Bacteriological evaluation of a down-draught necropsy table ventilation system

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Abstract

Aims—To evaluate the microbiological efficacy of a down-draught necropsy table ventilation system (which surrounds the cadaver with a "curtain" of air under continuous extraction) during post mortem procedures.

Methods—Air sampling was carried out both in the presence and absence of staff and cadaver and during a full post mortem procedure, with functioning and non-functioning table air extraction. The penetration of the air "curtain" was also examined during the use of an oscillating bone saw by means of a tracer organism, Bacillus subtilis var nigric, painted on to the skull.

Results—There was little difference between bacterial counts obtained in the presence of staff only, staff plus cadaver, or during a post mortem examination. With all counts obtained, however, there was a two to three-fold reduction when the ventilation was in operation compared with when the extract duct was occluded. Using the tracer organism, a two to three log reduction in counts was shown when the "curtain" was in operation during the use of the oscillating bone saw.

Conclusions—These results suggest that the system provides potential protection for post mortem room staff against airborne infections.

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A few studies have highlighted the risk of infection to staff working in post mortem rooms.\(^1\) The most commonly recorded infections are pulmonary tuberculosis, hepatitis B, and non-A, non-B hepatitis. Wound and gastrointestinal infections occur less frequently. Other potential hazards include HIV infection, viral haemorrhagic fever, other viral infections, and Creutzfeldt-Jacob disease.

Most of these infections are probably acquired by direct contact or following puncture wounds, although dissemination of Mycobacterium tuberculosis in air and on to surrounding surfaces has been shown during the slicing of infected lung.\(^2\) Pulmonary tuberculosis in post mortem room staff is most likely acquired by inhalation. There is little evidence that blood-borne infections are spread in air, although there is some indirect evidence for aerosol transmission of hepatitis B.\(^3\) More recently, airborne dissemination of HIV has been demonstrated in an experimental model investigating possible aerosol transmission from high speed drills in the operating theatre setting.\(^4\) Consequently, there is some concern that certain procedures, including the use of mechanical saws in the post mortem process, might disseminate infectious agents.

One means of minimising potential airborne spread of infection is to use a post mortem isolator tent, but this is cumbersome, reduces visibility, and increases the time required for the procedure.\(^5\) Local exhaust extraction has been suggested as another means of hazard reduction. A ventilated post mortem table applies this principle on a large scale, surrounding the whole cadaver with a curtain of air under continuous extraction. There is little published evidence to show that their use induces significant reduction in the numbers of airborne organisms and so the extra cost of such tables has not been considered justified. The design of the first tables produced also made them difficult to clean. We describe our experience with a newly designed system recently installed at the Royal Free Hospital.

Methods

POST MORTEM ROOM VENTILATION SYSTEM

To overcome some of the problems associated with the previously available ventilated necropsy tables three 207.5 × 75.5 cm tables were constructed to a new design by Afos Ltd., Hull, in collaboration with one of us (JEMcL) and, together with three Afos ventilated dissection tables, these were installed in the refurbished post mortem room in 1989. The table design allows air to be extracted around the perimeter of the whole table through three rows of 1.27 cm diameter holes surrounding the working surface of the head half of the table and two rows surrounding the foot half. The general post mortem room ventilation system was designed in collaboration with Ronald Lear and Associates, Cheshunt, Herst., to provide a general downwards displacement of air in the room. Fresh air entry into the room is provided through a ceiling grid with the entire extract ventilation of the room through the six tables. This maintains the room at negative pressure and ensures that an air extract is always closer to the source of contamination than the operator. The extract from each necropsy table is linked to the extract from its associated dis-
section table and an adjustable damper is provided so that the entire extract ventilation can pass through two pairs of tables or all three. With air extraction switched through two pairs of tables, the flow rate through the necropsy table is set at 0.566 m³/second and through the dissection table at 0.212 m³/second. Some air is additionally extracted into an adjacent storage area at 0.271 m³/second, giving a total extraction rate for the post mortem room of 1.828 m³/second under all conditions. With this system the air exchange rate in the post mortem room is 30.6 air changes/hour.

The tables were designed to be easily cleaned with liquid waste draining into a sump below the working surface where it is diluted with running water and run off under gravity towards a sump drain. The pedestal is fitted with a hydraulic lift to allow the working height of the table to be easily adjusted.

**AIR SAMPLING**

Air samples were collected using a Casella slit sampler. The sampler was used to collect air samples at 2 minute intervals (1400 litres air per sample) on to blood agar plates. These were taken at the four corners of one of the post mortem tables which had been selected at random for the whole study. Above each point samples were taken from three different heights, table height (0 cm), cadaver height (20 cm), and post mortem staff head height (80 cm). Four samples were taken at each height and the mean colony counts obtained. Samples were taken in the following situations: (1) when the examining staff were alone in the room without the cadaver; (2) in the presence of staff and cadaver; (3) during the post mortem examination.

A complete set of samples was taken with the extraction port in the table occluded and a further set with it fully open.

The dispersal of bacteria when opening the skull was investigated using a tracer organism *Bacillus subtilis var niger* (NCTC 10073). A suspension containing 10⁹ spores/ml was painted on to the outer surface of the skull and the bone cut with a De Souther M 170 oscillating saw. Air samples were taken in duplicate above the corners of the head end of the table. Settle plates were also exposed at six different places in the post mortem room: (a) beneath the post mortem table; (b) on the post mortem table within the "curtain"; (c) on a trolley 2 m away at a height of 70 cm; (d) on the dissecting table 2.5 m away; (e) on a basin 3 m away at a height of 75 cm; (f) in a corner of the room at a distance of 3.6 m.

**MEASUREMENT OF AIR FLOWS**

Air flow above the table was measured using a hot wire anemometer (Airflow Developments Ltd., High Wycombe, Bucks). Air flow at table level was measured by placing an anemometer (Griffin and George, Loughborough, England) on the necropsy table so that the 7.0 cm diameter aperture of the instrument covered five ventilation holes at the head end and four at the foot end of the table. Additional measurements were made at the quarter and three-quarter points on each side, as well as at the mid-point of the foot and head end of the table. Air movement was monitored by means of smoke testing. Smoke was continuously produced by a smoke generator placed on the table until a steady state had been achieved. Cold smoke was also intermittently released from smoke tubes (MSA Ltd., East Shawhead, Coatbridge, Scotland) at a height of 20 cm above the table and 20 cm away from the edge at two points one-third and two-thirds of the distance along each side of the table.

The blood agar plates were incubated aerobically for 24 hours at 37°C and for a further 48 hours to detect fungi. All organisms were identified using standard laboratory methods.

**Results**

**AIR SAMPLING**

The mean total aerobic colony counts/1400 l of air at the different heights above the post mortem table with and without the ventilation in operation are shown in figs 1 and 2. The total counts did not exceed 144/1400 l air in the absence of air extraction and were reduced between two and three-fold when the ventilation was in operation.

The organisms isolated were coagulase negative staphylococci, *Staphylococcus aureus*, *Micrococcus* spp *Bacillus* spp and filamentous fungi. Gram negative bacteria were isolated on only one occasion.

The mean counts of the tracer organism *B subtilis var niger* are shown in fig. 3. The effect of the local air extraction was to reduce the counts from >10⁷/1400 litres air to 10⁶/1400 litres air. The settle plate counts increased from a maximum of 5 bacterial colonies during a sample period of 60 minutes when the room was empty to 140 colony forming units (all coliforms) on the plate beneath the table when a post mortem examination was in progress in the absence of ventilation.

**AIR FLOW**

Air was found to be moving downwards at a velocity of 0.2 m/second 20 cm above the surface of the table and smoke testing showed

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**Figure 1** Post mortem room air samples: ventilation on (mean total aerobic counts).

Staff only
- Staff + cadaver
- Necropsy in progress

<table>
<thead>
<tr>
<th>Level of sample above table</th>
<th>Heads (80 cm)</th>
<th>Cadaver (20 cm)</th>
<th>Table (0 cm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Staff only</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Staff + cadaver</td>
<td>40</td>
<td>40</td>
<td>40</td>
</tr>
<tr>
<td>Necropsy in progress</td>
<td>160</td>
<td>160</td>
<td>160</td>
</tr>
</tbody>
</table>
that smoke released 20 cm above the table was readily drawn in towards the air inlets and disappeared within 10 seconds. Smoke produced at the level of the table was contained within the perimeter of the table. At the level of the table the mean air velocity was 2.240 cm/second where there were three rows of holes, 1.928 m/second where there were two, and 3.383 m/second over the drainage ports giving a mean air extraction rate of 0.644 m/second. A possible reason for the measured value being higher than the figure specified in the design is that, in use, the working surface of the linked dissection table is 40% covered by a cutting-up board and organ tray.

Discussion

The potential benefits of a down-draught ventilation system in the post mortem room would appear to be two-fold: a reduction in the ambient concentrations of formalin; and other volatile substances, and the provision of some protection against the microbiological hazards of post mortem procedures.

Eradication of unpleasant odours, as well as objective evidence of reduced concentrations of formalin has already been demonstrated during use of ventilated specimen tables of similar design.11 This has been borne out by the considerable experience with the full AFOs table system since its installation at the Royal Free Hospital, and this aspect alone is felt by the users of the system to be of considerable benefit. Until now, however, there has been no evidence that the system will reduce the numbers of micro-organisms associated with post mortem procedures. A previous study examined airborne micro-organisms in a variety of different post mortem rooms including one facility with two Plum down-draught mortuary tables. There was no significant effect on the bacterial content of sampled air, but no attempt was made to examine the dissemination of organisms from within the "air curtain".12

A recent detailed study of post mortem rooms using an identical sampling method10 to ours showed very little difference in the counts obtained in rooms with different air-change frequencies (including those with no mechanical assisted ventilation). Indeed, the authors noted that the total counts related more to the number of persons present than to the procedure performed. Our study, likewise, showed little difference in the total counts whether the cadaver was present or not, or whether or not a procedure was under way. The use of the table ventilation did produce a two or three-fold reduction in total counts at all levels above the table, but it is doubtful whether such a reduction in inoculum would prevent transmission of infection even if these organisms had all been contributed by the cadaver.

The organisms isolated when the ventilation was in use were almost all skin organisms (coagulase negative staphylococci, Bacillus spp and Micrococcus spp), confirming Babbs's findings and providing further evidence that staff organisms were the major contributor to the total counts. The use of the cranial saw might be expected to generate relatively high velocity particles which could escape the ventilation "curtain". But, the ventilation appeared to prevent passage of all tracer organisms except at cadaver height (the point at which the velocity of particles at the "curtain" would be expected to be highest).

Overall, the system achieved a two to three log fall in counts, suggesting a significant protective benefit for post mortem room staff. It is possible that similar results could be achieved by using a saw with integral extraction ventilation, but such a unit introduces further equipment to the post mortem room which is itself difficult to clean.

We have found these tables to be no more difficult to clean than other types of stainless steel table. During use the constant washing of the sump keeps the under surfaces clean and at the end of the post mortem examination the simple design facilitates the final cleaning.

No attempt was made in this study to examine the effect on viruses or mycobacteria, but it is reasonable to speculate that parti-
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The transmission of these organisms should be reduced in a similar manner by this system. There is no conclusive evidence that either hepatitis B or HIV-1 infections have caused infection in post mortem room workers via the airborne route, but there is support for transmission of pulmonary tuberculosis to post mortem room staff via this route. Fortunately, the numbers of staff who have been infected are small, but this fact is likely to preclude a prospective clinical study to examine the microbiological efficacy of these tables. The data obtained from this study certainly suggest potential efficacy and it does not seem appropriate for these ventilated tables to be dismissed as “unnecessary.” Whether they represent an economical means of providing low level exhaust ventilation and working surfaces will largely depend on whether the design of the post mortem suite permits inexpensive installation of the necessary ceiling and underfloor ducting.

Although the use of a ventilated post mortem table seems to provide potential health and safety benefits for staff it must be borne in mind that the most important route for transmission of blood borne infections is via penetrating sharps injuries. All attempts should be made to avoid these by adopting procedures recommended in the current Health and Safety Advisory Committee guidelines (HSAC 1991).

We are grateful for the help and cooperation given by Mr C. Marriage and Mr J. Levinski during the course of this study. We additionally thank Mr J. M. Caygill and Mr C. Bristow for providing the smoke generator.

**Book Review**


This book, which conforms to the usual high standards of IARC monographs, provides a detailed account of the biophysics of ultraviolet and solar irradiation and their biochemical actions on DNA. There is much about their intriguing biological effects on cellular responses and repair, and a critical review of their tumorigenic, local immunosuppressive, and other harmful effects—for example, in causing cataract and “solar keratosis”. Fluorescent lighting is reviewed and is regarded as an unproved carcinogenic risk for man, but improper use of sunbeds is shown to be harmful. Sun screens are briefly discussed, too.

There is rightly much concern about the risks to man of excessive ultraviolet and solar irradiation. This book provides the facts to support the drive to protective measures, and at the same time it is a valuable source of information about the neglected areas of radiation pathology and toxicology.

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**Notices**

Quality '93

Raising quality in the NHS: what progress?

11 November London

Quality '93 follows the success of Raising Quality in the NHS held in March 1992.

Quality '93 is being organised by the BMA, the BMJ, the Kings Fund, the College of Health and Quality in Healthcare. The meeting will review progress with raising quality in the NHS and also look at what’s new in raising quality. A particular focus will be on involving patients in raising quality.

The meeting is open to doctors, nurses, all health professionals, managers, politicians, researchers, policy makers, and members of the public.

For further details contact:

Pru Walters, BMA House, Tavistock Square, London WC1H 9JP.

Telephone: 071 383 6518.

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**The Royal College of Pathologists**

**MEETINGS**

Advances in Breast Pathology — Relevance to Management and Prognosis

Wednesday 17 November 1993

Diagnostic Developments in Leukaemia and Lymphoma

Wednesday 19 January 1994

Perspectives in Good Laboratory Management

Tuesday 22 February 1994

all at

The Royal College of Pathologists,
2 Carlton House Terrace,
London SW1

The meetings are open to members and non-members of the College.

Further details and application forms can be obtained from

Scientific Meetings Officer, RCPath,
2 Carlton House Terrace, London SW17 5AF

Tel: 071 930 5862 ext 24/26.

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4th Breton Workshop on Autoimmunity

Brest, 15–16 April 1994

Siögren’s syndrome (J F Bach, H M Moutsopoulos and N Talal); systemic lupus erythematosus (E M Tan, D Alarcon-Segovia and G Friou); rheumatoid arthritis (R N Maini, I M Roitt and J Sany). Official languages: English and French, with simultaneous translation.


Secretariat: Laboratory of Immunology, Brest University Medical School Hospital.

Tel: (33) 98 22 33 84.

Fax: (33) 98 80 10 76.

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Junior members receive the Journal of Clinical Pathology each month. Other benefits include reduced registration fees to attend ACP scientific meetings, all the documents required to full members of the Association including ACP News, which has a regular column for juniors, and the twice yearly summary of pathology courses included in the ACP programme of postgraduate education. Junior members have their own representative body, the Junior Members’ Group, which has a direct input to Council.

For Junior Membership apply to: The Honorary Secretary, Association of Clinical Pathologists, 221 Preston Road, Brighton BN1 6SA. (0273) 561188.

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**Errata**

The following errors appeared in:

Bacteriological evaluation of a downward necksac table ventilation system by McLaughlin et al, J Clin Pathol 1993; 46:746-9: the third paragraph of the Results section should read:

The mean counts of the tracer organism B subtilis var niger are shown in fig 3. The effect of the local air extraction was to reduce the counts from >102/1000 l air to 13/1400 l.

The settle plate counts increased . . .

On page 748:

The mean air velocity at the level of the table should read 2.240 m/second and not 2.240 cm/second as currently printed.