adjacent to the fusion peptide and a c-terminal HR region (HR2) close to the transmembrane anchor. Peptides derived from the HR2 and proximal membrane regions of HIV-1, SIV, HRSV, human parainfluenza virus type 3, measles virus, and a coronavirus have all been shown to be able to block virus infection [28,46–48]. A twelve mer peptide derived from the proximal membrane region of FIV gp41, QLQKWEDWWVL, was a potent (low nanomolar) inhibitor of FIV infection [49]. Here we observed that peptides derived from the proximal membrane region of HIV-1, HIV-2, Ebola virus, SARS coronavirus, and coronavirus 229E act as antagonists to the formyl peptide receptor. The coronavirus peptides are the most potent antagonists and these peptides exhibited an enhanced affinity for W190/N192 FPR. Our observation that tryptophan-rich peptides act as antagonists is consistent with the observation of Bae et al., who observed that WRWWWW was a potent antagonist for FPRL1 and FPRL2 [50,51]. WRWWWW did not affect FPR [50], and it has been shown that DP178/T20 does not bind to FPRL1 [4], indicating that these receptors have slightly different ligand binding specificities but that both receptors may bind tryptophan rich peptides. A recently discovered FPRL2 agonist is also tryptophan rich [52] and the C-terminus of that peptide, WPWQVL, is remarkably similar to WPWWVVL which we found to be a potent antagonist for FPR.

Figure 9. Down regulation of FPR surface expression in response to various ligands.

Surface expression of FPR in W190/N192 FPR expressing cells, R190/K192 FPR expressing cells, or R190/N192 FPR expressing cells was determined after exposure for 1 h at 37°C to either 10 μM formyl-Nle–Leu–Phe, 10 μM ETYIKWP–WWVVL, or 10 μM formyl-Nle–Leu–Phe +10 μM ETYIKWPWWVL. ETYIKWPWWVL inhibited receptor downregulation in response to formyl-Nle–Leu–Phe by 16±2% in R190/N192 FPR expressing cells, 20±2% in R190/K192 FPR expressing cells, and 64±6% in W190/N192 FPR expressing cells. (P<0.001 W190/N192 vs. either R190/N192 or R190/K192).
situation. Klestadt et al. [57] observed that activation of FPR in THP-1 cells markedly downregulated the expression of CD4 and CD13, the receptors for HIV-1/HIV-2 and coronavirus 229E, respectively [11,58,59]. In addition, activation of FPR downregulates the expression of CCR5 and CXCR4 [60], the co-receptors of HIV-1 and HIV-2 [9,58]. Thus activation of FPR could serve as protection against viral infection by downregulation of viral receptors, and presumably FPR antagonists would prevent down-regulation. Ueda et al. noted that the HIV-1 envelope gp41 was able to down regulate the expression of FPR and other chemokine receptors at low nanomolar concentrations, but that the down-regulation of FPR and the chemokine receptors was dependent upon expression of CD4 [61]. Thus, there are a complicated set of interactions between FPR and the receptors and co-receptors of viruses, and FPR may play an important role in viral infections.

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