The Mutagenicity of Electrocautery Smoke

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Careful analysis of electrocautery smoke produced during breast surgery has found organic compounds that are unidentifiable with current analytical techniques. The purpose of this study was to determine the potential mutagenicity of the smoke produced by the electrocautery knife during reduction mammoplasty. Multiple air samples were collected in the operating room during two reduction mammoplasty procedures. Airborne smoke particles were tested for mutagenic potential in both tester strains of Salmonella typhimurium (TA98 and TA100) using the standard Salmonella microsomal test (Ames test). All testing was performed by the Hazard Evaluations and Technical Assistance Branch of the National Institute of Occupational Safety and Health.

The smoke produced with the electrocautery knife during reduction mammoplasty was found to be mutagenic to the TA98 strain. The Ames test, an established technique for evaluating the mutagenicity of a substance, was convincingly positive for the smoke collected during the breast surgery. Whether the smoke represents a serious health risk to operating room personnel is not known. Development of techniques to limit electrocautery smoke exposure in the operating room appears to be needed, and surgeons should attempt to minimize their exposure.

Occupational hazards of health care workers have recently received a great deal of attention. Understandably, exposure to contagious diseases from contact with infected patients has generated the most interest. The operating room can no longer be considered a safe, secure environment for the health care worker. An increased awareness of the potential health risks to the operating room team is needed.

Most airborne particles are not effectively filtered or evacuated by present operating room equipment. Surgical masks filter only large airborne particles; evacuation and laminar airflow systems are employed sporadically. Safety efforts have been limited to reducing anesthetic gas exposure. Little energy has been directed toward identifying and limiting ambient health risks to the operating room team.

Visible smoke is produced when the electrocautery knife is employed for tissue dissection and vessel cauterization. The smoke can have a noxious odor and can often be a source of irritation to the surgeon and surgical assistants. Little is known about the composition of the smoke produced by the electrocautery. A previous study failed to identify known carcinogens in the smoke. However, the majority of the smoke produced by the electrocautery knife during reduction mammoplasty is unidentifiable by present analytical techniques.¹

Reduction mammoplasty has become one of the more common surgical procedures performed by plastic surgeons, with a recent estimate placed at 35,000 yearly. Use of the electrocautery knife for breast tissue dissection and vessel hemostasis has become popular. Should the smoke produced by the electrocautery knife contain potentially harmful elements, a health risk to a large number of health care workers may exist.

A request was made to the Hazard Evaluations and Technical Assistance Branch of the National Institute of Occupational Safety and Health (NIOSH) for an evaluation of the smoke exposure risk to operating room personnel. This report concerns the on-site evaluation by NIOSH of the mutagenic potential of the smoke produced by the electrocautery knife during routine reduction mammoplasty.

**Materials and Methods**

Multiple air samples were collected in the operating room during two routine reduction
mammaplasties in which the electrocautery knife was used exclusively for breast tissue dissection and excision. The inferior pedicle technique was employed to support the nipple-areola complex. Injection of the breast tissue with an epinephrine solution was not done. The ages of the two patients were 51 and 16 years, respectively. Both procedures and on-site testing were performed at the Bryn Mawr Hospital.

Air samples were collected and analyzed by the Hazard Evaluations and Technical Assistance Branch of NIOSH. Technical information regarding the testing and results of these studies are available in NIOSH Report No. HETA 85-126.2

Sample Collection and Extraction

Airborne particles were collected on glass-fiber filters (type A/E, 4-in diameter) using Hi-Vol pumps (General Metal Works, Cleves, Ohio 45002) at flow rates between 17 and 24 ft³/min. Filters were changed if the flow dropped below 17 ft³/min.

Samples from the first patient were extracted with 150 ml methylene chloride (DCM) and then with 150 ml acetone plus methanol (A+M). Samples from the second patient were divided (because of the large quantity of particles). Half was extracted in the same manner as with the first patient, and the other half extracted with an XAD-2 resin column. Each extract was filtered and concentrated to a final volume of 0.45 and 0.3 ml in dimethyl sulfoxide for the first and second patients, respectively.

Sampling Sites

Similar sampling sites were employed for both patients. The operating room air samples (OR) were taken at locations approximately 2.5 to 3 ft above the operative field. Control air samples (CR) were taken in a separate side room approximately 0.5 ft above the floor.

Mutagenicity Assay

All extracts were tested for the mutagenic activity in both standard tester strains TA98 and TA100 of Salmonella typhimurium using the Salmonella microsomal microsuspension test.3 The system was motivated by adding increased numbers of bacterial cells (approximately 10⁸) in a concentrated suspension to airborne particle extracts with or without S-9. The motivator, S-9, was prepared from the livers of male Fischer rats pretreated with Aroclor 1254 (500 mg/per kilogram of body weight). After 90 minutes of preincubation at 37°C, the mixture was processed according to the standard Ames test protocol.4 The mutagenic activity was scored in tester cells from histidine dependence to histidine independence.

To test the stability of the smoke extract, an in situ assay, or “delay assay,” was utilized. Samples were taken from the trapping media at intervals of 2, 4, and 6 hours after surgery for this in situ assay. These samples were plated on the appropriate agar plates to determine survival and mutation frequencies. Plates were scored after incubation at 37°C for 2 days.

The smoke extract was considered mutagenic if the number of histidine revertants (His Rev) was twofold or greater than the control and showed a dose-related response. Four dilutions or concentrations (undiluted, 1:2, 1:4, and 1:8) of the smoke samples were utilized to test this dose-related response.

A known mutagen (and carcinogen), 2-aminoanthracene, was used as a “positive control” for comparison. “Control room” air was tested as well as no addition to the strains, as a “negative control.”

Results

The electrocautery smoke particles collected on the glass-fiber filters from both patients were found to be mutagenic to the TA98 strain of Salmonella. The TA98 strain did undergo alteration of its histidine dependence when exposed to the smoke extracts.

Smoke samples from the first patient (Table I) showed a positive response only with the S-9 activation. Samples from the second patient

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<thead>
<tr>
<th>TABLE I</th>
<th>Patient I: DCM Extraction</th>
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<tr>
<td></td>
<td>Histidine Revertants for Each Strain</td>
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<td></td>
<td>Particles: µg/plate</td>
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<tr>
<td>Negative control</td>
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<td>Filter control</td>
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<tr>
<td>Air from control room</td>
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<td>18</td>
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<td>37</td>
<td>8</td>
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<td>73</td>
<td>8</td>
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<td>Smoke from operating room</td>
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<tr>
<td>155</td>
<td>6</td>
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<tr>
<td>310</td>
<td>10</td>
</tr>
<tr>
<td>620</td>
<td>10</td>
</tr>
<tr>
<td>Positive control*</td>
<td>2.5</td>
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*2-Aminothracene.
(Table II) showed a slight response even without S-9 activation. The mutagenic response of smoke samples from the organic solvent (DCM) extraction of the second patient was higher than that from the XAD-2 column extraction (Table II). No significant mutagenic response was found with the in situ assay. It appears that the smoke particles are unstable and lose their mutagenic potential within 2 hours.

The TA100 strain of Salmonella typhimurium did not appear to be significantly altered by the smoke.

**DISCUSSION**

The findings of this preliminary study show that the smoke produced by the electrocautery knife during a reduction mammoplasty is mutagenic. The solvent extracts of the smoke changed the genetic makeup of the Salmonella typhimurium. The Ames test is a well-recognized and routinely utilized method of identifying compounds that are mutagenic. Standard collection and analytical techniques were employed by NIOSH in evaluating the smoke. The results of the smoke retrieved from the operative field during our two reduction mammoplasties are convincingly positive.

The difference in the results of the two patients suggests that the mutagenic potential may vary among patients. The young patient in our study had dense, firm breast tissue that produced smoke in greater concentrations and with greater mutagenic response. Additional clinical study is needed to confirm this impression.

We are obligated to discuss the significance of finding mutagens in the electrocautery smoke produced during surgery on our two patients. All smoke samples were found to contain mutagens, and the fact that only two patients were involved does not lessen the importance of these positive findings. Granted, concentrated smoke extracts were used in the testing. However, some significance must be granted to the mutagenicity potential found in the smoke samples.

Safe levels of ambient mutagens have not been determined and probably never will be. Variability among human subjects and the exposure times necessary to produce an ill effect will make it difficult to place accurate limits on exposure of known mutagens to people in an uncontrolled environment such as an operating room. Generally, the exposure time to smoke during a reduction mammoplasty is relatively short, and much of the visible smoke dissipates quickly as it rises from the operative field.

However, previous studies have demonstrated that all personnel in the operating room are exposed to a measurable amount of smoke when the electrocautery knife is used. The operating surgeon and the assistant surgeon have registered the greatest smoke exposure. Currently employed surgical masks are not designed to filter operating room air to render it suitable for safe aspiration. The small particle size of the smoke components may make it difficult to filter with a simple face mask. It may be possible to develop evacuation systems for the operative field to remove the smoke as it is generated.

Concern for overestimating health risks by extrapolation of experimental data has recently been raised. It is easy to overreact when concern for safety in the workplace is at issue. We do not wish to take our findings and make assumptions about the actual health risk of electrocautery smoke exposure during a reduction mammoplasty. Our aim is not to further alarm surgeons who must already work in an environment that, at times, appears dangerous or hostile.

It is not known whether the smoke produced by the electrocautery knife during a reduction mammoplasty represents a serious health hazard. However, the smoke extracts collected in our operating rooms during surgery were able to alter the genetic makeup of the tester Salmonella bacteria. These findings cannot be ignored when evaluating the relative safety of electrocautery...
smoke. The NIOSH report, filed after the smoke studies, made suggestions for health care workers who participate in operations where the electrosurgery knife is utilized. We concur with their conclusions and feel it is prudent to support the development of techniques to limit electrosurgery smoke exposure in the operating room. Surgeons should be cognizant of this potential health risk and should attempt to limit their smoke exposure.

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REFERENCES