Pulmonary Haptoglobin and CD163 Are Functional Immunoregulatory Elements in the Human Lung

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Abstract
Background: The acute-phase protein haptoglobin (Hp) and its receptor CD163 serve as immunomodulators and possess anti-inflammatory besides antioxidant functions. Objectives: To further understand the role of the recently described pulmonary Hp (pHp) and its receptor CD163 in case of inflammation and infection, pHp and CD163 were investigated on mRNA and protein level to gain insight into the cellular events taking place upon stimulation with the inflammatory mediators LPS, Pam3, cytokine IL-6 and dexamethasone, and upon infection with respiratory pathogens (Haemophilus influenzae, Streptococcus pneumoniae and Chlamydia pneumoniae) by use of a human ex vivo tissue culture model and cell cultures of A549 and alveolar epithelial cells type II. In addition, pHp and CD163 expression in COPD and sarcoidosis was assessed. Methods: We conducted experiments using 942 ex vivo cultured lung samples applying immunohistochemistry, immunocytochemistry, in situ hybridization, immunofluorescence, real-time PCR, RT-PCR, slot and Western immunoblot analyses with tissue lysates and culture supernatants as well as ELISA and cytometric bead array analyses. Results: This study describes for the first time the expression, regulation and secretion of pHp and its receptor CD163 in the human lung. The release of soluble mediators from A549 cell line and human monocyte-derived macrophages was observed indicating that Hp differentially activates the release of soluble mediators and major chemoattractants. Conclusions: The findings indicate a native function of pHp and CD163 as functional pulmonary defense elements due to local expression, regulation and secretion during lung infection and as part of the inflammatory immune response of the respiratory system.

Introduction

During the acute phase of inflammatory processes, elevated expression of acute-phase proteins (APP) is characteristic. One of those molecules is hepatic haptoglobin (Hp) occurring in different phenotypes. Its main role is
capturing hemoglobin to prevent both iron loss and kidney damage during hemolysis [1]. Little is known about the role of APP during local inflammation. In general, they are capable to suppress nonspecific inflammation via suppressed release of pro-inflammatory cytokines in response to inflammatory stimuli; on the other hand, inflammatory cells express receptors for Hp [2, 3]. In human hepatic cells, Hp expression requires glucocorticoids and IL-6 [4], whereas in hepatic rat cells Hp expression requires glucocorticoids as well as both IL-6 and IL-1β [5]. During inflammation, Hp expression and regulation vary in different tissues and species [6, 7]. Recently, we described for the first time a pulmonary Hp molecule (pHp) which is synthesized locally in the human lung [8].

It has been shown that CD163, which was identified as an Hp-hemoglobin (Hp-Hb) scavenging receptor, is synthesized as a 130-kDa glycoprotein belonging to the scavenger receptor cysteine-rich superfamily [1]. It is expressed exclusively in cells of the monocye lineage; high levels of CD163 were detected in a number of different mature tissue macrophages such as alveolar macrophages (AM) [1]. The regulation of CD163 expression by pro- and anti-inflammatory mediators and the secretion of anti-inflammatory cytokines in response to ligand binding to CD163 strongly suggest an immunomodulatory function of CD163. By contrast, TNF-α, IFN-γ and LPS suppress CD163 [9]. Interestingly, IL-6 induces both Hp and CD163 expression [9]. Up to date, the role of Hp synthesis in extrahepatic tissues is poorly understood. There may be more functions of Hp and CD163 in the lung than binding to hemoglobin during hemolysis or signaling related to the inflammatory response.

To address these questions, we investigated expression and regulation of pHp and CD163 mRNA and protein upon stimulation with pHp and CD163 mRNA and protein upon stimulation with Gram-positive and Gram-negative bacteria-simulating, microbial antigens (Pam3 and LPS), inflammatory cytokine IL-6, glucocorticoid (dexamethasone, DEX) and respiratory pathogens (Haemophilus influenzae, Streptococcus pneumoniae and Chlamydia pneumoniae), applying a human ex vivo tissue culture model (short-term stimulation of tissues, STST) in combination with the HEPES-glutamic acid buffer-mediated organic solvent protection effect (HOPE) fixation technique [10]. The HOPE technique is a novel formalin-free fixation procedure for cells and tissues which results in better preservation of nucleic acids and proteins compared to conventional formalin fixation, while the morphology of HOPE-fixed tissues is comparable to formalin-fixed materials. In addition, the release of soluble mediators and chemoattractant cytokines upon Hp co-stimulation was investigated. To further illuminate the role of pHp and CD163 in the context of lung inflammation, tissues from patients with chronic obstructive pulmonary diseases (COPD) and sarcoidosis were included. These tissues were not cultivated and diagnoses were confirmed by two pathologists. A functional role for glycoproteins in the regulation of inflammatory processes had already been shown [11, 12]. The present study holds the hypothesis that synthesis and secretion of high levels of pHp and CD163 during infection of tissues and cell cultures is an integral part of respiratory immune reactions.

Material and Methods

Sample Origin and Collection

Eighty-two human lung tissues were obtained from patients directly after lobectomy/pneumonectomy. Nine hundred forty-two definitely tumor-free lung tissues of up to 1 cm³, resected distant from the tumor, were collected from those patients and subjected to ex vivo tissue cultivation and stimulation including the respective medium-controls. Further, two pulmonary cell types (alveolar epithelial cells type II, AEC II, primary cells, obtained from the University of Freiburg, Department of Pneumology, and A549 cell line obtained from the Medical Clinic Borstel) were included in this study. For additional characterization of pHp and CD163 in chronic diseases, we also analyzed 11 COPD and 11 sarcoidosis cases which were not cultivated in STST (see below). Ten HOPE-derived single sections of lung tissue stimulated with inflammatory stimuli were produced additionally for in situ hybridization to verify immunohistochemistry (IHC) results.

This study was approved by the ethical committee of the University of Lübeck (reference No. 03/158) and is in compliance with the Helsinki declaration.

Short-Term Stimulation of Tissues

We investigated the 942 lung tissue specimens as previously described [13] within the STST model in the following way. For pHp investigations, 14 specimens were stimulated with IL-6 (50 ng/ml; Sigma-Aldrich Chemie GmbH), 80 specimens with LPS (10, 50 and 200 ng/ml; EMC Microcollections GmbH), 57 specimens with Pam3 (200 ng/ml; EMC Microcollections GmbH) and 63 specimens with DEX (10 and 50 ng/ml; Merck) using medium controls (culture medium without stimulus). Ninety-four specimens were pre-incubated for 3 h in medium to wash the tissue from blood and pre-existent pHp which was not synthesized due to applied stimulation.

For investigation of CD163 we stimulated 101 specimens with LPS, 67 specimens with DEX, 75 specimens with Pam3 and 14 specimens with IL-6 accompanying medium controls. Additionally, ex vivo infection was performed by application of H. influenzae, C. pneumoniae and S. pneumoniae into the culture medium (10 specimens each). Culture conditions were set as previously described (37°C and 5% CO₂) applying different concentrations
and incubation periods for each stimulus [14]. Stimulation was performed for 10 min, 30 min, 2 h and 24 h. In each STST experiment two specimens were incubated in parallel dedicated either for HOPE- or formalin fixation.

Several STST culture supernatants were analyzed using slot immunobLOTS (see below) to identify secreted pHp and CD163.

**Tissue Microarrays and Single Sections**

After stimulation all specimens were formalin-fixed and arranged in tissue microarrays (TMAs) as described previously [10]. Sections from these paraffin blocks were mounted on SuperFrost+ slides (Menzel-Gläser). Twenty-one TMAs were produced containing all differently stimulated tissues.

**IHC and Immunocytochemistry**

IHC stainings were performed with formalin-fixed and paraffin-embedded tissue. TMAs or single sections were used as previously described [8, 10]; the primary mouse anti-human antibodies utilized in this study was a monoclonal anti-Hp antibody (clone HG-36, dilution 1:100; Abcam) and a monoclonal mouse anti-human anti-CD163 antibody (clone 10D6, dilution 1:100; Di-HOPE). Immunocytochemistry was carried out with either HOPE- or formalin-fixed cytospins as previously described above. Amplification was performed using a Light Cycler® based system (Roche Molecular Biochemicals) with lightycler software 4 LL.2.0 for data analysis. Specific primers against human Hp were used (QuantiTect primer assay; Qiagen). GAPDH was used as housekeeping gene. Primers were as follows: 5'-GGAGTCGGACCCCATATCAT-3' (forward) and 3'-CTGAGAGTCAAGGCAGCAC-5' (reverse). Relative mRNA quantification was done by using standard curves for Hp and GAPDH. The mRNA level in each sample is described as a relative expression value and is compared to the expression level of infected and stimulated lungs out of STST.

**Collection of Supernatants**

To investigate secretion of pHp and CD163 from ex vivo tissue culture, A549 cell line and AEC II, 2 ml of the culture supernatants were collected and centrifuged at 6,000 rpm for 5 min before storing at −80°C. For pHp investigation 45, 32, 22 and 12 supernatants were analyzed following LPS, DEX, IL-6 and Pam3 stimulation, respectively. For CD163 analyses 16, 12, 12 and 21 supernatants were used, respectively.

Slot and Western immunoblot analyses were conducted subsequently after application of different concentrations of stimuli (10, 50 and 200 ng/ml) and incubation times (10 min, 30 min, 2 h and 24 h); total protein analyses were performed using Bradford Coomassie Protein Assay (Pierce). Optical density was measured at 595 nm using a Helios β-photometer (Unicam).

**Slot Immunoblot Analyses of Culture Supernatants**

Immunoblotting of supernatants from ex vivo tissue culture, A549 cell line and AEC II, was performed with PR 60 and PR 648 slot immunoblot filtration manifolds and a vacuum pump (Amersham Biosciences); 300 µg per slot were applied, the procedure included four main steps starting from setting up the slot immunoblot, applying samples, removing the blot and immunostaining (analogue to SDS-PAGE Western blot staining). For evaluation of band intensity, a software was used (Band Leader).

**Cellular Transcription of pHp within STST Model (in situ Hybridization)**

For confirmation of pHp transcription in human lung tissues, we analyzed 10 IL-6- stimulated specimens and medium controls for comparison by in situ hybridization (ISH) to detect cellular transcripts of the pHp mRNA within the tissues. ISH targeting the 341-bp Hp transcript was carried out using the following Hp-specific primers: forward 5'-AGGCATTATGAAGGCAGGACAC-3', reverse 3'-CTTCAGGGCTGAATCTTGC-5'. Slides were kept in moist chambers overnight at 46°C. Probes were generated and hybridized like previously described [14].

**Conventional RT-PCR**

Total RNA was extracted from 10 human, *H. influenzae*-infected, STST-cultured, HOPE-fixed lung tissues which showed upregulation in IHC stainings (following RNeasy Mini kit; Qiagen). RT-PCR was performed conventionally according to Abdullah et al. [8] using Hp-specific primers: forward 5'-AGGCATTATGAAGGCAGGACAC-3', reverse 3'-CTTCAGGGCTGAATCTTGC-5'; for CD163, specific forward primer 5'-AGACCTGGACTTTAATGC-3', reverse primer 3'-CGTTGTGAGTCAATGCTTAT-5'. In parallel, specific primers for the housekeeping gene glyceraldehyde-3-phosphate-dehydrogenase (GAPDH; forward primer 5'-AGAACGGGAAGCTTGTCATC-3', reverse primer 3'-TGCTGA TGATCTTGAGGCTG-5') were used as controls. For densitometry, Band Leader software was used.

**Real-Time RT-PCR**

RNA isolation and cDNA synthesis were done like mentioned above. Amplification was performed using a Light Cycler® based system (Roche Molecular Biochemicals) with lightycler software 4 LL.2.0 for data analysis. Specific primers against human Hp were used (Quantitect primer assay; Qiagen). GAPDH was used as housekeeping gene. Primers were as follows: 5'-GGAGTCGGACCCCATATCAT-3' (forward) and 3'-CTGAGAGTCAAGGCAGCAC-5' (reverse). Relative mRNA quantification was done by using standard curves for Hp and GAPDH. The mRNA level in each sample is described as a relative expression value and is compared to the expression level of infected and stimulated lungs out of STST.

**Measurement of Chemokine and Soluble Mediators via ELISA and Cytometric Bead Array**

To investigate the role of pHp during the initial local response to the inflammatory stimulus LPS (200 ng/ml), A549 cell’s chemoattractant and monocyte chemoattractant protein-1 (MCP-1) and CXC (IL-8) were measured. A549 cells were cultured up to 30 h in the presence or absence of LPS (200 ng/ml) and Hp protein (150 ng/ml) (lyophilized Hp; Abcam). Supernatants were collected at the end of incubation and stored at −80°C until use. Chemokine levels in supernatants were determined using a commercially available ELISA kit for MCP-1 (R&D Systems GmbH). For IL-8 determination the BioSource CytoSet kit was used. Optical density measurements were performed with a microplate reader model 450 nm and values were analyzed with Microplate Manager/PC Data Analysis software (both from Bio-Rad Laboratories). Human monocyte-derived macrophages were generated and used for cytometric bead array (CBA) analysis as previously described [16]. Supernatants from macrophage cultures were analyzed by CBA (BD Biosciences), as recommended by the manufacturer.

**SDS-PAGE and Western Blotting**

We performed Western blotting using STST-cultured, HOPE-fixed material. Supernatants of cell cultures from the A549 cell line as well as from bronchoalveolar lavage (BAL) stimulated with inflammatory stimuli were additionally blotted. Dепаразиниза-
tion, rehydration, extraction and immunostaining were performed as described previously [8]. Protein concentrations were determined in extraction buffer using a spectrophotometer with Bradford Coomassie Protein assay kit (Pierce); 10–15 µg of protein were used, mixed with sample buffer (200 mM Tris-HCl pH 8, 20 mM EDTA, 25% glycerol, 2% SDS and 0.02% bromophenol blue) either containing 1% DTT as reducing agent or without DTT for nonreducing conditions. The percentages of supernatants showing a clear induction of the secretion compared to medium controls are displayed. χ² tests were used for statistical analyses (* p < 0.05; ** p < 0.01; *** p < 0.001).

**Cell Culture and Cytospin Preparation**

Cytospin preparations were prepared as previously described [15], performing HOPE and formalin fixation in parallel for immunocytochemistry and immunofluorescence stainings (described below), Human A549 cells (American Type Culture Collection) and primary AEC II were maintained in modified high glucose DMEM (4.5 g/l) medium with L-glutamine (PPA) supplemented with 1% of penicillin/streptomycin solution (Gibco/Invitrogen) and 10% heat-inactivated FCS (PPA). BAL cells/fluuid was further cultured in RPMI 1640 with L-glutamine (PPA) applying inflammatory agents, inflammatory cytokines and DEX. After stimulation, cells were washed 3 times in PBS and harvested after 30 h. Finally, cells were resuspended in PBS reaching a concentration of 2 × 10⁶/ml. Around 50,000 cells were attached to SuperFrost+ microscope slides (Menzel-Gläser) by centrifugation for 5 min at 450 rpm using high acceleration in a Cytospin 2 centrifuge (Shandon). Afterwards, cells were dried for 10 min at 37°C. After overnight fixation at 4°C in HOPE solution, cells were incubated with acetone/glyoxal for 1 h, and dehydrated 6 times for 10 min with acetone (all at 4°C), followed by two incubation periods in isopropanol (10 min at 60°C, 2 min at 60°C) and subsequent air-drying. Rehydration was achieved by incubation in 70% (v/v) acetone for 10 min at 4°C and DEPC water incubation for 10 min (also 4°C). Slides were then air-dried again. For comparative studies, dried cytospins were fixed in 4% formalin for 10 min at 4°C.

**Immunofluorescence Staining**

Immunofluorescence stainings of cytospin preparations from BAL (formalin- and HOPE-fixed) were carried out using the same primary antibodies as in the previously described IHC procedure [10]. As a secondary antibody, we applied goat anti-mouse fluorescein isothiocyanate (FITC, Alexa Fluor 488; Invitrogen) in a dilution of 1:200 (45 min in the dark) according to the instructions of the manufacturer. Counterstaining was performed using DAPI (Vector Laboratories) for examination with a fluorescence microscope (Nikon Eclipse 80i); slides were stored at 4°C in the dark.

**Statistics**

We have included standard errors of the means and p values from χ² tests (fig. 1, 4, 6, 10).

**Results**

**pHp Is Expressed and Regulated by Microbial Antigens, IL-6 and Glucocorticoids in the Human Lung**

The numbers of investigated cases (n) and the percentages of cases which showed upregulation of pHp upon stimulation with different stimuli are shown in figure 1a. All stimuli caused upregulation of pHp in different intensities and were determined for LPS in 67/80 cases (83.7%), for DEX in 54/63 (85.7%), for IL-6 in 12/14 (85.7%) and for Pam3 in 41/57 cases (71.9%). The baseline of pHp expression in a medium control is displayed in figure 2e, as well as examples of various other stimulation experiments.
The main sites of protein expression were observed in AEC II and AM. Dose- and time-dependent induction of pHp was observed with DEX and LPS. Pam3 and IL-6 were tested each in one concentration only and induced pHp time dependently. Generally, a granular staining of pHp inside the cytoplasm was observed (fig. 2a–e).

**CD163 Is Differently Regulated in Human Lung Cells by Microbial Antigens, IL-6 and Glucocorticoids**

Figure 1b shows the numbers of investigated cases and the related percentages of cases showing an induction of CD163 being differently regulated upon different stimuli in STST. Increased expression of CD163 was caused by DEX, IL-6 and Pam3 in 77.6% (52/67), 78.5% (11/14) and 46.6% (35/75) of cases. LPS downregulated receptor expression in 47.5% of tissues (48/101; fig. 1b). All effects except of Pam3 and IL-6 (only one concentration used) were dose and time dependent. Signals of CD163 were exclusively localized in the cell membranes of AM.

**pHp and CD163 Get Actively Secreted from Lung Tissues upon Stimulation and Enhance the Secretion of Soluble Mediators**

Figure 1c, d shows the total number of investigated cases and in how many cases (in percent) secreted pHp and CD163 could be detected in culture mediums derived from STST. After 10 min of STST, activation of pHp protein secretion occurs compared to the medium control (pre-incubated in medium for 3 h; fig. 3a). pHp signaling on slot immunoblots after 24 h of incubation showed intense bands of pHp caused by LPS in 31/45 supernatants (68.8%), by DEX in 28/32 (87.5%), by IL-6 in 7/12 (58.3%) and by Pam3 in 11/22 supernatants (50%). Strong signals appeared related to specimens which were not incubated in medium before stimulation (fig. 3b). An increase in signal intensity correlates with the length of the respective incubation period. Secretion of CD163 was induced by application of LPS in 10/16 (62.5%), of DEX in 10/12 (83.3%), of IL-6 in 10/12 (83.3%) and of Pam3 in 7/21 (33.3%) cases (fig. 1d). As expected, no secretion was observed in AEC II and A549 culture supernatants as CD163 is associated only to AM.

**Hp Induces the Release of Immunomodulatory Molecules in Lung Cells**

Supernatants analyzed from A549 cell line showed that application of pure Hp protein and LPS as well as co-stimulation with Hp and LPS enhances the release of the soluble chemoattractants IL-8 and MCP-1 after 30 h of stimulation. LPS and Hp alone increase concentrations of...
these chemoattractants dose dependently. Costimulation with LPS and Hp shows a comparably small effect with 10 ng/ml LPS and 10 μg/ml Hp, while stimulation with high doses of both (200 ng/ml LPS and 150 μg/ml Hp) result in extremely elevated IL-8 and MCP-1 concentrations (fig. 4a, b).

In human monocyte-derived macrophages, treatment with 10 μg/ml pure Hp protein for 24 h led to the release of TNF-α, IL-6 and MCP-1. In contrast, IL-12p40 and RANTES formation were only detectable in response to LPS but not in response to Hp protein (fig. 4c). In addi-
tional in vitro co-stimulation macrophages were treated with Hp in the presence of LPS. However, the addition of Hp did not modulate the macrophage response towards these macrophage-activating stimuli (data not shown).

**Cellular Transcription of pHp**

By application of ISH, pHp transcripts were found in AM and AEC II of the lung tissues, negative controls did not show any signals (fig. 5a–c). This is in accordance with the cell types found to be expressing pHp protein by IHC.

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Transcriptional Regulation of pHp and CD163 in Response to Different Stimuli

In real-time PCR upregulation of pHp mRNA by DEX, LPS, Pam3 and IL-6 was determined; hereby only DEX affected tissues in a dose- and time-dependent manner (fig. 6a). Concerning LPS, the amount of pHp mRNA was recorded after stimulation with 200 ng/ml for 24 h and showed the highest increase in pHp mRNA. Furthermore, CD163 mRNA expression increases dose dependently according to RT-PCR data in DEX-stimulated lungs (fig. 7a). No variation was observed after Pam3 application; LPS caused downregulation of CD163 which verifies the IHC results (data not shown).

Transcriptional Regulation of pHp and CD163 in Response to Infection and Chronic Diseases

RT-PCR showed equal levels of pHp mRNA in all cases. CD163 was significantly downregulated by *H. influenzae* (fig. 7b). The same is true in real-time PCR, which again shows a downregulation of CD 163 mRNA due to *H. influenzae* and *C. pneumoniae* (fig. 6b). A significant upregulation of pHp was observed in the chronic diseases COPD and sarcoidosis (fig. 6c).

**pHp and CD163 Are Induced by Experimental Respiratory Infection**

IHC results revealed elevated expression of pHp and downregulation of CD163 expression in 11 specimens after 24-hour infection with *H. influenzae*. Expression of pHp was more frequent in macrophages than AEC II. *S. pneumoniae* caused comparable elevated expression of pHp and CD163 in AEC II and AM (all 11 specimens). Strong upregulation of pHp and its receptor upon *C. pneumoniae* infection was observed after 24 h, again in AEC II and AM (fig. 8a–d).

Patterns of pHp and CD163 Expression in Inflammatory Lung Diseases

For comparison of pHp and CD163 expression in COPD and sarcoidosis, 11 cases were analyzed by IHC. Areas around inflamed airways showed strong pHp and CD163 expression in AM and AEC II from COPD patients (fig. 9a, b). By contrast, weak expression of pHp and extremely high expression levels of CD163 from cells forming granuloma lesions of sarcoidosis have been observed (fig. 9c, d).

Western Blot Analysis of Stimulated Tissues Targeting pHp and CD163

Western blotting of lysates from STST-cultured tissues and the corresponding supernatants from tissue cultures showed comparable amounts of pHp protein concerning the different stimuli. Bands showed a molecular weight around 160 kDa (representing Hp 2-2 according to non-reducing conditions and as described in the data sheet; data not shown). In cases of BAL supernatant signaling could be generated only after 35 h stimulation but not after 24 h. Furthermore, Western blotting targeting CD163 in supernatants of BAL culture and cell lysates revealed upregulation due to DEX and IL-6 stimulation and down-regulation with LPS (data not shown).

**pHp Stainings of Cytospin Preparations Prepared from Primary AEC II**

After stimulation with 200 ng/ml LPS, AEC II showed time-dependent expression of pHp. Equal results were
found after stimulation with DEX and Pam3 (not shown). Figure 10 shows the medium control (fig. 10a) as well as pHp expression after 24 and 35 h (fig. 10b, c). Figure 10d shows a statistic evaluation of the time-dependent induction of pHp by LPS.

Immunofluorescence Microscopy Reveals Secretion of pHp via Vesicles due to Stimulation with LPS, DEX and IL-6 in BAL and AEC II Cytospins

Immunofluorescence microscopy of BAL cytospins showed secretion of pHp, the targets were located in vesicles (fig. 11). Further, the immunocytochemistry-positive cytospins prepared from primary AEC II showed pHp induction by DEX, LPS and IL-6 (data not shown, Pam3 was not tested). Expression and secretion in vesicles were observed and clearer in HOPE material compared to formalin; signals were more present after 35 h compared to 24 h of stimulation.

Discussion

The environment of the lung generally needs strong anti-inflammatory properties due to exposure to the atmosphere and a large number of potentially incoming chemical agents, microorganisms, organic and inorganic material or other antigens. Therefore, there have to be effective intra- and extracellular antioxidant defense and other protection mechanisms [17]. If the fast synthesis of gene products related to those purposes is not ensured, lung tissues underlie permanent stress and inflammation. In cases of acute pathogen contact the expression of acute-phase proteins is immediate and local expression becomes inevitable. Recently, pHp has been identified as a part of the lung’s local, acute protection system being synthesized locally in lung adenocarcinomas and in the
tissue surrounding squamous cell carcinomas. pHp expression could further be shown in AM and AEC II of tumor-free lung tissue [8].

The enhanced production of APPs in response to inflammatory stimuli suggests that these proteins might augment the acute inflammatory response.

**Regulation of pHp**

In our study, induction of pHp was shown with inflammatory agents LPS, IL-6 Pam3 and DEX on protein and mRNA level. The results have been verified by data from stimulation-experiments with A549 cells and primary AEC II (IHC).

The induction is caused by many factors and the regulation of Hp is varying among species and cell types. IL-6, IL-1, TNF-α and glucocorticoids are capable of upregulating the synthesis of Hp in rat and bovine hepatic cell lines [18].

The activity of DEX on pHp and CD163 is of pharmacologic relevance since DEX is a frequently applied drug in pneumology. In humans, Hp is mainly regulated by IL-6 and DEX which is in agreement with results of this study.

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**Fig. 8.** pHp synthesis is induced by microbial infection with *H. influenzae*, *C. pneumoniae* and *S. pneumoniae* within ex vivo tissue culture after 24 h of incubation. Here, HOPE technique was applied: medium control without infection (**a**), infection with *H. influenzae* (**b**), *C. pneumoniae* (**c**) and *S. pneumoniae* (**d**). Staining was observed in AM and AEC II. All images 200×.

**Fig. 9.** Detection of pHp and CD163 in chronic lung diseases using IHC. **a**, **b** COPD, **c**, **d** Sarcoidosis. Concerning COPD, pHp expression was shown within obstructive bronchitis (upper arrow in **a**) and AM surrounding the obstructive bronchitis (lower arrow in **a**) while highly elevated CD163 expression from AM (upper arrow in **b**) was observed. Lower arrow (**b**) points on the negative obstructive bronchitis for CD163. In case of sarcoidosis, pHp is expressed weakly in granuloma cells as pointed by lower arrow (**c**) and most of AM and AEC II cells surrounding the granulomatous lesions express pHp (upper arrows in **c**). CD163 is highly over expressed from cells which form the granulomatous lesions (**d**). **a**, **b**, **d** 200×. **c** 400×.
At the first view it seems curious that pHp is upregulated and released upon apparently 'controversial' stimuli, which moreover takes place in the same time frames. We conclude that pHp release is not only involved in anti-inflammatory effects of glucocorticoids, but also helps to limit the inflammatory reactions upon bacterial infection. Therefore, we speculate an additional role of pHp in 'keeping chronic infectious processes chronic' which is supported by the data obtained from COPD and sarcoidosis.

IL-6 is a well-known inflammation-related cytokine, trigger of the acute-phase response and mediator for Hp gene regulation in all species studied so far [19, 20]. Hp increases approximately 3- to 6-fold in the plasma of mice receiving LPS injections; the regulation of Hp response to inflammation has been carefully studied in vitro using hepatoma cells which depend on an interaction between specific DNA sequences and a nuclear binding protein (C/EBP-β) for IL-6 serving as transcription factor for Hp (a sequence in the 5'-regulatory region of the human Hp gene) [5].

We have shown that exogenous pHp is stored in monocytes and macrophages within the cytoplasm (granular staining) and other groups reported that it is further secreted by these cells during phagocytosis of Candida albicans [21]. This indicates that pHp protein levels enhance locally at the sites of inflammation or due to induction by stimuli to modulate granulocyte activity as we have shown in supernatants of ex vivo tissue cultures. Furthermore, specific binding of Hp to neutrophils leads to an inhibition of respiratory burst activity [3].
In addition, it has been reported that Hp suppresses macrophage function such as LPS-induced production of TNF-α, proliferation and cytokine production by T cells, and proliferation of B cells. This study shows that there are more differential functions of pHp to activate monocytes and macrophages and in cell lines as shown in stimulated human derived macrophages. This is in agreement with Arredouani et al. [22]. Hp has further been shown to bind to mono- and polynuclear cells through CD11/CD18, CD22 and other yet unidentified surface antigens. Therefore, due to the wide distribution of receptors, Hp modulates functions of these cells and causes endocytosis of Hp with exerting its immunomodulatory and anti-inflammatory effects through receptor-mediated signaling [2, 22].

**Regulation of CD163**

This study also describes an induction of CD163 by DEX and IL-6; on the other side, CD163 gets downregulated in AM due to LPS stimulation. This is in agreement with other studies dealing with CD163 in cell lines [9]. The upregulation of CD163 observed in this study is also in agreement with Ritter et al. [23] who showed the presence of multiple glucocorticoid response elements in the promoter region of the human CD163 gene and by describing several consensus binding sites in the sequence. This might be important for the strong expression of CD163 mRNA during IL-10 and IL-6 treatment [11, 23]. In contrast to anti-inflammatory stimuli, pro-inflammatory substances decrease CD163 mRNA levels. LPS represents a potent inflammatory stimulus and has been shown to mediate some of its action by members of the Rel-family transcription factors, although there are no consensus sequences for Rel located in the proximal promoter region of CD163 [24].

Recently, CD163 was identified as an Hp-Hb scavenger receptor [1]. Therefore, increased synthesis of pHp and CD163 by AM as reaction on inflammatory agents and DEX could contribute to the removal of Hb and thus protect the lower respiratory tract against Hb-related oxidative damage and inflammation.

Interestingly, CD163 mRNA and protein level show downregulation upon LPS stimulation but high protein levels in supernatants. This effect can be explained by LPS-caused shedding of CD163 from the surface of macrophages (known as soluble form of hemoglobin scavenger receptor sHbSR) [25, 26]. Upregulation caused by DEX and IL-6 highly increased the levels of CD163; this was observed in patients with myelo-monocytic leukemia and infection [25]. An elevated expression of pHp and CD163 from a local source, and, additionally, systemic Hp at the site of inflammation or due to induction via inflammatory agents seems to be a scavenging pathway for fast elimination of pro-inflammatory oxidative agents and for the modulation of granulocyte activity.

**Fast Secretion of pHp as a Regulatory Strategy**

Transcription data of pHp mRNA and IHC data differ slightly for low-dose LPS (10 and 50 ng/ml) stimulation because time and dose dependence could only be observed in IHC. By contrast to LPS, DEX possesses specific responsive elements in the promoter region of the human Hp and CD163 genes [11, 19]. Elevated transcription of pHp was recorded during high-dosed LPS stimulation (200 ng/ml) after 24 h, which is in agreement with the results of IHC and slot immunoblot analyses showing fast secretion in a dose- and time-dependent manner. But this might be influenced by secretion of pre-existing pHp [21], so tissues were washed from pre-existing pHp by pre-incubation in medium to prove pHp synthesis definitely due to stimulation. We found that after 10 min of stimulation secretion starts and increases up to 24 h.

**Secreted pHp Induces Immunomodulatory Molecules**

The significant role of Hp for the secretion of chemoattractants and soluble mediators from the A549 cell line and human monocyte-derived macrophages indicates an important involvement in chemoattractant effects of human lungs and differential activation of the release of pro-inflammatory mediators by human macrophages. This is in agreement with studies which observed that pro-inflammatory cytokines such as TNF-α induce AEC II to produce chemoattractants that could provoke the transendothelial migration of CD3+ lymphocytes and CD14+ monocytes [27]. Recent studies also suggest that the alveolar epithelium plays a role in modulating immune responses; the immortalized lung epithelial cell line A549 can produce MCP-1, RANTES and IL-8 following stimulation with TNF-α and other pro-inflammatory cytokines [28, 29]. Rat type II pneumocytes produce MCP-1 in response to IL-1 [29] and can be stimulated to produce neutrophil chemoattractants [30]. Our results prove for the first time that pHp in human AEC II contributes to the development of local inflammatory responses by production of the major chemoattractant cytokines MCP-1 and IL-8 which are responsible for attracting monocytes, lymphocytes and polymorphonuclear cells and soluble mediators. The potentiation of the effects obtained with high doses of Hp+LPS seems to be a possible way for immediate reaction upon acute and massive inflammation.
pHp and CD163 in COPD and Sarcoidosis

This study additionally suggests a role of pHp and CD163 in chronic diseases (such as COPD and sarcoidosis) where strong expression has been observed. Indications summarized lead to speculations that there are differential histological changes in lesions between two chronic cases as well as due to increased levels of chemokines which are responsible for the recruitment of monocytes and for forming chronic inflammation. This is characterized by continuous recruitment and activation of immune cells such as monocytes in response to a persistent stimulus and by keeping the disease in chronic condition because both Hp and CD163 are known to serve as immunomodulators during inflammation with elevated levels in the healing phase of acute inflammatory reactions, in chronic inflammatory diseases and during wound healing processes [11, 12].

pHp and CD163 in Pulmonary Infections

Besides antibacterial effects through depleting hemoglobin, which is a main source for bacteria requirements as found in recent studies, deletion of a set of three Hp-Hb binding protein genes from a nontypeable strain (86-028NP) of *H. influenzae* attenuated virulence in the chinchilla otitis media model of noninvasive disease [31]. CD163 has been shown to bind both Gram-positive and Gram-negative bacteria and is activated through TLRs. Furthermore, regulation of CD163 during an innate immune response implies an important role for this molecule in host defense against infection [26, 32]. We found pHp and CD163 downregulated in lung tissues upon *H. influenzae* infection showing decreased mRNA synthesis; the observed high-protein levels might be explained by accumulation of pHp protein in cellular vesicles. We provide evidence that local pHp synthesis and secretion are increased after inflammatory induction in STST and alveolar cell lines. The secretion of pHp by cells is a comparably fast mechanism since pHp was measured in the supernatants already after 10 min of stimulation, which suggests a high degree of independence from other pathways, for example that of TLRs. Due to the comparably large amounts of quickly secreted pHp it should be considered that a substantial amount of immunoregulatory events in the human lung are taking place in a pHp-enriched environment.

Further studies will focus on the interaction of CD163 with bacterial ligands, intracellular signaling pathways and actual roles of CD163 and pHp during bacterial infection in vivo and by use of human materials and mouse models (knockout experiments). Furthermore, experiments with knockdown or inhibition of pHp and CD163 in cell systems are underway to further enlighten the role of this complex in the human lung.

Finally, the results of this study strongly suggest a role of pHp in antioxidant, inflammation-influencing and immunomodulatory control mechanisms of the human lung. The complete integration of these new results into the complex immunoregulatory network of the lung can contribute to the development of novel treatment regimens facing acute or chronic infections. Finally, these findings confirm a function of pHp and its receptor as native local pulmonary defense elements with influence beyond the yet known immunomodulatory and antioxidant properties directly within the respiratory system.

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References
