Histatin 5 binds to *Porphyromonas gingivalis* hemagglutinin B (HagB) and alters HagB-induced chemokine responses

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Histatins are human salivary gland peptides with anti-microbial and anti-inflammatory activities. In this study, we hypothesized that histatin 5 binds to *Porphyromonas gingivalis* hemagglutinin B (HagB) and attenuates HagB-induced chemokine responses in human myeloid dendritic cells. Histatin 5 bound to immobilized HagB in a surface plasmon resonance (SPR) spectroscopy-based biosensor system. SPR spectroscopy kinetic and equilibrium analyses, protein microarray studies, and I-TASSER structural modeling studies all demonstrated two histatin 5 binding sites on HagB. One site had a stronger affinity with a $K_D = 1.9 \text{ mM}$ and one site had a weaker affinity with a $K_D = 60.0 \text{ mM}$. Binding has biological implications and predictive modeling studies and exposure of dendritic cells both demonstrated that $20.0 \text{ mM}$ histatin 5 attenuated (p < 0.05) $0.02 \text{ mM}$ HagB-induced CCL3/MIP-1α, CCL4/MIP-1β, and TNFα responses. Thus histatin 5 is capable of attenuating chemokine responses, which may help control oral inflammation.

Saliva contains an elaborate mixture of peptides, proteins, chemokines, cytokines, growth factors, and antimicrobial peptides. Some peptides and proteins play key roles in oral lubrication, mastication, digestion, and the demineralization and remineralization of teeth, whereas others play key roles in innate immune defense of oral tissues. Among the latter are the histatins, a group of low molecular weight histidine-rich peptides and peptide fragments produced by the human parotid, submandibular, and sublingual salivary glands. Although 24 histatin fragments are known, there are 12 primary histatins arising from *HIS1* and *HIS2* gene products. Histatins 1 and 3 arise by initial proteolytic events and histatin 5 is a cleavage product of histatin 3. Histatins bind to specific microorganisms like the periodontal pathogen *P. gingivalis*. They also have broad-spectrum antibacterial activity against other oral species of *Streptococcus*, *Actinomyces*, and *Porphyromonas*; antifungal activity against species of *Candida*, *Saccharomyces*, and *Cryptococcus*; bind copper and zinc; inhibit proteases and toxins; inhibit hemagglutinating activity of some oral bacteria and inhibit co-aggregation of other oral bacteria; inhibit lipopolysaccharide-mediated gelation of *Limulus* amoebocyte lysate; reverse the anti-complement action of lipopolysaccharide or lipid A; and enhance wound closure. Some histatin fragments also have anti-microbial, DNA-binding, and anti-inflammatory activities. Histatin 5 is one of these and pretreatment of outer-membrane protein from *P. gingivalis* with $10.0 \text{ µg/ml}$ (e.g., $3.3 \text{ mM}$) synthetic histatin 5 causes a 37.0% inhibition of IL6 and a 47.0% inhibition of IL8 production in human gingival fibroblasts.

Hemagglutinin B (HagB) is a major virulence factor of *P. gingivalis* involved in non-fimbrial adhesion of the microorganism to host cells. It is a 49.0 kDa protein composed of 350 amino acid residues with a pI of 8.42.
Recombinant HagB induces a robust chemokine and cytokine response in dendritic cells\(^{16}\). In this study, we hypothesized that histatin 5 binds to recombinant HagB and attenuates HagB-induced chemokine responses in dendritic cells. Our objectives were to i) characterize the interaction of histatin 5 with immobilized HagB by surface plasmon resonance (SPR) spectroscopy using kinetic and equilibrium analyses; ii) identify histatin 5 binding sites on HagB using protein microarrays and I-TASSER molecular modeling; and iii) determine the ability of histatin 5 to attenuate a HagB-induced chemokine and cytokine response using in silico dendritic cell functional simulation modeling of signaling pathways and direct exposure of dendritic cells (Figure 1).

**Results**

**Binding of histatin 5 to immobilized HagB.** Our first objective was to characterize the interaction of histatin 5 with immobilized HagB by kinetic and equilibrium analyses of SPR spectroscopy data (Figure 1). HagB was immobilized at two surface densities to a COOH\(_2\) sensor chip using a standard amine coupling method. A third flow channel (FC) was activated and deactivated to serve as a reference in data analysis. Initial binding tests were performed to assess the specificity of histatin 5 for immobilized HagB. In the first test, performed in running buffer without CM-Dextran, notable nonspecific binding of histatin 5 was observed to the reference channel. However, when 0.5 mg/ml CM-Dextran was introduced to the buffer, the non-specific binding became negligible. Subsequent studies on the kinetic binding of histatin 5 to HagB were performed in running buffer with 0.5 mg/ml CM-Dextran. Histatin 5 was observed to fully dissociate from both the high density HagB surface (Figure 2a) and the low density HagB surface (Figure 2b) without the use of regeneration.

The binding responses of histatin 5 demonstrated apparent heterogeneous characteristics, which were thought to be due to the presence of two independent binding sites on immobilized HagB. (Table 1) Equilibrium analysis of a fixed concentration histatin 5 assay (Figure 3a) showed that the dose response plot conformed well to a two-site model (Figure 3b). The maximum binding responses (Rmax) of these data further confirm a multi-site interaction by being larger than the theoretical Rmax. For example, a 1:1 interaction of analyte (MW 3,034.0 Da) and ligand (49.0 kDa) the expected Rmax for FC 2 is 88.0 RU while in Figure 2b responses of >95.0 RU are observed. While individually these results do not conclusively identify a heterogeneous ligand, their agreement suggests this as a valid hypothesis.

Two-site analysis by fitting equilibrium models to the fixed concentration assay indicated the high affinity site to have a KD of 1.9 \(\mu\)M and the weaker affinity site to have a KD of 60.0 \(\mu\)M. Kinetic analysis using the OneStep\(^{\text{TM}}\) injection, which is modeled by Taylor Dispersion theory, enabled a more accurate estimation of the rate constants for each interaction site\(^{16,18}\). In this injection an internal dispersion line was used to create an analyte gradient of histatin 5, which was then flowed over the FC from low to high concentration (Figure 3c). The gradient binding data were analyzed using a kinetic two-site model, which estimates rate constants for the two individual interactions. (Table 2) The \(K_D\) values determined from the kinetic model fit were 340.0 ± 10.0 and 260.0 ± 10.0 nM from the high and low-density HagB surfaces, respectively. The \(K_D\) values determined were 11.3 ± 0.07 and 15.2 ± 0.2 \(\mu\)M from the high and low-density HagB surfaces, respectively.

A final objective for the SPR spectroscopy experiments was to compare HagB binding of HBDs with histatin 5. HBD1 and HBD3 were observed to bind the sensor chip with more non-specific binding than histatin 5 (data not shown). The concentration of CM-Dextran was therefore increased to 2.0 mg/ml for the HBD assays (Figure 4). HBD1 was observed to have a weak affinity interaction with HagB as expected while HBD3 demonstrated a heterogeneous

![Flowchart](https://www.nature.com/scientificreports/srep03904/flowchart.png)

**Figure 1 | A flow chart on the study design.** SPR spectroscopy kinetic and equilibrium analyses, protein microarray studies, and I-TASSER structural modeling studies all demonstrated two histatin 5 binding sites on HagB. - One site had a stronger affinity with a KD1 of 1.9 \(\mu\)M and one site had a weaker affinity with a KD2 of 60.0 \(\mu\)M. Predictive modeling studies and exposure of dendritic cells both demonstrated that 20.0 \(\mu\)M histatin 5 attenuated (p < 0.05) 0.02 \(\mu\)M HagB-induced CCL3/MIP-1\(\alpha\), CCL4/MIP-1\(\beta\), and TNF\(\alpha\) responses.
binding response, which was not able to be fit with a standard interaction model. The NaCl concentration of the buffer was increased to 0.3 M and the assay was repeated for both defensins (Figure 5). HagB binding of HBD1 appeared to be slightly modified by the change in salt concentration. HagB binding of HBD3 however was significantly affected by the change in salt concentration such that an equilibrium model could be fit to the dose response plot (Figure 5d).

Identification of binding sites on HagB. Identification of histatin 5, HBD1, and HBD3 binding sites on HagB began with their differential binding to the 350 different, overlapping 15mer peptides in the HagB peptide microarrays. Histatin 5-Cy5-NHS, HBD1-Cy5-NHS, and HBD3-Cy5-NHS bound to select 15mer peptides in each array resulting in higher fluorescent intensities than that of adjacent overlapping 15mer peptides on either side. To identify a binding site we assumed that for two overlapping peptides, the significant increase or decrease of fluorescent intensity of one peptide compared with the other adjacent peptide is due to the addition or deletion of a new or old residue. For example, in the histatin 5-HagB microarray, the fluorescent intensity increased from 0.0 to 22.0 from the 15mer peptide (31–45, NIAKLNPKLPELEKA) to peptide (32–46, IAKLNPKLPELEKAI). Based on our assumption, the increase would be due to the addition of the residue I46.

Histatin 5, at 1.0 μg/ml, bound to HagB residues I32, I46, Y76, F88, D117, N154, R165, I242, E256, and Y291. These residues can be mapped to two potential binding sites using the I-TASSER structural model (Figure 6a). The first site contained residues I32, I46, Y76, and F88 (colored in red) and the second site contained residues D117, N154, and R165.

Table 1 | Affinity constants for HagB analytes by equilibrium analysis

| Analyte | Langmuir Model | $K_D1$ (μM) | $K_D2$ (μM) | $R^2$ of fit |
|---------|----------------|-------------|-------------|--------------|
| Histatin 5 | One-site | 9.7 ± 0.8 | N/A | 0.9830 |
| Histatin 5 | Two-site | 1.9 ± 0.6 | 60.0 ± 0.4 | 0.9940 |
| HBD1 | One-site | 1000.0 ± 1000.0 | N/A | 0.9809 |
| HBD3 | One-site | 35.0 ± 8.0 | N/A | 0.9911 |

Figure 2 | SPR spectroscopy data showing histatin 5 binding to immobilized HagB. (a,b) Referenced SPR spectroscopy data showing specific binding of histatin 5 to HagB at a higher surface density of HagB (a) and a lower density of HagB (b).
N154, and R165 (colored in blue); the other three residues (I242, E256, and Y291, colored in green) might belong to the second binding site but they are too far away from the first binding site. This might be caused by the inaccurate structure modeling at the C-terminus (modeled as loop), where the residues I242, E256, and Y291 are located.

HBD1, at 1.0 mg/ml, bound to HagB residues Y17, L26, K38, Y76, D89, D103, Y213, V261, R273, Y291, and L350. The residues L26, K38, Y76, D89, D103, and Y213 (in red color) likely form one binding site according to the I-TASSER model (Figure 6b). Residues V261, R273, Y291, and L350 are located the loop region of the I-TASSER model, which have low model quality and thus it is difficult to make a conclusion for these residues.

HBD3, at 1.0 mg/ml, bound to HagB residues N31, N54, K118, L147, E152, D199, L212, A281, and G295 and HBD3 at 2.0 mg/ml bound to HagB residues V3, N13, D69, A98, K118, K131, K141, E152, D160, L212, K271, D285, and S338. Using the I-TASSER model and the consensus of the two experiments, we conclude that HBD3 binds to residues K118, K131, L147, E152, and D160 (colored in red in Figure 6c) and this may represent a possible binding site.

**Predicted influence of histatin 5 on a HagB-induced dendritic cell response.** The *in silico* dendritic cell functional simulation model predicted that histatin 5 would reduce the HagB-induced IL8 response by 31.8%, the CSF2/GM-CSF response by 55.9%, the CCL3/MIP-1α response by 42.0%, the CCL4/MIP-1β response by 44.4%, and the TNFα response by 46.5% (Figure 7a, b).

**Influence of histatin 5 on a HagB-induced dendritic cell response.** None of the solutions, media, or inocula containing 0.2, 2.0, or 20.0 μM histatin 5 or 0.02 or 0.2 μM HagB induced any cytotoxicity in the dendritic cell cultures at 16.0 hours.

0.02 μM HagB induced strong IL8 responses at 2.0 and 4.0 hours that peaked at 8.0 hours and began to decline at 16.0 hours post exposure. Although histatin 5 appeared to be able to attenuate a HagB-induced IL8 response at 8.0 hours in some experimental replications, it was not consistent in all primary dendritic cell cultures and often lost by 16.0 hours (not shown).

0.02 μM HagB also induced strong CCL3/MIP-1α, CCL4/MIP-1β, and TNFα responses and a weaker CSF2/GM-CSF response (Figure 8). At 16.0 hours, a 100:1 micromolar ratio of histatin5:
HagB, histatin 5 induced a 9.3% decrease in CCL3/MIP-1α, 12.5% decrease in CCL4/MIP-1β, a 25.1% decrease in TNFα, and a 42.0% decrease in CSF2/GM-CSF (Figure 8). At 16.0 hours at a 1,000 m M. Histatin 5 therefore has a more favorable affinity for HagB than in 5 micromolar ratios of histatin 5 in CCL4/MIP-1β, and a 42.4% decrease in TNFα.

**Discussion**

Histatins are a group of low molecular-weight histidine-rich peptides produced in abundance by the human parotid, submandibular, and sublingual salivary glands. Concentrations in parotid saliva range from 7.0 to 28.0 μg/ml for histatin 1, 6.0 to 43.0 μg/ml for histatin 3, and 10.0 to 43.0 μg/ml for histatin 5 and concentrations in submandibular and sublingual saliva range from 28.0 to 122.0 μg/ml for histatin 1, 15.0 to 75.0 μg/ml for histatin 3, and 26.0 to 90.0 μg/ml for histatin 5.

Histatins have a variety of innate immune functions. They bind to bacteria, inhibit co-aggregation of oral bacteria, and have broad-spectrum antimicrobial activity against bacteria and fungi (e.g., C. albicans). For example, 125I labeled histatin 5 binds to P. gingivalis 381: binding of histatin 5 is rapid, reversible, saturable and specific; the number of histatin 5-binding sites per cell was estimated to be 3,600; and the dissociation constant (Kd) was calculated to be 1.5 μM. The binding of histatins to virulence agonists alters the physiologic properties of the agonist and neutralizes toxins, inactivates proteases, and neutralizes the activities of lipopolysaccharide. Histatin 5 also has anti-inflammatory properties. Histatin 5 is known to suppress the production of IL6 and IL8 to P. gingivalis outer-membrane protein by human gingival fibroblasts.

In this study, we show that histatin 5 binds to P. gingivalis HagB. The interaction of histatin 5 with immobilized HagB was characterized by kinetic and equilibrium analysis of SPR spectroscopy data. Evidence for a two-site interaction was observed and considered when choosing the appropriate models. A higher affinity binding site was identified with a Kd of 340.0 or 260.0 nM when measured at two HagB surface densities. A weaker affinity site was calculated to have a Kd of 11.3 or 15.2 μM when measured at two HagB surface densities. We therefore conclude that histatin 5 has a moderate affinity for HagB that includes a weaker affinity interaction that could be due to the peptide’s basic isoelectric point. Analysis with OneStep™ provided additional confidence in the kinetic and affinity constants obtained for both interaction events.

We also showed that histatin 5 attenuates HagB-induced CCL3/MIP-1α, CCL4/MIP-1β, and TNFα responses at 16.0 hours post-inoculation of dendritic cells. Attenuation more readily occurs at 100:1 and 1,000:1 micromolar ratios of histatin 5:HagB than at 1:1 and 10:1 micromolar ratios of histatin 5:HagB (Figure 8). It is worth noting that 2.0 μM (6.1 μg/ml) and 20.0 μM (60.7 μg/ml) of histatin 5 are well within the physiologic concentration of histatin 5 reported above in the oral cavity.

HBD1 demonstrated a very weak affinity for HagB compared with histatin 5 and HBD3. HBD3 demonstrated a strong affinity with HagB that was highly charge based. After the non-specific binding of HBD3 was reduced by increasing the NaCl concentration, the resulting interaction affinity was estimated with a KD value of 35.0 μM. Histatin 5 therefore has a more favorable affinity for HagB than the defensin peptides and which is apparently more specific than just electrostatic attraction.

The exact mechanism for histatin 5 attenuation of the HagB-induced response is not yet fully known. It is easy to speculate that histatin 5 simply binds to HagB and inhibits its attachment to dendritic cell surfaces. If that were the case, we would expect to see attenuation of all of the HagB-induced chemokine and cytokine responses. More likely, histatin 5 may be first altering the binding of HagB to the...
surface of dendritic cells like that seen by HBD3 and then selectively altering HagB-induced signal transduction and transcription factors (Figure 7a), also like that seen by HBD3. The MAPK pathways are important in controlling the type and magnitude of the inflammatory response to *P. gingivalis* and its extracellular products including HagB, and histatin 5 may work via a similar mechanism.

We believe that histatin 5 may now be grouped with a family of anti-inflammatory components present in the oral cavity. These include salivary gland derived peptides, fibronectin, LL-37, lactoferrin, human neutrophil peptide (HNP) α defensins, and HBDs. Therefore, we believe that histatin 5, in the physiologic range of human saliva, has the ability to significantly attenuate chemokine and cytokine responses in dendritic cells exposed to HagB, a virulence factor of the periodontal pathogen *P. gingivalis*. Together with this family of anti-inflammatory peptides in saliva, histatin 5 may complement existing mechanisms to maintain oral health.

**Figure 5** | SPR spectroscopy data showing binding of HBDs with immobilized HagB. The assay buffer was modified to contain increased NaCl (0.3 M) and 2.0 mg/ml CM-Dextran. (a) HBD1 dilution series binding HagB. (b) Dose response plot of the interaction of HBD1 with HagB. A one-site Langmuir isotherm is shown applied to the data. (c) HBD3 dilution series binding HagB. (d) Dose response plot of the interaction of HBD3 with HagB. A one-site Langmuir isotherm is shown applied to the data.

**Figure 6** | The mapped binding sites of peptides to the predicted 3D I-TASSER structural model of HagB. (a) Histatin 5 binds to residues I32, I46, Y76, and F88 (colored in red) and to residues D117, N154, and R165 (colored in blue). (b) HBD1 binds to residues Y17, L26, K38, Y76, D89, D103, and Y213 (colored in red) likely forming one binding site. (c) HBD3 binds to residues K118, K131, L147, E152, and D160 (colored in red) likely forming one binding site. Histatin 5, HBD1, and HBD3 bind to HagB in different regions: histatin 5 and HBD1 bind more centrally to the HagB molecule whereas HBD3 binds near the end of the loop area. However, one binding site for histatin 5 is near the binding site for HBD3. This may account for the ability of histatin 5 to attenuate HagB-induced chemokine and cytokine responses in dendritic cells similar to that for HBD3, although much higher concentrations of histatin 5 are needed than HagB. For high-resolution images of the binding domains see Supplementary Figures 1a, b & c.
Figure 7 | The predictive in silico virtual dendritic cell platform generated results showing the percentage change with respect to the control reference baseline. (a) A high-level view of the HagB and histatin 5 regulation in the dendritic cell platform. HagB provides activating signals to CD14 and activin receptors. On binding to CD14 and activin receptors, it stimulates the release of pro-inflammatory cytokines like CCL3/MIP-1$\alpha$, CCL4/MIP-1$\beta$, CSF2/GM-CSF, IL8, and TNF$\alpha$ via activation of MYD88/TRAF6 and SHC1/ERK pathways, respectively. Both activin and CD14 signals converge at the activation of MAPKs, AKT, and PKC to activate transcription factors like NFKB, AP1, SP1, CREB1, NFAT, IRFs, etc., which transcribes different cytokines. Histatin 5 exerts inhibitory effect on dendritic cells. Histatin 5 negatively regulates the pro-inflammatory signal by inhibiting CD14 receptors. The dotted red lines in the figure indicate inhibitory links and dotted fluorescent green lines indicate activation links. The downstream effects and activation links of each transcription factor are shown by uniquely colored solid lines. (b) The illustrated effect of over-expression of histatin 5 along with HagB stimulation on CCL3/MIP-1$\alpha$, CCL4/MIP-1$\beta$, CSF2/GM-CSF, IL8 and TNF$\alpha$. Compared to HagB stimulation alone on the referenced baseline, the levels of CCL3/MIP-1$\alpha$, CCL4/MIP-1$\beta$, CSF2/GM-CSF, IL8 and TNF$\alpha$ had significantly reduced.
homeostasis and control and contain oral infection and inflammation by down regulating chemokine and pro-inflammatory cytokine responses.

In summary, histatin 5 behaves like HBD3. Both histatin 5 and HBD3 bind to HagB and both attenuate HagB-induced chemokine and cytokine responses in dendritic cells when co-incubated together at 37°C for 30 minutes with HagB. Although the trends of histatin 5 with HagB are reproducible, they are much more subtle than the trends seen with HBD3 and HagB and much higher concentrations of histatin 5 are needed. One explanation may be the difference in binding domains of histatin 5 and HBD3 on HagB (Figure 6): HBD3 binding occurred near the end of the loop whereas histatin 5 and HBD1 binding occurred more centrally to the HagB molecule. Future work will focus on identifying the chemokine and cytokine active domain on HagB and the binding domain of antimicrobial peptides on HagB.

Figure 8 | Influence of histatin 5 (H5) on a P. gingivalis HagB-induced CCL3/MIP-1α, CCL4/MIP-1β, TNFα, and CSF2/GM-CSF dendritic cell response. In one experiment (top line graph, each panel), dendritic cells were exposed to 0.01 M PBS, pH 7.2; 0.02 μM HagB; 2.0 μM histatin 5; or 2.0 μM histatin 5 + 0.02 μM HagB. Aliquots of cell culture media were removed at 0.0, 2.0, 4.0, 8.0, and 16.0 hours. At 16.0 hours at a 100:1 micromolar ratio of histatin 5:HagB, histatin 5 induced a 9.3% decrease in CCL3/MIP-1α, 12.5% decrease in CCL4/MIP-1β, a 25.1% decrease in TNFα, and a 42.0% decrease in CSF2/GM-CSF. In a second experiment increasing the concentration of histatin 5 10-fold (bottom bar graph, each panel), dendritic cells were exposed to 0.01 M PBS, pH 7.2; 0.02 μM HagB; 20.0 μM histatin 5; or 20.0 μM histatin 5 + 0.02 μM HagB. Aliquots of cell culture media were removed at 0.0, 2.0, 4.0, 8.0, and 16.0 hours. At 16.0 hours a 1000:1 micromolar ratio of histatin 5:HagB, histatin 5 induced a 49.8% decrease in CCL3/MIP-1α, 39.6% decrease in CCL4/MIP-1β, a 42.4% decrease in TNFα, and a 55.3% decrease in CSF2/GM-CSF. * = p < 0.05.
Methods

Reagents. All solutions, media, and cell culture inocula were prepared as previously described. 100 mM sodium phosphate buffer (pH 7.4), 0.14 M NaCl, pH 7.2 (Sigma-Aldrich, St. Louis, MO), 0.05% Tween 20, and 0.05 M NaCl contained 0.5 mg/ml CM-Dextran and 2.0 mg/ml CM-Dextran from 25.4 to 0.4 μM NaCl and 2.0 mg/ml CM-Dextran from 0.01 M PBS, pH 7.2 to 0.05% Tween 20, and 1% bovine serum albumin (BSA) with slight shaking for 1.0 hour at room temperature to reduce non-specific interactions with histatin 5, HBd1, or HBd3. 1.0 μg/ml histatin 5-C5- NHS, 1.0 μg/ml HBd1-C5-NHS, and 1.0 and 2.0 μg/ml HBd3-C5-NHS in 0.01 M PBS, pH 7.4, 0.05% Tween 20 with 0.1% BSA were added to the slide chambers. 1.0 μg/ml control peptide-Cy5-NHS; mouse monoclonal anti-α-HA antibody-Cy5; and mouse monoclonal anti-FLAG (M2) antibody-Cy3-NHS in 0.01 M PBS, pH 7.4, 0.05% Tween 20, with 0.1% BSA were added to control slide arrays. All slides were incubated with slight shaking overnight at 4°C. After 16.0 hours, the arrays were washed three times for one minute each with 0.01 M PBS, pH 7.4, 0.05% Tween 20.

The slides were scanned in a Typhoon 9410 Variable Mode Imager and the extent of histatin 5, HBd1, and HBd3 binding was determined from the corresponding images. Spot intensities were quantified with PepSlide® Analyzer. Based on average median foreground intensities, an intensity map was generated and binders in the peptide map highlighted by an intensity color code with red for high and white for low spot intensities. Peptides fragments on the array containing arginine and lysine-rich regions like PKKKKKK (e.g., HagB amino acid residues 309–314) tended to interact with labeled samples due to ion-exchange effects with the charged dye molecules and thus were regarded as non-specific.

Modeling the structure of HagB. The state-of-the-art protein structure prediction method I-TASSER34–36 was used to predict the structure and function of HagB. First, the HagB sequence was threaded over the PDB library37 by the meta-threading algorithm FTSETs37 to identify all structurally related templates from the threading templates were then reassigned into full-length models by replica-exchange Monte Carlo simulations with the threading unaligned regions (mainly loops) built by ab initio folding. The lowest free-energy conformation was selected by clustering the Monte Carlo simulation structures using SPICKER38. Next, fragment ensemble simulation was performed starting from the SPICKER38 generated ensembles of energetically feasible conformations, where the spatial restraints collected from both the LOMETS templates and the analog PDB structures by TM-align39 were used to guide the replica-exchange simulations. Finally, the models were refined in the atomic-level by the fragment-guided molecular dynamics (FG-MD) simulations40. To examine the biological functionality of the HagB sequence, the COFACTOR program41 was used to match the I-TASSER-derived and the protein function library BioLiP42 by local and global structural comparisons.

The overall topology of the HagB structure model was a three-helix bundle attached by a long loop at the C-terminus (Figure 6). The confidence score of the I-TASSER structure prediction was −2.52 with the estimated TM-score = 0.42 ± 0.14, which corresponds to an approximately correct topology. The major threading templates were selected by I-TASSER include human APPL1BAPR domain (PDBID: 4h8aA), nucleotide exchange factor Son of Sevenless (1zd4A), and BAR-PH domain of APPL1 (2q13A).

Modeling of the dendritic cell response of histatin 5 on HagB. An in silico dendritic cell functional simulation model was used to predict the effect of histatin 5 on the HagB-induced IL-8, CCL2/CM-CSE, CCL4/C5/MIP-1α, CCL4/C5/MIP-1β, and TNFα responses of dendritic cells43. This model provides a dynamic and transparent view of dendritic cell physiology at the functional proteomics abstraction level and predicts outcomes for different experimental scenarios in an automated high-throughput system. The model was manually aggregated, and is currently updated, from published research and collaborative data on signaling pathways, intermediates, transcription factors, enzyme kinetics and gene regulations comprising 3,860 biological species with over 15,433 cross-talk interactions44. The technology has been recently used to predict a) elastin triggered transient pro-inflammatory responses by human dermal fibroblasts45; b) anti-aggregation of alpha-synuclein in M17 dopaminergic cells46; c) TNFα activity of heat shock protein 90 in tumor cells47; d) effects on endothelial cells, and stromal cells48; e) effects on osteosarcoma cells and COX2 inhibitors on progesterone receptor isoforms A and B in myometrial cells49; d) ability of bicuspid to inhibit STAT3 expression in HepG2, SNU-387, and PLC/PRF5 human hepatocellular carcinoma cells50; e) the potential effects of isorhamnetin on the peroxisome proliferator-activated receptor gamma signaling cascade51.

For this study, HagB was thought to activate dendritic cells through TLR and also through C5 receptor signaling52. It was thought that the effect of HagB on dendritic cells by TLR signaling. HagB was introduced with a concentration of 8.0 × 10^4 μM along with a 5-fold over-expression of histatin 5 and the effect on downstream markers were analyzed compared to HagB stimulation alone on the referenced baseline. The first stimulation protocol response consisted of control simulation for 15,000 seconds for the simulated stimulation with 8.0 × 10^4 μM HagB, and trigger stimulation for 50,000 seconds. This response was compared with a second stimulation protocol response consisting of control simulation for 15,000 seconds, stimulation with 8.0 × 10^4 μM HagB, trigger simulation for 50,000 seconds, histatin 5 inhibition on CD14 activation flux, and drug simulation for 75,000 seconds.
Dendritic cell response of histatin 5 on HagB. Mononuclear cell-derived human myoid dendritic cells (Lanza Walkersville, Inc., Walkersville, MD and StemCell Technologies, Inc., Vancouver, BC Canada) were used and contained cells staining 31.5% CD1c+ (79.1% CD1c+ and 54.4% CD141*) (79.1% CD1c+) (Flow cytometry kit FMC016, R&D Systems, Inc., Minneapolis, MN). Adhered dendritic cells in 24 well plates (4.1 × 10^5 cells/cm²; Corning, Tewksbury MA) were treated with 0.01 M PBS, pH 7.2 (37 °C, 5% CO²; control); 2.0, or 20.0 μM histatin 5 in LGM-3 (histatin 5 control), or 0.02 μM HagB in LGM-3 (HagB control). Dendritic cells were also treated with 0.02 μM HagB + 2.0 μM histatin 5, previously mixed together at 37 °C for 30 minutes or 0.02 μM HagB + 20.0 μM histatin 5, previously mixed together at 37 °C for 30 minutes. After treatment, cells were incubated at 37 °C in 5% CO². At 0.0, 2.0, 4.0, 8.0, and 16.0 hours, 0.1 ml was removed from each well and 0.1 ml of LGM-3 was added back. Samples were centrifuged at 13,200 g (16,100 rpm) at 25 °C for 5 minutes to pellet non-adherent cells and the supernatants were removed and frozen at −80 °C.

Cytotoxicity. After collection of the last timed sample, LGM-3 was removed and fresh LGM-3 containing Alamar blue (AlamarBlue, Invitrogen Corp., Frederick, MD) was added, and the cells were incubated for 16 hours. The fluorescence intensity of the resorufin was determined using an excitation wavelength of 590 nm (SpectraMax M2 Multi-Mode Microplate Reader, Molecular Devices, LLC, Sunnyvale, CA).

Determination of chemokines and cytokines. Briefly, 25.0 μl of dendritic cell supernatant was added to anti-human multi-cytokine magnetic beads (Milliplex immunoassay, Millipore, Billerica, MA USA) and incubated at 4 °C for 18.0 hours. Unbound material was removed by aspiration (ELX405TS magnetic plate washer, BioTek, Winooski, VT USA); anti-human multi-cytokine biotin reporter was added, and the beads were incubated at room temperature for 1.5 hours. The day Streptavidin–phycoerythrin was then added and the plates were incubated at room temperature for an additional 30 minutes. Stop solution was added, and the plates were read (LumineX model 100 IS, Austin, TX). Standard curves for each cytokine were prepared from 2.3 to 10,000.0 pg/ml of chemokines and cytokines in each sample were interpolated from standard curves (Spong et al. 5,1, Luminex, Austin, TX USA; MILLIPLEX Analyzer v5.1, Millipore, Billerica, MA USA).

Statistical analysis. A log10-transformation was applied to ILS, CSF2/M-CSF, CCL3/MIP-1α, CCL4/MIP-1β, and TNFα concentrations detected in the tissue culture supernatants, as recently described. The log transformation attenuates the positive skew in the distributions of the chemokine concentrations and makes the data fit the log-transformed concentrations. Pairwise group comparisons were conducted using the method of Tukey’s Honest Significant Differences (HSD). A 0.05 level was used to determine statistically significant differences. All analyses were conducted using JMP (Version 10.0, SAS, Cary, NC).

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Author contributions
In this study, we assessed the interaction of histatin 5 with immobilized HagB (SPR spectroscopy-based biosensor, A.M.); determined the histatin 5, HBD1, and HBD3 binding sites on HagB (HagB protein microarray, J.V.H., E.N.R. and K.A.B.); modeled the preliminary histatin 5, HBD1, and HBD3 binding domains on HagB (TASSER imaging, J.Y. and Y.Z.); assessed the histatin 5 attenuation of HagB-induced chemokine and cytokine response in human dendritic cells (Luminex 100 IS, D.S.B., C.L.F., E.N.R., A.P.F., D.D. and K.A.B.); and predicted the pathways or transcription factors altered by histatin 5 (P.R.N., R.V., S.C. and S.V.). J.E.C. oversaw the statistical analysis and D.S.B. and K.A.B. wrote the main manuscript.

Additional information
Supplementary information accompanies this paper at http://www.nature.com/scientificreports

Competing financial interests: D.S.B., J.R.V.H., J.Y., C.L.F., E.N.R., A.P.F., D.D., J.E.C., Y.Z. and K.A.B. declare no competing financial interests in the findings of this study or with Cellworks Group, Inc., Saratoga, CA; Cellworks Research India Pvt Ltd.; or with SensiQ Technologies Inc., 800 Research Parkway, Suite 100, Oklahoma City, OK 73104, USA. A.M. works for SensiQ Technologies Inc., 800 Research Parkway, Suite 100, Oklahoma City, OK 73104, USA. P.R.N., R.V. and S.C. work for Cellworks Research India Pvt Ltd., Bangalore, India and S.V. works for Cellworks Group, Inc., Saratoga, CA.

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