The mechanism whereby pepsin abolishes the background staining of connective tissue has not been investigated but it is probably related to the removal of ground substance protein molecules rather than to an effect on collagen fibres per se because bacterial collagenase is ineffective in abolishing background staining.

I should like to thank Professor J. O'D. McGee, of the Department of Pathology, Gibson Laboratories, Radcliffe Infirmary, Oxford, for his help and encouragement.

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Letters to the Editor

2-Hydroxyethyl methacrylate for sectioning bone marrow

Dancey *et al.* (1976), in their paper entitled 'Section preparation of human bone marrow for light microscopy', seem to me to have been unfortunate in the difficulties encountered when using hydroxyethyl methacrylate for this purpose. Perhaps they did not read my paper on the subject (Green, 1970). We have used the method given therein for about eight years with good results, without trouble, and without the use of special equipment. I agree with their finding that Ruddell's (1971) embedding mixture containing pyridine as an activator instead of N,N-dimethyl aniline used previously (Ruddell 1967a, 1967b) is unreliable. In fact in our own trials we could not make it work. However, the N,N-dimethyl aniline mixture is entirely satisfactory for aspirated marrow particles and also for large pieces of tissue. We regularly section blocks, measuring up to 2 x 2 cm, of a wide range of tissues including decalcified (by EDTA) bone marrow slabs. The polymerisation failures experienced by Dancey *et al.* do not occur.

The most important technical hints appear to be the use of a cooling waterbath during polymerisation (which occurs within 30 min), and the fact that impregnation of large tissue blocks by monomer can be prolonged more or less indefinitely at 4°C without partial polymerisation, so ensuring complete impregnation. In sections undue staining of the plastic matrix with basic dyes can be reduced by treating the monomer with basic ion exchange resin beforehand (North, 1971).

Many valuable and relevant points about this easy-to-use plastic embedding medium for 'ordinary' hospital laboratories are given by Sims (1974), who used steel knives for sectioning, and Bennett *et al.* (1976), who present some original and stimulating information on the use of large glass knives and ordinary (rather than ultra-) microtomes. Although we use a base sledge microtome, Murgatroyd (1976) has reported good results with a rotary microtome. The great improvement in histological morphology in hydroxyethyl methacrylate sections compared with paraffin sections makes its use for selected tissues well worth while.

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References


Letters to the Editor


Macrophage electrophoretic mobility test

The report by Crozier et al. (1976) that the macrophage electrophoretic mobility test (MEM) is ineffective in the diagnosis of cancer is in full agreement with that of our group at Weston Park Hospital, Sheffield (Shelton et al., 1975). While the MEM test may be of value in the hands of highly specialist workers it is certainly of no present value outside such centres. It is interesting that a simple macrophage migration inhibition test to encephalito-genic factor gave similar results to the complex and capricious MEM test. It seems likely that further exploration and refinement of the simpler technique is a more hopeful approach to cancer diagnosis than the MEM test.

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References


Plasma paracetamol estimation

Spooner et al. (1976) claim that the method of Routh et al. (1968), for the measurement of plasma paracetamol, is relatively insensitive and more susceptible to interference from other drugs than the method of Dordoni et al. (1973). Although we agree that the former procedure is less sensitive, we feel that this is not a practical problem in the investigation of suspected paracetamol overdose. In fact, the improved sensitivity which these authors obtain limits the linearity of the method to 1000 µmol/l, necessitating some modification to the procedure in order to measure higher levels, which are often encountered in overdosage.

With regard to specificity, we have found no interference by phenobarbitone in the Routh procedure provided that the extraction is performed as originally described, that is, from ether into sodium bicarbonate solution before adjusting the pH. If this is omitted and the ether is shaken with sodium hydroxide solution instead, then barbiturate is extracted and does interfere.

Phenylbutazone is a problem in both procedures, and scanning of spectra is essential to minimise the risk of reporting false plasma paracetamol values (Wiener et al., 1976). Simply reading at one wavelength gives no indication of the presence of interfering compounds. Interference by phenylbutazone does not occur in the rapid colorimetric method described by Glynn and Kendal (1975) several modifications of which are now in use. In fact this method seems relatively free from the effects of other drugs with the exception of some sulphonamide preparations which could interfere at high concentration (Wiener, 1976).

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