Obesity, leptin and host defence of *Streptococcus pneumoniae*: the case for more human research

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Shareable abstract (@ERSpublications)
Use of human challenge models could enable a better understanding of the interactions between *S. pneumoniae* exposure, colonisation and subsequent mucosal and systemic immunity in humans with obesity. https://bit.ly/3RpMIfd

Cite this article as: Hales C, Burnet L, Coombs M, et al. Obesity, leptin and host defence of *Streptococcus pneumoniae*: the case for more human research. *Eur Respir Rev* 2022; 31: 220055 [DOI: 10.1183/16000617.0055-2022].

Abstract

Pneumococcal pneumonia is the leading cause of community-acquired pneumonia. Obesity is a risk factor for pneumonia. Host factors play a critical role in susceptibility to pulmonary pathogens and outcome from pulmonary infections. Obesity impairs innate and adaptive immune responses, important in the host defence against pneumococcal disease. One area of emerging interest in understanding the complex relationship between obesity and pulmonary infections is the role of the hormone leptin. There is a substantive evidence base supporting the associations between obesity, leptin, pulmonary infections and host defence mechanisms. Despite this, there is a paucity of research that specifically focuses on *S. pneumoniae* infections, which are the leading cause of community-acquired pneumonia hospitalisations and mortality worldwide. Much of the evidence examining the role of leptin in relation to *S. pneumoniae* infections has used genetically mutated mice. The purpose of this mini review is to explore the role leptin plays in the host defence of *S. pneumoniae* in subjects with obesity and posit an argument for the need for more human research.

Introduction

The global prevalence of obesity is increasing, yet little is known about the impact of obesity on bacterial pneumonia [1]. Community-acquired pneumonia (CAP) remains the leading cause of hospitalisation and mortality worldwide with *Streptococcus pneumoniae* the most prevalent pathogen [2]. Obesity is considered a risk factor for pneumonia due to impaired B- and T-cell-mediated immune responses that are important in the host defence against pneumococcal disease [3]. Additionally, the association between obesity and other major chronic conditions, such as diabetes and cardiovascular disease, increases the risk of CAP [3]. These host factors play a critical role in determining both the susceptibility to viral and bacterial pathogens and outcome from pulmonary infections.

Obesity has been frequently reported as a risk factor for adverse outcome from 2009 H1N1 influenza [4, 5] and severe acute respiratory syndrome coronavirus 2 [6] but, paradoxically, improved outcome in acute bacterial pneumonia [7–10], with studies on critically ill obese patients reporting no difference or reduced mortality compared to nonobese patients [11, 12]. Similarly, outcome data on the association between obesity and the risk and severity of pneumonia from bacterial infections is inconsistent; obesity has been linked to an increased risk of hospitalisation [13] and alternatively as acting as a protective factor during infections [2, 7]. One reason for this variability is the lack of data on the actual cause of CAP, which is not usually determined in clinical studies [2, 7, 13]. Determining the pathogen in CAP is important to advancing understanding of host defence mechanisms on patient outcomes. A bacterial infection that is highly dependent on neutrophil-mediated clearance may have a poor outcome in patients with obesity.
compared to normal weight individuals. Another important consideration is whether or not bacterial pneumonia was preceded by a viral infection. Studies in human [14] and murine models [15] have shown that the precedence of pathogen exposure may determine pneumococcal infection outcomes and disease severity; pneumococcal infection after influenza may exacerbate infection, whilst pneumococcal infection prior to influenza may reduce mortality in mice [15].

One area of emerging interest in understanding the complex relationship between obesity and pulmonary infections is the role of the hormone leptin [16] – a protein produced primarily by adipocytes [17] that has cytokine-like properties with similar structure and function to cytokines and cytokine receptors of the interleukin (IL)-6 superfamily [18]. As well as regulating appetite and energy expenditure, leptin has been shown to play multiple roles in innate and adaptive immunity in mice and humans [19], and more specifically in pulmonary infections [20].

Whilst there is a substantive evidence base supporting associations between obesity, leptin, pulmonary infections and host defence mechanisms, there is a paucity of research that specifically focuses on S. pneumoniae infections. There is emerging evidence of the role of leptin in mice infected with S. pneumoniae, but there remains limited understanding of the role leptin plays in the susceptibility of obese humans to pneumococcal disease and nasopharyngeal colonisation. This mini review explores the current evidence of the role of leptin in the host defence of S. pneumoniae in obese subjects.

**Leptin signalling and immunological functions**

Leptin is an adipocyte derived hormone, with cytokine like properties, whose function is mediated by binding to a leptin receptor encoded by the LEPR gene. Whilst there are six different isoforms of the leptin receptor, only the long isoform (LepRb) can completely transduce leptin signalling [21]. LepRb is predominantly expressed in the hypothalamus and is present in all types of immune cells, involved in both innate and adaptive immunity [22].

Most of the biological functions of leptin occur via the Janus kinase 2–signal transducer and activator of transcription 3 (JAK2-STAT3) pathway (figure 1). Leptin binds to the long isoform LepRb which causes dimerisation and stimulates JAK2 autophosphorylation as well as phosphorylation of tyrosine residues (Tyr974, Tyr985, Tyr1077, Tyr1138) within the receptor. Leptin receptor–phosphorylated tyrosine 1138 mediates the interaction with STAT3 which dimerise and translocate to the nucleus to activate gene transcription. Suppressor of cytokine signalling 3 (SOCS3) acts as a negative feedback signalling mechanism during prolonged stimulation of LepRb [23, 24]. Leptin induces the activation of the mitogen-activated protein kinase pathway through activation of extracellular signal-regulated kinase 1/2. This pathway is activated following JAK2 activation where Src homology-2 domain-containing protein tyrosine phosphatase-2 recruits growth factor receptor-bound protein 2 leading to activation of the signalling cascade (figure 1). Leptin further induces the phosphatidylinositol-3 kinase/protein kinase B (PI3K/Akt) pathway through insulin receptor substrate 1 phosphorylation following initial JAK2 activation [21, 25] (figure 1). Both these pathways result in cell proliferation and survival.

Almost all immune cells express leptin receptors. Leptin regulates both innate and adaptive immunity through the modulation of immune cell survival and proliferation. In innate immunity, leptin increases the cytotoxicity of natural killer cells and promotes the activation of granulocytes, macrophages and dendrite cells. This activation leads to the production of pro-inflammatory cytokines (e.g. tumour necrosis factor-α (TNF-α), IL-6, IL-12) which facilitate the shifting of T-cells towards type 1 T-helper cell (Th1) priming [22, 25, 26]. In adaptive immunity, leptin increases the proliferation and maturation of naïve T- and B-cells whilst decreasing the inhibitory effects of regulatory T-cells on the immune response [22, 25]. Leptin promotes pro-inflammatory Th1 cytokines rather than anti-inflammatory Th2 cytokines and facilitates Th17 responses. Lastly, leptin regulates B-cell development and activates B-cells to secrete cytokines [25].

**Leptin deficiency, starvation and host defence**

Much of the evidence examining the role of leptin in relation to pneumococcal infections has used genetically mutated nonobese mice with leptin (ob/ob), leptin receptor (db/db) or leptin signalling (s/s, LysM-LepRb-KO and CPE–/–) deficiencies. Studies using leptin-deficient (ob/ob) mice have consistently shown reduced pulmonary clearance following S. pneumoniae challenge when compared to wild-type mice [27, 28]. Administration of exogenous leptin has improved survival, bacterial clearance and reduced bacteraemia [27]. Following infection with lower doses of S. pneumoniae, ob/ob mice exhibited lower pulmonary levels of pro-inflammatory cytokines such as TNF-α and chemokines. Leptin deficiency has not been shown to affect bacterial growth in the lungs [28]. It is further recognised that leptin receptor deficiencies (db/db) and subsequent signalling (s/s) impairment affects pulmonary host defence in

https://doi.org/10.1183/16000617.0055-2022
mice [29, 30]. Disruption to different signalling pathways produces opposing effects on the host defence. Disruption to leptin receptor (LepRb) JAK2/STAT3 signalling enhanced leukotriene production and pulmonary bacterial clearance [29]. Whereas ablation of the leptin receptor (LepRb) in myeloid cells impaired *S. pneumoniae* pulmonary clearance and alveolar macrophage bactericidal function [30].
How this murine work applies to the regulation of bacterial pneumonia inflammation in the human lung needs greater consideration.

A further area of study has been the application of acute starvation models to understand how the metabolic regulatory role of leptin influences the host defence during pneumococcal infection [31]. In studies of nonobese mice, acute starvation during S. pneumoniae infection rapidly decreased leptin levels impairing host defence with associated reduced bronchoalveolar lavage fluid (BALF) neutrophil counts and levels of IL-6 and macrophage inflammatory protein 2. Leptin administration restored bacterial clearance in the lungs and increased levels of BALF neutrophils, cytokines, alveolar macrophages and leukotriene B4 synthesis.

The studies described above focus on leptin deficiency, receptor and signalling defects or conditions of starvation and not obesity per se. These distinctions are important because the pathogenesis of obesity directly influences our understanding of the role of leptin in pneumococcal disease: genetic mutations that cause obesity [27–30] and diet-induced obesity [1, 32], which is the main cause of obesity in humans. New evidence suggests that prolonged overnutrition leading to diet-induced obesity, hyperleptinaemia and leptin resistance, a hallmark of obesity, play a critical role in the immune response to bacterial pathogens [1, 20, 33].

**Overnutrition, hyperleptinaemia and host defence**

In diet-induced obesity, subjects have sustained higher serum leptin levels (hyperleptinaemia) and leptin resistance from prolonged high fat diet overnutrition than lean humans [34]. One study [1] examining the effect of obesity on the host defence against S. pneumoniae noted that CPEΔcarb-mice (carboxypeptidase E enzyme deficient mice; enzyme critical to processing prohormones and proneuropeptide regulation of appetite and energy expenditure) developed hyperglycaemia, raised serum leptin and triglyceride levels, and increased blood neutrophil count prior to S. pneumoniae infection. These findings, in conjunction with emerging evidence of hospitalised patients with severe pneumonia, suggest that it is not body weight per se but chronic hyperleptinaemia that is the link between obesity and increased risk of pulmonary infection [20].

Therefore, leptin resistance and the impact of altered inflammatory signalling in diet-induced obesity [35] need further consideration. Mutation in the leptin gene or the leptin receptor as described in the murine studies above are not the main causes of leptin resistance [35]. Instead, leptin resistance is induced by altered leptin transportation across the blood–brain barrier, deterioration in function of the leptin receptors accompanied by hypothalamic inflammation, endoplasmic reticulum stress and defective autophagy [34, 36, 37]. In high fat diet fed mice, leptin transport across the blood–brain barrier is substantially decreased [38]. In obese humans with severe hyperleptinaemia, leptin levels in the cerebrospinal fluid are only slightly increased [36]. Therefore, it is possible that pulmonary host defence may be impaired differently, in various models of obesity, such as diet-induced obesity with or without leptin resistance.

**Clinical studies**

There are very few clinical studies examining the relationship between leptin levels in obesity and S. pneumoniae infections in humans. Disz et al. [32] specifically examined the relationship between leptin and the outcomes of hospitalised patients with confirmed diagnoses of CAP. This study showed no difference in the serum leptin levels between hospitalised CAP patients and the healthy control group, when adjusting for body mass index. At the time of admission, approximately 5 days from symptom onset, a linear relationship between leptin levels and body mass index, body fat and muscle mass was observed suggesting that leptin acts as a nutritional marker, rather than an inflammatory reactant. However, the role of leptin in the host defence of pneumococcal disease and pneumococcal nasopharyngeal colonisation, in the absence of infection-induced starvation, remains unknown.

Determining the susceptibility or risk of infection in relation to obesity requires carefully controlled human infection studies that investigate pre and post exposure to S. pneumoniae. Similarly, the timing of infection in regards to leptin levels is critical to understand the severity of illness. In mice, leptin levels rise immediately after infection and decline after 24 h [31]. In humans, the time between infection and hospital admission is likely to be greater than 24 h and vary considerably between patients.

Often clinical studies involve obese patients who have multiple comorbidities, known to increase susceptibility to pneumococcal disease, thus making direct links with obesity difficult. Clinical data supports the notation that type 2 diabetes is a risk factor for pneumococcal and other types of CAP [39, 40]. Since approximately 90% of patients with type 2 diabetes are overweight or obese [41], it is difficult to determine if obesity or diabetes is responsible for impaired host defence in humans. Research
that includes normal weight patients with diabetes is needed to address this issue. Likewise, the question of
greater rates of colonisation in humans with obesity requires carefully controlled human infection studies.
Patients with and without type 2 diabetes would need to be included in these studies since this is an
important distinction that will affect understandings of host defence.

Body mass index measurements are frequently used in clinical studies as a proxy for adiposity despite not
directly measuring fat mass. Given that leptin is primarily produced in adipocyte cells, collecting data on
waist circumference, bioelectric impedance and dual energy X-ray absorptiometry scans will provide more
accurate data. Furthermore, 10–30% of people with obesity have normal metabolic parameters
(metabolically healthy obesity) [42]. Recording metabolic syndrome or measures of metabolic status will
provide a more accurate assessment of metabolic health and assist in the distinction between the host
defence mechanisms resulting from obesity and diabetes.

Conclusion

Collectively, the evidence suggests that there are both metabolic and immune responses that influence the
role of leptin in the host defence of \textit{S. pneumoniae} in obese subjects; the effects of these responses being
intricately related to the pathogenesis of obesity (genes or diet), the acute phase of illness, nutritional
status, and the severity of the disease. Furthermore, the evidence suggests that not only leptin deficiency
but prolonged hyperleptinaemia and leptin resistance may impair the pulmonary host defence as observed
in diet-induced obesity. As most of our understanding about the impact of obesity on the host defence to
pneumococcal disease comes from research in genetically mutated mice models, direct comparisons of
murine models with diet-induced obesity in humans may not be feasible. There is a paucity of clinical
research specifically examining pneumococcal nasopharyngeal colonisation, the prerequisite for
pneumococcal infection, in obesity. Further research is required to understand how \textit{S. pneumoniae}
colonisation is regulated in obese populations and informs our understanding of susceptibility to
subsequent pneumococcal disease; and how hyperleptinaemia in diet-induced obesity influences
nasopharyngeal colonisation and lung bacterial burden. The distinction between the influence of obesity
and type 2 diabetes on host defence to \textit{S. pneumoniae} exposure, colonisation and pneumococcal disease
needs further exploration. The use of human challenge models [43–45] in people with an extreme body
mass index could enable a better understanding of the interactions between \textit{S. pneumoniae} exposure,
colonisation and subsequent mucosal and systemic immunity in humans with obesity.

Provenance: Submitted article, peer reviewed.

Author contributions: C. Hales: conceptualisation; data curation; formal analysis; funding acquisition; methodology; project administration; writing – original draft; writing – review and editing; L. Burnet: data curation; formal analysis; writing – original draft; writing – review and editing; M. Coombs: data curation; formal analysis; methodology; writing – original draft; writing – review and editing; A.M. Collins: conceptualisation; writing – review and editing; D.M. Ferreira: conceptualisation; writing – review and editing

Conflict of interest: C. Hales has nothing to disclose. L. Burnet has nothing to disclose. M. Coombs has nothing to disclose. A.M. Collins has nothing to disclose. D.M. Ferreira has nothing to disclose.

Support statement: This review was funded by a Research Excellence Award, and Research Faculty Grant, Te Herenga Waka, Victoria University of Wellington, Aotearoa New Zealand (grant numbers: 221390 and 400195). Funding information for this article has been deposited with the Crossref Funder Registry.

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