Effect of neuromedin U on allergic airway inflammation in an asthma model (Review)

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Abstract. Asthma is a major inflammatory airway disease with high incidence and mortality rates. The Global Initiative for Asthma released a report called ‘The Global Burden of Asthma’ in 2004. However, the specific pathogenesis of asthma remains unclear. An increasing number of studies have demonstrated that neuromedin U (NMU) plays a pleiotropic role in the pathogenesis of asthma. NMU is a highly structurally conserved neuropeptide that was first purified from porcine spinal cord and named for its contractile effect on the rat uterus. NMU amplifies type 2 innate lymphoid cell (ILC2)-driven allergic lung inflammation. The NMU receptors (NMURs), designated as NMUR1 and NMUR2, belong to the G protein-coupled receptor family. NMUR1 has also been found in immune cells, including ILC2s, mast cells and eosinophils. In view of the important roles of NMU in the pathogenesis of asthma, the present review evaluates the potential mechanisms underlying the impact of NMU on asthma and its association with asthma therapy.

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Abbreviations: Areg, amphiregulin; CysLTs, cysteinyl leukotrienes; CGRP, calcitonin gene-related peptide; DAG, diacylglycerol; DC, dendritic cell; ECM, extracellular matrix; ERK, extracellular signal-regulated kinase; FcεRI, Fc epsilon receptor I; GPCR, G protein-coupled receptor; HDM, house dust mites; IL, interleukin; ILC2s, type 2 innate lymphoid cells; IP3, inositol 1,4,5-trisphosphate; KO, knockout; NFAT, nuclear factor of activated T cells; NMU, neuromedin U; NMUR, neuromedin U receptor; PGD2, prostaglandin D2; PLC, phospholipase C; Th, helper T cell; TSLP, thymic stromal lymphopoietin

Key words: allergy, asthma, type 2 innate lymphoid cells, neuromedin U, mast cells, eosinophils, neuroimmunity

1. Introduction

Asthma is a heterogeneous disease that is usually characterized by chronic airway inflammation with airway hyper-responsiveness, airway remodelling and disordered mucosal immunity (1-3). Most asthma-associated deaths occur in low- and low-middle-income countries. According to the latest World Health Organization estimates released in December 2016, approximately 235 million individuals currently suffer from asthma, and 383,000 deaths occurred due to asthma in 2015 (4). The strongest risk factors for developing asthma are exposure to inhaled substances and particles, such as pollen and house dust mites (HDM), which may provoke allergic reactions or irritate the airways (5,6). Patients with asthma suffer recurrent episodes of wheezing, coughing, chest tightness and shortness of breath. These episodes are usually associated with airflow obstruction within the lung, which is often reversible either spontaneously or with treatment (7). Additionally, the specific pathogenesis of asthma remains unclear. Therefore, the investigation of potential molecular mechanisms will improve our understanding of the pathogenesis of asthma and help us to identify new effective therapeutic targets.

Neuromedin U (NMU) is a multifunctional neuropeptide with pleiotropic effects, including the mediation of intestinal peristalsis and the modulation of the sense of satiety, body weight, circadian oscillation, bone formation, insulin production, cancer development, energy balance and metabolism (8-12). However, these effects will not be addressed in the present review. Recently, reports have demonstrated that the neuropeptide NMU enhances ILC2-driven allergic lung inflammation (13-15). Therefore, the effect of
NMU on the pathogenesis of asthma will be evaluated in the present review.

2. Overview of asthma

Asthma is an airway disease and is characterized by four treatable traits: Airflow limitation, altered cough reflex sensitivity, airway infection and airway inflammation (1,16) (Table I). Airflow limitation is caused by several factors, including the sensitization of airway nerves, the accumulation of mast cells, the repeated obstruction of airway smooth muscle, inflammatory mural oedema, the decreased production of bronchoprotective factors and structural changes to the airway (17,18). Airway hyper-responsiveness is another important factor in the development of asthma and is a result of an imbalance in the autonomic nervous system. It involves various inflammatory cells, mediators and cytokines, damage to the epithelial airway, and the exposure of the subepithelial nerve terminals (19). Thirdly, viral infections aggravate airway inflammation in asthma and can even lead to asthma attacks (20). Finally, airway inflammation is heterogeneous among asthma patients. Eosinophilic airway inflammation is the predominant type of granulocytic inflammation because it is recognizable and treatable (21). In patients with eosinophilic asthma, there are two different pathogenic pathways (22).

In allergic eosinophilic airway inflammation, specialized dendritic cells present allergens to steer the differentiation of naive T lymphocytes towards the formation of Th2 cells, which produce cytokines such as interleukin-4 (IL-4), IL-5, IL-9 and IL-13 that result in IgE switching in B cells (23,24).

The released IgE molecules bind to the Fc epsilon receptor I (FcrRI) on mast cell surfaces (25). Exposure to allergens and the cross-linking of receptors by allergens, which bind to high-affinity IgE result in mast cell degranulation (26). Type 2 innate lymphoid cells (ILC2s) are activated by cysteinyloleukotrienes (CysLTs) and prostaglandin D2 (PGD2) secreted by activated mast cells (27). Once activated, ILC2s rapidly expand and secrete large amounts of IL-5 and IL-13. ILC2s contribute to allergic airway inflammation by directly interacting with Th2 cells to promote the release of cytokines, mucus production and airway eosinophilia (28-30).

In nonallergic eosinophilic asthma, air pollutants and pathogens induce the release of epithelium-derived and macrophage-derived cytokines, including IL-33, IL-25, TSLP and IL-1β, which activate ILC2s in an antigen-independent manner via their respective receptors (Fig 1) (31,32). It was previously shown that IL-13 released by ILC2 disrupts the bronchial barrier integrity of the airway epithelium in asthmatic patients, therefore maintaining type 2 inflammation in the airways (33). In summary, the role of ILC2s in airway epithelium has received more attention.

In the treatment of asthma, inhaled glucocorticosteroids are currently the most effective anti-inflammatory medications for the therapy of persistent asthma. Their efficacy in decreasing airway hyper-responsiveness has been demonstrated, which decreases asthma symptoms and controls airflow inflammation (34). Additionally, long-acting inhaled β2-agonists are more effective when combined with inhaled glucocorticosteroids (35). However, long-term exposure to high doses of inhaled glucocorticosteroids leads to adrenal suppression, easy bruising and decreased bone mineral density (36-38). The key role of IL-5 in eosinophilic airway inflammation make it a crucial drug target, which has driven the clinical development of two monoclonal antibodies against IL-5, reslizumab and mepolizumab (39). Nevertheless, these antibodies are ineffective for non-phenotyped patients with persistent asthma who have already received treatment with inhaled corticosteroids (Table I) (22).

Asthma is an inflammatory airway disease involving a variety of cells and cytokines. Due to the phenotypic heterogeneity of asthma, its pathogenesis is complex; the treatable traits are not separate, but are rather interrelated. For this reason, airway disease emerges as an intractable disease, without the development of novel therapies. NMU acts as a multifunctional neuropeptide in allergic responses. In the following sections, the discovery, distribution and function of NMU are to be discussed. Furthermore, the role of NMU in allergic airway inflammation and its potential for clinical application will be discussed.

3. Biology of neuromedin U

NMU, which was first isolated from porcine spinal cord and named for its potent contractile effect on the rat uterus in 1985, is a highly conserved peptide secreted by cholinergic neurons (40). NMU was found in rabbits, dogs, frogs and chickens; a 23-amino-acid version was identified in rats, and nonapeptides were detected in guinea pigs and chickens (41-43). NMU is therefore widely conserved throughout the animal kingdom and shows almost complete conservation of its amidated C-terminal pentapeptide, indicating that there is a strong evolutionary pressure to conserve this peptide. NMU also has widespread distribution in the peripheral and central nervous system (9,44). NMU-like immunoreactivity (NMU-LI) protein and mRNA are distributed in the stomach, ileum, spleen, pancreas, heart, lung, kidney, prostate, pituitary gland, adipose tissue, bone, bone marrow and lymphocytes in humans (45,46). NMU-LI is also widely distributed in the central nervous system, including the cingulate gyrus, thalamus, locus coeruleus, medulla oblongata, hypothalamus, substantia nigra and medial frontal gyrus in humans (45,46). Due to the high affinity of NMU and the saturable and specific binding sites for NMU-23 in rats, NMU has been previously characterized as a cognate ligand for the designated ‘orphan’ class A G-protein-coupled receptors (GPCRs) (47). Two different receptors exist for NMU, termed NMUR1 (also known as GPR66 and FM-3) and NMUR2 (also known as TGR-1 and FM-4), which are encoded by genes located in human chromosomes 2 and 5, respectively (48-50). NMUR1 is mainly expressed in peripheral tissues, such as the intestine, pancreas, uterus, lung and kidney. NMUR2 is predominantly found in specific regions of the central nervous system, including the spinal cord, dorsal root ganglia and medulla oblongata (51). In the rat spinal cord, NMU-like immunoreactivity protein levels are greater in the dorsal than in the ventral horn, suggesting a sensory role for NMU (41). More recently, using chimeric G proteins, it was demonstrated that NMUR1 primarily signals through the Gq/11 proteins, whereas NMUR2 signals through the Gq proteins (50,52,53). Moreover, NMU mRNA has been detected in antigen-presenting cells,
particularly monocytes and dendritic cells, and NMUR1 mRNA was detected in T cells and natural killer cells (54,55). Therefore, NMU has the potential to serve as a target for the treatment of asthma.

Thus, NMU is a multifunctional neuropeptide with several roles in different cells and tissue types, which relays signals to the central nervous system (CNS) to stimulate organs. NMU also affects cells directly, by increasing cell proliferation and migration, and inducing the release of hormones and autocrine/paracrine factors. These functions are mostly mediated via the NMUR1 and NMUR2 receptors, although other alternative receptors have been described. It is also possible that NMU may also elicit its effects by binding and signalling through unknown receptors. Furthermore, NMU may also serve as an ideal target for the treatment of certain disorders, although its multiple roles should be taken into consideration when inhibiting its functions for therapeutic purposes (10).

4. NMU acts as a multifunctional neuropeptide in the pathogenesis of asthma

**NMU and ILC2s.** ILC2s have recently been identified as effector cells that are key early regulators of immune responses in airway barrier surfaces (33). ILC2s are activated by cell-derived exogenous cytokines, such as IL-25, IL-33 and thymic stromal lymphopoietin (TSLP). Activated ILC2s produce type 2 cytokines, such as IL-5 and IL-13, to initiate allergic inflammation at mucosal surfaces (28,56). However, the specific molecular pathways that mediate the response of ILC2s to alarmins such as IL-25, IL-33 and TSLP remain unclear (29). Notably, NMUR1 is largely specific to ILC2s, according to single-cell RNA sequencing and flow cytometry (13). Moreover, it is highly expressed in ILC2s at baseline and following the induction of airway inflammation with HDM, in contrast with its expression in other lung-resident cell populations. Furthermore, NMU expands IL-25-driven inflammation. IL-25 combined with NMU led to increased expression of IL-5 and IL-13 in the lung and bronchoalveolar lavage fluid, whereas IL-25 alone only modestly increased their expression. NMU combined with IL-25 alters a non-pathologic dose of IL-25 into a pathogenic dose. However, the number of ST2+ ILCs following HDM challenge was decreased in NMU-knockout mice compared with the number in wild-type mice, which indicates that NMU promotes ILC activation and effector function. Moreover, in NMUR1-knockout mice, ILC2 frequency was markedly reduced after HDM challenge compared to that induced by PBS, reflecting the effects also observed in wild-type mice (13). In addition, NMUR1-knockout ILC2s demonstrated lower average inflammatory score than wild-type ILC2s following HDM challenge, which is consistent with the fact that NMU-NMUR1 signalling promotes ILC2 responses in vivo (13-15). In regard to the signalling pathway, through which NMU activates ILC2s, it was shown that NMU activates phospholipase C (PLC), which catalyses the conversion of the phospholipid inositol to diacylglycerol (DAG) and inositol 1,4,5-trisphosphate (IP3). Subsequently, IP3 induces Ca2+ release from intracellular stores (55). Cardoso et al (14) found that NMU triggered extracellular signal-regulated kinase (ERK) phosphorylation and regulated innate type-2 cytokine downstream of a Ca2+/calcineurin/NFAT cascade, leading to the expression of the type 2 cytokine genes IL-5, IL-13 and amphiregulin (Areg) in ILC2s (Fig. 2), (57). Thus, the neuropeptide NMU can activate NMUR1 in an ILC2-mediated manner, resulting in the uniquely potent and immediate production of innate type 2 cytokines.
Figure 1. Physiopathology of asthma. In allergic asthma, dendritic cells present allergens to naive T lymphocytes to induce Th2 cells, which produce IL-4 to induce IgE switching in B cells. The released IgE molecules bind to FcεRI on mast cell surfaces. CysLTs and PGD2, which are secreted by activated mast cells, are activators of ILC2s. ILC2s support Th2 cells by inducing a type 2 response, airway eosinophilia and mucus hypersecretion. In non-allergic eosinophilic asthma, air pollutants and pathogens induce the release of epithelium-derived and macrophage-derived cytokines, including IL-33, IL-25, TSLP and IL-1β, which activate ILC2s in an antigen-independent manner via their respective receptors. This leads to the secretion of type 2 cytokines by ILC2s, including high amounts of IL-5 and IL-13, which leads to eosinophilia, mucus hypersecretion and airway hyperreactivity.

Figure 2. The signalling pathway through which NMU activates ILC2s. NMU activates ILC2s via NMUR1, which is a GPCR. The activated G protein receptor activates PLC, which catalyses the conversion of the phospholipid inositol to DAG and IP3. Subsequently, IP3 elicits Ca2+ release from intracellular stores. The increased Ca2+ influx triggers ERK phosphorylation and activates the Ca2+ calcineurin/NFAT cascade, inducing the increased expression of the type 2 cytokine genes IL-5, IL-13 and Areg. ILC2s, type 2 innate lymphoid cells; GPCR, G protein-coupled receptor; ERK, extracellular signal-regulated kinase; NFAT, nuclear factor of activated T cells; NMU, neuromedin U; NMUR, neuromedin U receptor; p-, phosphorylated; PLC, phospholipase C; DAG, diacylglycerol; IP3, inositol 1,4,5-trisphosphate.
NMU interacts with mast cells. Mast cells are important participants in the early stage of allergic inflammation and are derived from haematopoietic stem cells. Mast cells affect intercellular communication during inflammation by secreting cytokines that contain several mediators (58). In addition, the severity of airway hyperresponsiveness was strongly correlated with mast cell activation (59). IgE binds to FcεRI on mast cells and is crosslinked with specific antigens on the cell surface, which can induce mast cell degranulation and the synthesis of chemokines and cytokines (60). Specifically, mast cells proximal to nerve fibres contain, secrete and respond to several neuropeptides (61). NMUR1 is also highly expressed in mast cells. NMU combines with NMUR1, resulting in mast cell degranulation and the subsequent release of preformed mediators, most notably histamine and chemokines, which leads to early-phase inflammation characterized by vasodilation, extravasation, edema and smooth muscle contraction. Neutrophils also aggregate at reaction sites as a result of the cytokines released by mast cells. The levels of NMU-induced Ca$^{2+}$ release and mast cell degranulation are nearly comparable with those induced by IgE receptor cross-linking. In contrast, the subsequent infiltration of neutrophils is completely inhibited in NMU-deficient mice (62). NMU activates mast cell-mediated inflammation; therefore, NMU receptor antagonists could be novel targets for the pharmacological inhibition of mast cell-mediated inflammatory diseases.

NMU contributes to the accumulation of eosinophils. The activation of mast cells by NMU at an early stage might trigger airway inflammation and the release of eosinophilic chemotactic factors, which attract activated eosinophils during allergen exposure and lead to progressive allergic inflammation (63,64). NMUR1 is also expressed in a mouse eosinophil cell line. NMU elevates intracellular Ca$^{2+}$ levels and ERK phosphorylation and activation, which promotes cell adhesion to components of the extracellular matrix (ECM), and eotaxins contribute to eosinophil accumulation. Inflammatory cells adhere and interact with components of the blood cell wall and the ECM, which aids their ability to extravasate and migrate into inflamed sites (65). NMU acts directly on eosinophils to play an important role in cell activation, adhesion and migration. On the contrary, it was observed that the absence of Gq signalling in Gq-deficient mice blocked the accumulation of eosinophils in the lungs, following an allergic challenge (15,66). NMU directly acts on eosinophils to induce cell adhesion to fibronectin and collagen type I. Additionally, eosinophil chemotaxis is induced by NMU at high concentrations comparable to that induced by eotaxin, which is also known to be involved in integrin activation and the adhesion of eosinophils. Thus, NMU is an important mediator of eosinophil-mediated inflammation, and a potential therapeutic target for bronchial asthma/eosinophil-mediated inflammatory diseases (67). To date, only limited data have been obtained regarding the possible importance of NMU and eosinophils. Therefore, the association between NMU and eosinophils requires further study.

Role of NMU in neuroimmunity. Evolution has generated multiple mechanisms to defend against external and internal sources of danger. For example, the immune system eliminates various threats through a variety of immune cells and antibodies. The nervous system promptly inputs information into the CNS and produces complex defence behaviours. The immune system and the nervous system cooperate with each other to defend against danger. In this, they share a common ‘language’ comprised of cytokines, receptors, and neuropeptides, which enable mutual communication (68). Immune cells are found in close proximity to the nerve terminal processes in the mucosal surfaces of the airways, which are then poised for interaction (23). The lung is extensively innervated via sensory fibres, most of which express markers of nociceptors (69). Asthmatic patients have a denser network of these fibres around small airways and a low threshold for their activation in response to airborne irritants (70,71). This indicates the excess activity of peptidergic sensory fibres. Upon exposure to allergens, nociceptor peripheral terminals release neuropeptides, such as substance P and calcitonin gene-related peptide (CGRP), resulting in neuroimmunity (68,72,73). Furthermore, NMU has been found in the spinal cord, dorsal root ganglia and medulla oblongata via radioimmunoassays and immunohistochemistry (41,74). The distribution of NMU is consistent with that of neurons involved in nociception. One of the physiological roles of NMU may be its involvement in nociception (75). Neurons that secret NMU can be found in the ventromedial hypothalamic regions and in some nuclei of the caudal brainstem regions, which are involved in nociceptive transmission and pain modulation (44,45,76). NMU has been demonstrated to markedly and selectively enhance the excitability of nociceptive neurons in spinal dorsal horns in a dose-dependent manner (77). Hyperactivity of nociceptive dorsal horn neurons induced by NMU could mediate pain-associated behavioural changes and several neuroendocrine functions (49). This expression pattern of NMUR2 mRNA corroborated with the hypothesis that its ligand, NMU, is a sensory transmitter/modulator (77). Compared with the wild-type mice, the nociceptive reflexes were decreased in the NMU KO mice, indicating that endogenous NMU may play an important role in reflexes and in adaptation to environmental stimuli (78). The mRNA expression NMU is increased in the spinal cord but not in the hypothalamus following a pain stimulus, suggesting that pain may stimulate the synthesis of NMU. Thus, NMU is involved in nociceptive reflexes (78). Nociceptor activation upon allergen exposure is a very early event in the development of inflammation. Appreciation of the immune and nervous systems as part of a holistic, coordinated defence system provides new insights into inflammation and exciting opportunities for managing acute and chronic inflammatory diseases (73). In short, these studies suggest that NMU is important for nociceptive reflexes and allergen exposure, although more detailed genetic and mechanistic investigations of NMU and its role in vivo and in vitro are still required.

5. Clinical implications of NMU involvement in asthma

In view of the importance of allergic airway inflammation in the clinical manifestations of allergic diseases, it is of note that NMU can induce immune cell-driven inflammation. Upon inhaling pollutants, microbes and glycolipids, the nervous system rapidly processes information and triggers these processes. NMU acts as a mediator between sensory neurons
and immune cells to potentiate or initiate inflammation. Overall, NMU provides a novel neuroimmune target for the treatment of asthma. At present, there are no investigative reports regarding the development of NMUR subtype-selective antagonists. The natural products EUK2010, EUK2011 and EUK2012 have been identified as NMUR2-specific agonists, and icariin from Herba epimedii has been described as an NMUR2-selective agonist (79,80). Structure-activity relationship study identified the more potent hexapeptide 5d that exhibits NMU1 agonist activity similar to that of Hmnu (81). Moreover, scientists have discovered two synthetic low molecular weight non-selective NMU receptor agonists (45).

Regarding asthma therapy based on the function of NMU, NMU amplifies allergic airway inflammation in an asthma model. This indicates that NMU might be a meaningful therapeutic target for the treatment of allergic airway inflammation in asthma. For example, ketotifen, as a mast cell membrane stabilizer, can protect mast cell membranes to decrease membrane metamorphosis and the release of allergic inflammation mediators (82). Additionally, butyrate ameliorates allergic airway inflammation by limiting eosinophil trafficking and survival (83). Considering the constant progress in pharmaceutical development and molecular biology, screening for novel molecules that act on targets in the human airway and immune cells will be conducted, leading to new treatments for asthma.

6. Conclusion

In the context of the complexity and intricacy of airway inflammation, NMU acts as a multifunctional neuropeptide in the pathogenesis of asthma. An understanding of the function of NMU aids in improving the understanding of the mechanism underlying the pathogenesis of asthma. Considering the constant improvements in organoid culture systems and transcriptomic techniques, NMU receptor antagonists will likely be a novel target for the pharmacological inhibition of asthma in the near future, which may significantly improve the clinical outcomes of patients with asthma. However, the precise regulation of NMU in asthma still requires further study, in order to be used in clinical applications.

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Authors' contributions

XR participated in the entire review process and prepared the manuscript. FD and YZ contributed to collecting the relevant literature. YW and WM conceived the review and modified the manuscript. All authors read and approved the final version of the manuscript.

Ethics approval and consent to participate

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Patient consent for publication

Not applicable.

Competing interests

The authors declare that they have no competing interests.

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