Use of dichloro-difluoromethane aerosols in preparing frozen sections

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The advent of the cryostat had significantly advanced frozen section techniques. It is now the standard method for producing tissue sections in histochemistry and immunofluorescence studies and has improved the preparation of frozen sections for rapid histological diagnosis. The best method for the initial freezing of the tissues before section is, however, still controversial. This difficulty of quick freezing also arises with the traditional frozen section apparatus in small hospitals without histology laboratories, and in places where cryostats are not available.

To satisfy a local demand for a highly portable frozen section apparatus several methods to give freezing facilities with a minimum of bulk and weight were tried. Ethyl chloride was discarded as too slow and inefficient, with the added hazard of using toxic inflammable vapour near operating theatres. Carbon dioxide cylinders of 2 lb capacity were used for some years but the weight of the cylinders (12 lb each) and the need to carry two cylinders with the apparatus made us look for a less bulky method. Thermomodule units working on the Peltier principle were tried with some success but, although they permit easier sectioning, the transformer required with these units weighed 14 lb and the cold running water needed to cool the freezing head was not always conveniently available.

Eighteen months ago dichloro-difluoromethane was tried as a freezing agent, and for the last 12 months it has been used routinely, both with the cryostat in the main histology laboratory, and with the portable apparatus described here.

Dichloro-difluoromethane (Freon 12) is a gas used in the manufacture of refrigerators and is available in 12 oz aerosol cans under the trade name of Arctic Spray for use in the plumbing and electronic industries. It is non-inflammable, colourless, has little or no toxic effects except at very high temperatures, and is therefore completely safe in routine use (Thomas, 1965).

Each can provides approximately six minutes of continuous spray through a narrow polythene tube which can be accurately directed onto small pieces of tissue. In tissue up to 3 mm thick, temperatures of approximately minus 50° C can be achieved in five to 10 seconds. This rate and degree of freezing is comparable with carbon dioxide. There are no special storage containers such as

are required for liquid oxygen, liquid nitrogen, or solid carbon dioxide, and the cans can be kept at room temperature with an indefinite shelf life. Dichlorodifluoromethane is also economical and compares favourably in price with other freezing agents.

The technique for using this freezing agent is described with cryostats and with a specially designed portable frozen section apparatus.

FIG. 1. Portable frozen section apparatus. Lids opened to show packed contents.

1Available from Electronic Chemicals Ltd., 30, Notting Hill Gate, London W.11.

Received for publication 11 April 1968.
rigid wooden box (size 12 in. \times 12 in. \times 12 in.) fitted with a drop front on detachable hinges for easy access. A Leitz small freezing microtome is bolted to a rigid shelf in the centre of the box. The microtome clamp has been modified by placing a sawn-off cryostat chuck in the simple clamp provided with the microtome, insulated from the clamp by thin sheets of perspex and packed with polyurethane foam. Thus modified the stage will remain frozen for up to seven or eight minutes without further freezing. All apparatus required for dissection, fixation, cutting, and staining is packed in containers constructed out of black perspex. As can be seen in the illustrations, when the apparatus is in use the tray from beneath the shelf is laid alongside the box and heating for rapid formalin fixation is provided by an electric heating mantle. If electricity is not available a simple spirit burner could be substituted.

APPARATUS

<table>
<thead>
<tr>
<th>UPPER SHELF</th>
<th>LOWER DETACHABLE SHELF</th>
</tr>
</thead>
<tbody>
<tr>
<td>(1) Leitz small freezing Microtome (fixed);</td>
<td>(1) Heating mantle;</td>
</tr>
<tr>
<td></td>
<td>(2) slide tray;</td>
</tr>
<tr>
<td></td>
<td>(3) cork dissecting board (6 in. square);</td>
</tr>
<tr>
<td></td>
<td>(4) cover slips and mounting medium;</td>
</tr>
<tr>
<td></td>
<td>(5) instruments (knife, scissors, forceps, and slide diamond);</td>
</tr>
<tr>
<td></td>
<td>(6) water bath for floating sections;</td>
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<tr>
<td></td>
<td>(7) staining rack;</td>
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<tr>
<td></td>
<td>(8) blotting paper.</td>
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METHOD

Arctic Spray is introduced into and around the modified microtome stage and sprayed directly on the tissue in a series of 10 to 12 second bursts. Sectioning is then performed in the usual way.

REFERENCE

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*J Clin Pathol* 1969 22: 244-245
doi: 10.1136/jcp.22.2.244

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