Airway gene transfer in a non-human primate: Lentiviral gene expression in marmoset lungs

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Genetic therapies for cystic fibrosis (CF) must be assessed for safety and efficacy, so testing in a non-human primate (NHP) model is invaluable. In this pilot study we determined if the conducting airways of marmosets (n = 2) could be transduced using an airway pre-treatment followed by an intratracheal bolus dose of a VSV-G pseudotyped HIV-1 based lentiviral (LV) vector (LacZ reporter). LacZ gene expression (X-gal) was assessed after 7 days and found primarily in conducting airway epithelia as well as in alveolar regions. The LacZ gene was not detected in liver or spleen via qPCR. Vector p24 protein bio-distribution into blood was transient. Dosing was well tolerated. This preliminary study confirmed the transducibility of CF-relevant airway cell types. The marmoset is a promising NHP model for testing and translating genetic treatments for CF airway disease towards clinical trials.

The relentless progression of cystic fibrosis (CF) airway disease remains an unresolved problem although the rate of decline has been slowed by the range of treatments now available. The majority of morbidity and mortality relates to the chronic airway infection and inflammation that commences in early infancy and leads to premature death from lung failure.

Gene-based correction of defective airway epithelial cell function, by introduction of the functional CF transmembrane conductance regulator (CFTR) gene, is a rational approach designed to produce lasting therapeutic benefit and could be the basis for a cure. Practical success would restore CFTR ion-channel function, producing a physiologically-balanced treatment for CF airway disease. Additionally, CFTR gene transduction of airway stem/progenitor cells should produce extended correction because the daughter cells that re-populate the airway epithelium contain the corrected CFTR gene. Early correction, when CF lungs are considered to be functionally unaffected, has the potential to prevent the initiation of CF lung disease.

In mice our single-dose lentiviral (LV) airway gene transfer technique produces immediate as well as long-term airway reporter gene transfer, extending up to a mouse lifetime. Use of a lysophosphatidylcholine (LPC) airway pre-treatment improves gene transfer and may enable access to airway stem/progenitor cells able to support long-term gene expression. Importantly, this method has already shown its utility with the therapeutic CFTR gene; we have reported more than 12 months of functional CFTR gene expression in CF mouse airways. Furthermore, we have demonstrated robust reporter gene transfer in mouse lung and produced low-level proof-of-principle expression in sheep lung.

The aim of this pilot study was to test the potential of this method in the lungs of a non-human primate (NHP), the marmoset (Callithrix jacchus). The marmoset has a number of advantages as a NHP animal model for studying lung gene transfer. In particular, due to its small size, similar in body size to a very large rat, the modest LV volumes required are within the capabilities of current vector production methods.

Results
This preliminary study was designed to establish if our LV vector delivery system could transduce NHP conducting airway tissue prior to proposing and designing more extensive NHP gene transfer studies. For that reason in this short-term study only two animals (one male and one female) were used. No experimental controls were employed since published data and our previous studies in mice and sheep strongly indicate that none of these controls would elicit gene expression.
LPC and LV dose deliveries were uneventful. General behaviour (demeanour, feeding, respiration) and auscultation findings were within normal limits over the post-treatment study period. In both animals the gross appearance of the lungs was unremarkable when the chest was opened. Histological examination of lung tissue by a veterinary pathologist showed no abnormalities (including cellular infiltration) of consequence.

**LacZ gene expression: Histology.** One week post-dosing LacZ gene expression was visible in excised lung tissue from both marmosets, although the female showed a higher level of transduction (Figs. 1–3 from the female marmoset). Figure 1 shows an *en face* example of LacZ gene expression in lung tissue. LacZ gene expression was present primarily in surface and basal epithelial cells in the conducting airways. The patchy blue cell staining had a generally peribronchiolar distribution along conducting airways in several lobes (Fig. 1). In cross-sections of the trachea (Fig. 2) patchy transduction was observed across the full thickness of the epithelium. In the alveolar tissue (Fig. 3) there was substantially less cell transduction present compared to the conducting airways. The transduced cells in the distal lung airways included alveolar Type 1 and Type II cells (Fig. 3b–c) and inflammatory cells such as alveolar macrophages (Fig. 3d–e), and these were distributed across the lobes, again in a patchy pattern. The macrophages displayed blue LacZ staining indicating that both direct transduction and phagocytic capture of transduced cells could occur.

Histological analysis of the liver and spleen samples showed no LacZ gene expression.

**LacZ gene presence: PCR.** LacZ gene presence in the lung, liver and spleen was assessed by quantitative PCR (qPCR). The LacZ gene was detected in the lung tissue from the female marmoset (the *AC* values for the three tissue samples, performed in triplicate, were 11.91, 6.57 and 5.89), but not in the sampled lung tissue from the male. The LacZ gene was not detected in the liver or spleen tissue from either marmoset, nor in scavenged (untreated colony cull) marmoset control tissue.

**Discussion**

To our knowledge these are the first studies to test and show that airway gene transfer can be achieved *in vivo* using a VSV-G pseudo-
typed HIV-based LV vector in an adult NHP model. Our findings extend earlier work in human airway cell cultures and tracheal explants showing efficient transduction with an Ebola-pseudotyped HIV vector, and airway transduction in infant macaques after in utero parenchymal lung injection of a VSV-G pseudotyped HIV-based vector. The primary finding was that the marmoset lung can be transduced using our LV vector delivery system. Based on our behavioural assessments, gross post mortem organ examination, and histological analysis, no adverse effects of either LPC or the LV vector were apparent. As this was a limited observational pilot-study design there were insufficient numbers of animals to provide detailed quantitative comparisons.

The patchy gene expression observed in both marmosets has also been observed in mice and sheep when fluid-based gene vector dosing was used. This may be caused by multiple factors including, but not limited to, incomplete coverage of the large surface area of the lung airways by the relatively small dose volume, specificity of the vector for particular cell types, and a mismatch in the areas treated by the LPC and LV vector doses. Importantly, the patchiness that would accompany LV-CFTR vector dosing in the same way does not prevent physiologically functional CFTR gene expression occurring in our CF mouse nasal airway studies.

Ciliated airway surface epithelial cells are a primary target of CF gene therapies, but are terminally-differentiated. Gene expression will be lost when those cells undergo apoptosis as part of normal lung cell turnover processes. The lifespan of these epithelial cells in the marmoset is unknown, but transduction of stem/progenitor cells may be important for producing long-term gene expression. Studies in mice suggest that the epithelial stem/progenitor cells responsible for regular regeneration of the conducting airway epithelium reside, in part, in the epithelial basal-cell compartment. The transient permeabilisation that LPC imposes on epithelial cell tight junctions can provide access to those deep lying basal cells.

Importantly, we detected LacZ gene expression in these deep lying epithelial cells at 7 days, suggesting our LV vector can reach regions where some niches of adult stem/progenitor cells are expected to reside in mammalian airways. Similarly, alveolar epithelial cells and macrophages were also transduced, with the latter finding extending the results in rodents to a NHP model.

LacZ gene expression was not detected via X-gal staining or qPCR in any sub-epithelial tissues or in the liver or spleen. Although LacZ positive cells were visible in the lungs of both animals, the negative qPCR finding in the male may have been the result of the randomly selected tissue samples not containing any LacZ transduced cells. The finding of a transient presence of the vector protein component p24 in serum (Fig. 4) confirms that although LV vector or vector components can reach the circulation they are rapidly removed.

Although even conventional lipid-based lung gene transfer events can deliver vector DNA to non-target organs, further studies are required to elucidate the mechanisms and implications of any off-target organ transduction.

While the finding of gene transfer and expression in airway epithelial cells is clear, there are obvious limitations. The very small group size and absence of control animals (other than scavenged tissue for qPCR analyses) were deliberately chosen to prevent unwarranted use of additional NHPs should reporter gene transfer had not proven effective. With the success of this pilot study, future studies now warrant the use of larger group sizes including control animals. Studies should be designed to assess the longevity of gene expression, and should include a LV-only treatment group to directly determine the requirement for LPC in the production of both short and long-term lung gene expression. Additional detailed biosafety assays and physiological, behavioural, immunological and histological monitoring must also be included.

In summary, this pilot study demonstrated the successful extension of our LV delivery technique into a NHP lung. In the translation of basic and preclinical science into useful human therapies this initial success validates the use of LV gene delivery into a NHP lung and suggests marmosets could serve as a valuable pre-clinical model for translational studies in respiratory medicine.

Methods

This multi-institutional study was approved by the Animal Ethics committees of the Women’s and Children’s Health Network, South Australia; and Monash University, Victoria. Two marmosets, one female (F: ~275 g) and one male (M: ~300 g) were sourced from the marmoset colony of the NH&MRC National Primate Colony, Monash University. Animal husbandry, experiments and subsequent monitoring were performed under specialist NHP veterinarian supervision and care provided by Monash Animal Services, Victoria, Australia.

Airway pre-treatment. Airway pre-treatment with LPC (Sigma Aldrich L4129, prepared w/v in PBS) was given prior to a single airway gene transfer event. The LPC concentration (0.1% w/v) was derived from mouse studies where effectiveness and toxicity were previously published methods. Virus titre was $1.2 \times 10^8$ viral genome equivalents per ml as assayed by quantitative PCR. The LV vector volumes (F: 350 µl and M: 500 µl) were also estimated by scale-up from our mouse deliveries.

Pre-treatment and LV dosing. Anaesthetic induction was with 10 mg/ml alfaxalone (Jurox Rutherford, NSW, Australia) and animals were intubated with an endotracheal tube (Sheridan Uncuffed 2.0, Hudson RCI, USA) placed with the tip midway between the epiglottis and the carina. Anaesthesia was maintained using 1.25% – 1.5% isofluorane. Dosing was performed in a Class II biosafety cabinet with animals held supine throughout. The LV vector was delivered as a single fluid bolus to the trachea via the endotracheal tube over ten seconds, one hour after the LPC pre-treatment. Animals were held anaesthetised and supine in the cabinet for an additional hour after LV dosing before being extubated and placed in PC-2 caging facilities.

Monitoring. Vital signs were monitored throughout dosing and in the post-operative period. Body weight and general behaviour were monitored daily along with blood sampling (centrifuged at 13,000 rpm, with sera stored at -80°C until required).

Tissue harvesting. Animals were humanely killed at 7 days using an Alfaxalone induction followed by pentobarbital overdose. The lungs were inflation-fixed in situ in 2% paraformaldehyde/0.5% glutaraldehyde (PFA/ glut) in PBS at 4°C and a pressure of 30 cmH2O for 15 mins. Samples of liver and spleen were also harvested and fixed in PFA/Glut at 4°C.

LacZ gene expression: Histology. The lungs were then excised and submerged in fresh chilled PFA/Glut overnight, and were subsequently processed for LacZ expression by our standard X-gal staining method. The extent of LacZ expression in airway cells was first assessed en face in gross transversely-sectioned portions of lung. Blocks that showed blue LacZ staining were then prepared for routine histological sectioning with haematoxylin/eosin (H&E) or a Safranin-O counterstain.

LacZ gene presence: qPCR. Triplicate tissue samples (20 mg) were selected from random locations in the lungs, liver and spleen. Scavenged tissue from untreated animals was used as a negative control. All lung samples were then incubated at 55°C.
overnight in tail lysis buffer (Viagen Biotech, USA, Cat. # AB102-T) and 20 mg/ml Proteinase K (Promega, USA, Cat. # V3021) at a 1:50 ratio then heated to 80°C for one hour and crude lysates were stored at –20°C until analysis. All other samples were processed via the Wizard SV Genomic DNA Purification System (Promega, USA, Cat. # A2361) according to manufacturers specifications.

qPCR for the NLS-LacZ gene and marmoset GAPDH housekeeping gene was performed in a 96 well plate format (CFX Connect Real-Time PCR, Bio-Rad) according to the manufacturers standard protocol. Specific amplification was detected using SYBR Green (Fast SYBR Green Master Mix, Applied Biosystems USA, Cat. # 4381562). Cycle thresholds (Ct) for the LacZ gene were normalised with respect to the housekeeping gene and presented as ΔCt. The following primers were used: NLS-LacZ forward GCC ACT TCT TGA TGG ACC ACT T, NLS-LacZ reverse CCG CCA CGG ACA TCA TCT; GAPDH forward AAA GTG GAT GTC GTC GCC ATC AAT GAT and GAPDH reverse CTG GAA GAT GGT GAT GGG ATT TCC ATT (GeneWorks, Australia).

Vector particle dissemination. Sera were analysed using a HIV-1 p24 ELISA Kit (Perkin Elmer Life Sciences USA Ca# NEK050) performed as per manufacturer instructions.

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Author contributions
D.W.P. conceived and designed the study. D.W.P., D.M. and R.B. performed the gene transfer; M.D. wrote the manuscript. All authors reviewed the manuscript.

Additional information
Competing financial interests: The authors declare no competing financial interests.

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