Comparison of the haematoxylin basic fuchsin picric acid method and the fluorescence of haematoxylin and eosin stained sections for the identification of early myocardial infarction

HK AL-RUFAIE, RA FLORIO, EGI OLSEN

From the National Heart Hospital, Westmoreland Street, London W1M 8BA

SUMMARY A retrospective study has been carried out on the necropsy material from 30 patients who have died after a clinically diagnosed myocardial infarction. This study has been undertaken to compare the reliability of the fluorescence of infarcted myocardium when stained by haematoxylin and eosin and an adjacent section stained by the haematoxylin basic fuchsin picric acid (HBFP) method to detect early ischaemia. The results showed that the fluorescence technique is reliable, reproducible and coincides with the findings obtained by HBFP stain.

The diagnosis of myocardial infarction of less than six hours duration by histological, histochemical or biochemical methods has always been subjective. Morphological changes in the myocardium less than six hours after the clinical event of infarction are minimal, making it impossible at light microscopic level, to assess the degree of irreversible damage that has taken place. Reviews of the histochemical investigations to demonstrate the changes in fat, glycogen, succinic dehydrogenase, cytochrome oxidase, phosphorylase, uridine diphosphate, glucose-glycogen transferase, B hydroxybutyrate dehydrogenase and isocitrate dehydrogenase have been carried out.1,2 These methods can be difficult to interpret. Furthermore, fresh tissue, often impossible to obtain, is essential for histochemical analysis. This is also a prerequisite for evaluation at electron microscopic level.3

Biochemical methods, which include the changes in the Na:K ratio within the myocardium have been reported to be the most reliable techniques for the detection of early infarction.4 Other methods such as staining by acid or basic fuchsin have also been reported.5-7 These have the added advantage of permitting assessment on paraffin-embedded tissue. More recently the examination of tissue sections stained by haematoxylin and eosin by short wavelength (blue) light has also been reported.2

In the opinion of the present authors the only method which gave consistent and reproducible results in the detection of early infarction was hitherto the haematoxylin basic fuchsin picric acid (HBFP) stain.8 We have undertaken this study to compare the results obtained by this method with the fluorescence of haematoxylin and eosin stained sections.

Material and methods

A search through the necropsy records for the past three years yielded 30 cases of patients who had died after a clinically diagnosed myocardial infarction, with the period of survival varying between 30 min and 36 h. In this study of 30 cases there were 23 men and 7 women of ages ranging from 29 to 68 yr. The clinical diagnosis based upon history, physical findings and electrocardiographic evidence in all of the cases showed myocardial infarction as the most likely cause of death, no additional causes being identified at post mortem. The selection of only that material which was likely to be infarcted enabled the direct comparison of positive reactions, rather than a blind study to assess the already proven sensitivity of the two techniques. In every case the whole heart was available for study and transverse sections of both ventricles were analysed microscopically. All tissue had been fixed in 10% neutral buffered formalin, it was processed and embedded for paraffin sectioning, and cut at 5 μm thickness. Adjacent sections were stained by haematoxylin and eosin and HBFP stains.

Accepted for publication 6 January 1983
Comparison of the haematoxylin basic fuchsin picric acid method

The full methodology of the HBFP stain is quoted here. As difficulties have been experienced with this method by some workers, we have included some notes on the staining technique which we have found useful in obtaining reliable results.

1 Dewax sections and hydrate in water.
2 Stain nuclei with an alum haematoxylin (solution A), differentiate in 1% acid alcohol and blue by washing in running tap water for 5 min.
3 Stain in 0.1% basic fuchsin in distilled water (solution B) for 3 min (Note (i)).
4 Rinse slides in distilled water.
5 Rinse slides in absolute acetone.
6 Differentiate in 0.1% picric acid in absolute acetone (solution C) for 15–20 s (Note (ii)).
7 Rinse in absolute acetone.
8 Clear in fresh xylo, Inhibisol or CNP 30.
9 Mount in any synthetic resin mountant.

Ischaemic heart muscle, RBCs, collagen tissue: crimson red.
Nuclei: blue.
Other tissues and structures: yellow.
Notes
(i) The only suitable basic fuchsin found by the authors is available from Raymond A Lamb, 6 Sunbeam Road, London NW12 6JL. This solution must be freshly prepared.
(ii) At the differentiation stage it is better to treat the slides individually, face up on a slide rack, flooding the slide with the differentiating solution whilst gently rocking the slide for 15–20 s. Flood the slide again with fresh solution and then immediately wash in the absolute acetone. Finally wash the slide with fresh clearing agent. It is important to ensure that all solutions are fresh and uncontaminated.

Differentiation is completed when the red blood cells and collagen tissue in the section remain red whilst all other tissue components (other than ischaemic muscle) are yellow. If these parameters are not closely observed false positive or false negative results may be obtained. The decolouration of the red blood cells appears to be the most sensitive criterion.

The fluorescent examination was carried out on a Standard 14 Zeiss microscope fitted with an epifluorescence condenser model IV FL, illuminated with a 12 V 100 W tungsten halogen light source and viewed with Neofluar objectives. The filters used were a primary filter of 440–490 nm and a dichromatic reflector of 510 nm and a barrier filter of 520 nm. With this microscope it is possible to switch from conventional stage lighting to the epifluorescence illumination thus allowing for the examination of both HBFP stained sections and the fluorescence of haematoxylin and eosin stained sections with the maximum correlation.

Discussion
The first six hours postinfarction are the most difficult to diagnose due to the lack of visible morphological change or reliable histochemistry. The HBFP stain has been reported to give a positive staining reaction 30 min after coronary occlusion and remain positive up to 36 h. This technique has hitherto been the only reasonably reliable method of demonstrating early ischaemic changes. The difficulties of differentiation have led some workers to experience false positive or false negative staining, although this can be largely overcome if our modifications are observed. Careful examination of these 30 cases of clinically proven myocardial infarction by the fluorescent technique has also proved to give consistent reliable results, with extremely good correlation with the HBFP method. Some time needs to be spent on known positive material to appreciate the sometimes subtle shades of fluorescence such as those between yellow to red/brown with hazy border (infarcted muscle) and red/brown with a clearly demarcated border (normally perfused muscle), but no mistake can be made with the bright yellow fluorescence when it is present. Waviness as described by Bouchardy and Majno* was observed in some of the cases of both groups. These areas were invariably positive by both techniques. However, in other instances the wave pattern was absent in positive areas. This suggests that the change of waviness is not always reliable as an indication of ischaemia. The animal experiments of Lie et al** have shown that the HBFP method becomes positive after the first 30 min from occlusion. Whilst there are no animal studies to show exactly at which point the fluorescent method gives positive results, some of our patients had a clinical history of no more than 30 min from the onset of symptoms to death. These cases still gave unequivocally positive results.

It is concluded from our results that both haematoxylin and eosin fluorescence and the HBFP stain are reliable, reproducible methods of demonstrating early myocardial infarction.
Comparison of the haematoxylin basic fuchsin picric acid method

For those workers, who dislike or mistrust the results obtained with the HBFP stain, comparison with the fluorescent method would help to form a conclusive diagnosis.

References


Requests for reprints to: Dr EGJ Olsen, National Heart Hospital, Westmoreland Street, London W1M 8BA, England.