going innate immune airway responses in asthma. The relevance of this to clinical disease expression requires understanding.

S43

THE EXPRESSION OF CRACM ION CHANNELS IN HUMAN LUNG MAST CELLS

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Introduction and objectives The influx of extracellular Ca^{2+} into human lung mast cells (HLMC) is essential for the IgE-dependent release of preformed granule derived mediators and newly synthesised autacoids and cytokines. However the identity of the ion channels underlying the influx of Ca^{2+} remains unknown. Members of the recently discovered CRACM (also known as Orai) ion channel family are candidates for this function. The channels carry Ca^{2+} selective currents (I_{CRAC}) that are activated when endoplasmic reticulum Ca^{2+} stores are emptied. Here we investigate whether CRACM channels are expressed in HLMC and their possible contribution to Ca^{2+} influx in these cells.

Methods HLMC were purified from lung tissue by immunomagnetic affinity selection. Total RNA purified from cells was used in quantitative RT-PCR using TaqMan probes and primers (Applied Biosystems). For electrophysiology the whole cell method of patch clamp was used.

Results CRACM1, CRACM2 and CRACM3 mRNA transcripts were all readily detectable in HLMC. CRACM1 transcripts (4.50±0.51 copies/10³ beta-actin transcripts, n=7 HLMC donors) were more abundant than CRACM2 transcripts (1.79±0.67 copies/10³ betaactin transcripts, n=5) and CRACM3 transcripts (3.33±0.95 copies/ 10³ beta-actin transcripts, n=5). Consistent with the presence of CRACM transcripts, an I_{CRAC}-like current was detected in HLMC following IgE-dependent activation and also in cells dialysed with 30 μM inositol triphosphate to empty endoplasmic reticulum Ca²⁺ stores. The Ca²⁺-selective current obtained under both conditions was blocked by the addition of 10 μM La³⁺, a known blocker of CRACM channels. Thus following addition of anti-IgE to 11 HLMC, current at -120 mV increased from -12.1 ± 1.1 pA at baseline to -34.2 ± 2.5 pA (p<0.0001) and following addition of 10 mM La³⁺ this was reduced to -17.4 ± 2.1 pA. (p=0.0003). When HLMC were dialysed with IP₃ (n=12), current increased from -15.3 ± 1.2 pA to -40.2 ± 3.2 pA (p<0.0001), and decreased to -17.7 ± 1.5 pA (p<0.0001) by La³⁺.

Conclusions The presence of an I_{CRAC} -like current and CRACM1, -2 and -3 mRNA transcripts in HLMC supports a role for CRACM channels in the influx of extracellular Ca²⁺ into these cells following activation. Such a role would make CRACM channels a potential therapeutic target in the treatment of asthma.

S44

INCREASED EXPRESSION OF IMMUNOREACTIVE THYMIC STROMAL LYMPHOPOETIN IN SEVERE ASTHMA

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Background Thymic stromal lymphopoetin (TSLP) is a cytokine implicated in the pathophysiology of asthma through two pathways: a TSLP-OX40L-T cell axis and a TSLP-mast cell axis. Whether these pathways operate in human asthma is unknown.

Aims To investigate whether mucosal TSLP protein expression relates to asthma severity, and distinct immunological pathways.

Methods GMA-embedded bronchial biopsies from healthy subjects (n=12) and patients with mild (BTS step 1, n=8), moderate (BTS steps 2 and 3, n=12) and severe (BTS steps 4 and 5, n=16) asthma were immunostained for TSLP, OX40, OX40L, CD83, IL-13, and inflammatory cell markers. Extent of immunostaining was correlated with clinical data.

Results Specific TSLP immunoreactivity was evident in both the airway epithelium and lamina propria of both healthy and asthmatic subjects. TSLP immunoreactivity was not present in airway smooth muscle. TSLP epithelial expression was significantly elevated in asthma as a whole compared to healthy controls (p=0.0005), particularly in mild and severe disease. The number of TSLP+ cells in the lamina propria was elevated in patients with severe asthma relative to other groups (p=0.0058). Co-localisation studies in 6 severe asthmatic subjects showed that 5% of TSLP+ cells in the lamina propria were CD68+ macrophages and 20% were tryptase+ mast cells. There were significant inverse correlations between TSLP counts in both the asthmatic bronchial lamina propria and epithelium with the FEV₁/ FVC ratio ($r_s = -0.53$, p = 0.002 and $r_s = -0.40$, p = 0.037). Immunostaining for OX40, OX40L and CD83 in the airways was sparse, with no difference between asthmatic patients and normal control subjects. IL-13 staining was increased in non-epithelial cells within the airway epithelium in severe asthma (p=0.033).

Conclusions TSLP expression is elevated in severe asthma despite high dose corticosteroid therapy. Although we did not detect activity of the TSLP-OX40L-T cell pathway within asthmatic bronchial mucosa, it is possible that this pathway operates in secondary lymphoid organs such as draining lymph nodes. The close approximation of airway stroma and mast cells suggests that the TSLP-mast cell axis, in which TSLP and IL-13 operate in a positive regulatory loop, is active in asthmatic bronchial mucosa and may be important in maintaining chronic airway inflammation.

S45

FUNCTIONAL KCa3.1 K⁺ CHANNELS ARE REQUIRED FOR HUMAN FIBROCYTE MIGRATION

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Background Fibrocytes are bone marrow-derived CD34⁺ collagen I⁺ cells present in peripheral blood that are implicated in the pathogenesis of tissue remodelling and fibrosis in both asthma and pulmonary fibrosis. Ion channels play key roles in cell function, but the ion channel repertoire of human fibrocytes is not known. The aim of this study was to assess whether human fibrocytes express the $K_{\text{Ca}}3.1~\text{K}^+$ channel.

Methods Human fibrocytes were isolated from peripheral blood taken from both normal and asthmatic subjects and maintained in cell culture. Channel expression was examined using RT-PCR, Western blot, and patch clamp electrophysiology. Migration assays were performed using the Transwell system.

Results Fibrocytes expressed both $K_{Ca}3.1$ mRNA and protein. At baseline unstimulated fibrocytes (n=30 healthy cells from 7 donors, n=17 asthmatic cells from 4 donors) demonstrated heterogeneity of the resting whole cell current, with a small outwardly rectifying current in 91% of cells, a steeply inwardly rectifying current in 30% of cells, and a linear current in 9% of cells. The $K_{Ca}3.1$ opener 1-EBIO (100 mM) elicited typical $K_{Ca}3.1$ currents in 46/47 cells tested. Whole cell current at +40 mV increased from 122±25 pA pre-1-EBIO to 795±104 pA post 1-EBIO (p<0.0001). This was accompanied by a negative shift in reversal potential from -29.8 ± 3.0 mV to -61.8 ± 1.6 mV (p<0.0001). 1-EBIO induced a greater current in