DYSREGULATED IRON METABOLISM MEDIATED BY IRP2 MAY INFLUENCE LUNG CANCER PROGRESSION, PARTICULARLY IN THE CONTEXT OF CIGARETTE SMOKE EXPOSURE

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Background Iron is required for cell growth, and various cancers have been shown to proliferate more readily when iron replete. We have shown previously in lung cancer and further demonstrated that this was reduced by either iron chelation or knockdown of IRRE2, an iron regulatory gene. 1 Differences in iron content of bronchoalveolar lavage (BAL) fluid have been reported in smokers compared to non-smokers, 2 so we hypothesised that iron dysregulation might be an active mechanism of cancer progression in smokers.

Methods Two lung cancer cell lines were cultured with either ferrous (Fe2+) or ferric (Fe3+) forms of iron, or with cigarette smoke extract (CSE). Proliferation, apoptosis, necrosis and migration were assessed by BRDU assay, FACS and scratch wound assay respectively. Iron regulation was assessed by means of gene expression and Western blot for IREB2 (protein product IRP2), ferritin and transferrin receptor.

Resected lung cancers (n = 78) were stained for iron regulatory protein 2 (IRP2) and staining related to clinical features such as tumour size and survival. Cancer cells proliferated more in the presence of ferrous iron or 5% CSE (p = 0.045) and survival poorer (p = 0.079).

Conclusions Proliferation of cancer cells driven by iron dysregulation may be a clinically relevant mechanism in lung cancer, particularly in smokers.

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METHODS TO ISOLATE BASAL CELLS FROM THE RESPIRATORY EPITHELIUM

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Introduction and objectives Maintenance of a healthy respiratory epithelium is essential in the prevention of airway disease. Perturbations in airway homeostasis have been linked to the pathogenesis of airway disease including asthma, fibrosis and lung cancer. The ‘stem cell hypothesis’ describes how a change within cells responsible for airway maintenance and repair can lead to development of cancer. Basal stem/progenitor cells in the upper airways are suggested to represent the cell of origin in squamous cell carcinoma and therefore are of research interest.

Isolation of this cell type has been hampered because established airway enzymatic digestion methods destroy epitopes of interest on the surface of basal cells. We sought an optimised method of digestion for the isolation of viable basal cells from murine, and then human, airway epithelium.

Methods Conventional enzymatic digestion of the murine upper respiratory epithelium involves an overnight pronase incubation. Using flow cytometry, we compared this strategy to other reported methods: a dispase/trypsin digest, 3 collagenase incubation and a combination of these.

A method allowing selection of a pure basal cell population in the mouse trachea was subsequently translated to human airways to assess its efficacy.

RESULTS Following pronase digestion, only 2% of epithelial cells were basal cells, probably as a result of enzymatic epitope removal. Optimal extraction of murine basal cells involved removal of the epithelium through a dispase/trypsin incubation followed by incubation of tracheal remnants in collagenase to release the remaining basal cells from submucosal glands. We identified a well-defined basal cell population representing 30% of the airway epithelium, consistent with known airway basal cell frequency, which can be isolated by fluorescence-activated cell sorting.

Application of this strategy to digest human airways revealed a comparable population of basal cells and allowed sorting of a viable cell population.

Conclusions We optimised a method to facilitate the extraction of basal epithelial cells from both mouse and human airways. This strategy allows sorting of a pure, viable basal cell population for use in further assays.

REFERENCES
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MMP12 AND LMO7 ARE KEY GENES INVOLVED IN THE EARLY PATHOGENESIS OF SQUAMOUS CELL CARCINOMA OF THE LUNG

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Lung cancer is the most lethal cancer type worldwide. In order to improve patient survival it is important to enhance our understanding of the early changes associated with lung cancer progression. UCLH has a unique cohort of patients with pre-invasive lung squamous cell carcinoma (SCC) lesions. Within this cohort there is a discrepancy between the prevalence of pre-invasive lesions and the incidence of invasive lung cancer, which suggests that not all pre-invasive lesions progress to invasive carcinomas. The aim of this study was to identify and characterise key genes involved in the early pathogenesis of lung SCC.

We performed genome-wide gene expression Illumina Whole-Genome DASL® arrays in 19 regressive and 20 progressive pre-invasive lung SCC lesions. The expression of matrix metalloproteinase 12 (MMP12) and LIM domain 7 (LMO7) was also determined in the 39 pre-invasive lung cancer lesions by immunostaining analysis. The functional role of MMP12 and LMO7 in cell migration and invasion was demonstrated by MMP12 and LMO7-knockdown in different squamous cell carcinoma cell lines and human bronchial epithelial cells (HBEcs), respectively.

We found 939 genes significantly differently expressed between the progressive and the regressive pre-invasive lung SCC lesions. We identified a remarkably elevated expression of a spectrum of genes in the progressive lung SCC lesions involved in different