

MUCOCILIARY TRANSPORT BEHAVIOUR OF POLLUTANT AND MARKER PARTICLES ON LIVE MOUSE AIRWAYS

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BACKGROUND:

- Airborne pollutants contribute to the development of respiratory diseases such as cystic fibrosis (CF)
- We have previously developed synchrotron imaging techniques for examining the mucociliary transport (MCT) behavior of particulates in the nasal airways and trachea of anaesthetised mice
- Our previous studies delivered particulates in a fluid bolus, however the presence of fluid perturbed the airway surface and would have altered both the manner of deposition and post-deposition MCT behaviour
- The aim of this study was to develop a technique that can be used to detect, monitor and compare the deposition and MCT behaviour of pollutant and marker particles after dry deposition into the trachea of live mice

METHODS:

- Particles of fibreglass, quarry dust and lead ore, as well as reference 14 µm hollow glass beads were delivered to the trachea via the ET tube using a dry powder insufflator and air pump (PneumCentury, Wynnomatic, PA, USA)
- Images of the airway surface were captured at 1000 frames per second (fps) using a synchrotron X-ray beam at the SPring-8 synchrotron in Japan
- Images were obtained post-experimentally and analysed using software to generate movies showing particle motion and also analysed using X-ray velocimetry

RESULTS:

- The first movement of particulates tended to appear approximately 10-20 seconds after deposition
- As in previous studies the particle transit was heterogeneous: after deposition some particles did not move, while others transited the field of view rapidly
- The big and heavy particles of lead and tantalum moved substantially less than the other particulates after deposition
- Most particles did not follow a linear path along the airway: many followed seemingly random, tortuous paths
- There was more movement of particles along the bottom quarter of the tracheal surface than the remainder of the trachea, possibly due to the quantity of fluid present and gravitational effects
- In some animals we also saw radial and axial contraction of the tracheal airway throughout the imaging period

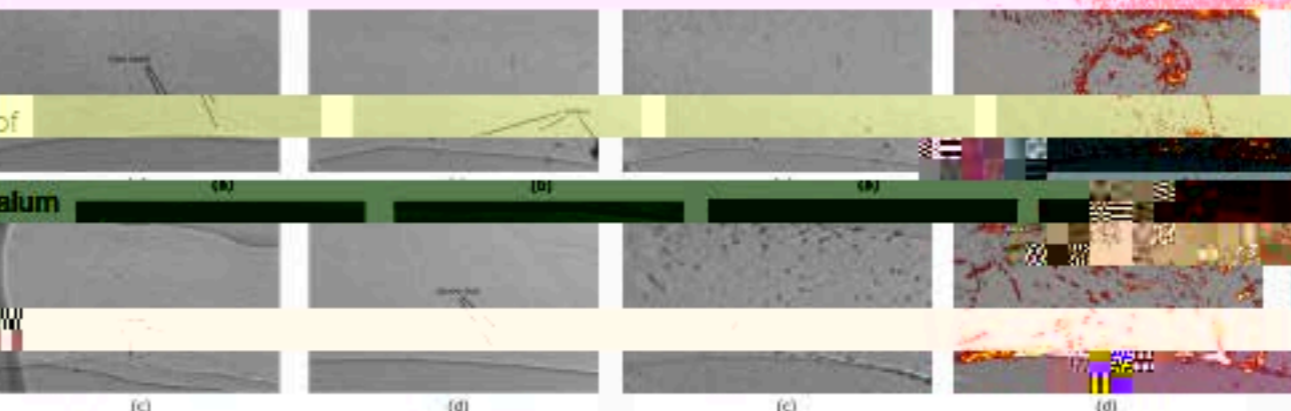


FIG 1: Particle visibility on the airway surface of live mice (a) silver coated hollow glass beads, (b) lead, (c) fibreglass fibres, and (d) quarry dust. In these static images it is hard to clearly identify some of the particles (excluding the large lead particles and fibreglass fibres), but in the image sequences the dynamic behaviour of the particulates is clear. Images are 1.8 x 1.2 mm and the imaging location is just above the main carina.

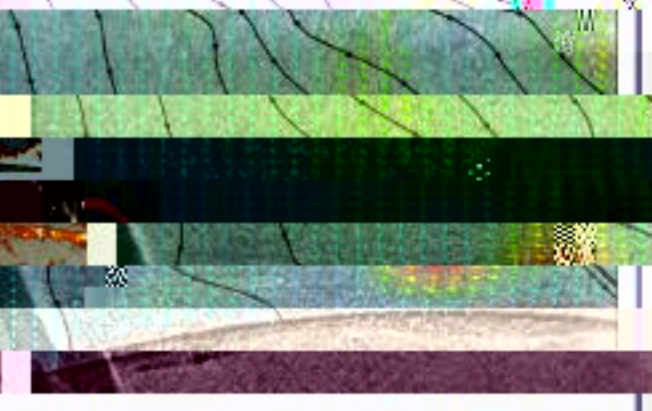


FIG 3: X-ray velocimetry (XV) analysis using a Mechanical Engineering flow measurement technique. On average, there is more particle MUC of hollow glass beads along and towards the dorsal quarter of the tracheal surface. A grid of the average movement vectors (green arrows) with black streamlines reveals all particle motion over the entire imaging period. Figure orientation and location is the same as in FIG 1.

CONCLUSION:

Synchrotron PCXI permits detection of particle transit via MCT along live mouse trachea. We are continuing with studies to improve our direct and non-invasive MCT assessment methods to assist our understanding and treatment of respiratory diseases such as CF.

ACKNOWLEDGEMENTS:

Support provided by NH&MRC Australia and the CureACE Foundation. SPring-8 experiments performed under proposal 2010B1137. Travel funding provided by the Australian Synchrotron SA Program. We thank Naoto Yoneda and Desugi for their assistance.