which takes a minimum of 46 hours. The implications of this rapid fixing/decalciﬁng protocol are that parafﬁn wax sections of a trephine biopsy specimen can be ready to examine the day after the biopsy specimen is taken. In practice, bone marrow trephine biopsy specimens can be processed after as little as three hours in isotonic TCA, with good results. However, to be certain of adequate ﬁxation we stipulate a minimum ﬁxation time of ﬁve hours.

Morphological preservation at 80 hours was better using isotonic TCA. Trichloroacetic acid is known to increase tissue volume, but the addition of NaCl to form an isotonic solution seems to reduce this effect. If there is a risk of excessive exposure to TCA, such as may occur if a biopsy specimen is left in TCA from Friday over the weekend to Monday, it is advisable to use an isotonic solution to prevent unwanted morphological deterioration.

Processing bone marrow trephine biopsy specimens in isotonic TCA not only cuts down on conventional processing time but also results in superior morphological preservation than that achievable using formalin ﬁxation. Shrinking artefact is reduced and nuclear detail enhanced thus more closely approximating the standard of plastic sections.


Two techniques for electron opaque staining of elastic ﬁbres using tannic acid in fresh and formalin ﬁxed tissue

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Abstract

Two electron microscopic staining techniques, one using tannic acid-glutaraldehyde as a ﬁxative, and the other using tannic acid-uranyl acetate solution as a stain on ultra-thin sections of glutaraldehyde ﬁxed material, were directly compared for elastic ﬁbre staining on several human and animal tissues. Various concentrations of tannic acid were compared using both techniques. The two techniques were also compared on formalin ﬁxed tissues.

The use of tannic acid-uranyl acetate solution as a stain on processed tissue is by far the more consistent technique and achieves equally good results on glutaraldehyde or formalin ﬁxed tissue.

It is suggested that the use of the term tannic acid technique/method should be reserved for this particular method to achieve a meaningful interpretation of results in scientiﬁc papers.

Elastic ﬁbres are components of the extracellular matrix and are present in most organs. They are synthesised by connective tissue cells.1 Morphogenesis of normal and abnormal elastic ﬁbres in adult and human fetal lung has recently been described using two different electron microscopic techniques2 3 involving the use of tannic acid, along with other methods.4 “The exact mechanism involved in the interaction between tannic acid and elastin is unknown, although Hayat has pointed out some possible mechanisms.5 These include ionic interactions between carboxylate anions from tannic acid and cationic sites on the protein. Alternatively, hydrogen bonding may be involved. At an alkaline pH there may be binding of anionic oxygen to a cationic group such as ammonium in elastin.6 Kageyama and colleagues,7 using a modiﬁcation of the original Kajikawa technique,8 have in fact shown that the maximum tannic acid staining of elastic ﬁbres is in the pH range 7.0–9.0. The claim that tannic acid enhances staining of connective tissue elements including elastic ﬁbres9 led us to compare these two tannic acid techniques9 10 to see whether they could be used interchangeably. The two methods were applied to both formalin and glutaraldehyde ﬁxed tissues.
Methods

Rat aorta and lung blocks and human fetal lung and kidney blocks were fixed in 3% glutaraldehyde in 0.1 M cacodylate buffer (pH 7.2) with the addition of 1%, 0.5%, or 0.25% tannic acid. Comparable blocks from these tissues were fixed in 3% buffered glutaraldehyde alone. These were post-fixed in 1% osmium tetroxide, dehydrated, and embedded in Araldite. Ultra-thin sections from glutaraldehyde fixed tissue were mounted on copper grids and stained with freshly made tannic acid-uranyl acetate solution, prepared as described previously. The sections were stained by floating them on single drops of this solution for 10 minutes, and after a thorough rinse in distilled water they were dried and stained with lead citrate for five minutes. The ultra-thin sections from tannic acid-glutaraldehyde fixed tissue were routinely stained for one hour in saturated alcoholic uranyl acetate followed by five minutes in lead citrate.

In the second part of this investigation 10% neutral buffered formalin was substituted for glutaraldehyde for comparison of the two techniques.

Results

Use of tannic acid-uranyl acetate on routinely fixed tissue gave precise and highly specific localisation of elastin aggregates and elastic fibres in all the tissues examined. Figures 1A and 2A show elastic fibres in the rat bronchus and the wall of the aorta respectively. The results were consistent with excellent reproducibility, and background non-specificity was not a problem. Substitution of 10% neutral buffered formalin for glutaraldehyde did not alter the outcome. Figure 3 shows clearly delineated inner elastic lamina in the vessel wall of a formalin fixed human fetal lung.

Figure 1A  Electron micrograph showing dense staining of the elastin in the wall of the rat bronchus (tannic acid-uranyl acetate stain).

Figure 1B  No staining of the elastin in a comparable area in the wall of the rat bronchus (tannic acid-glutaraldehyde fixation technique).

Figure 2A  Dense staining of the abundant elastin in the wall of the rat aorta (tannic acid-uranyl acetate stain).

Figure 2B  No demonstrable staining of the elastin in the wall of the rat aorta with tannic acid-glutaraldehyde fixation technique.
Our main criticisms of the tannic acid-glutaraldehyde fixation technique are that (i) the results are poor and inconsistent; and (ii) the non-specificity of the method can cause elastin apparently to localise in some areas and not in others, thereby facilitating mistakes. Therefore, we do not recommend its use as a technique for visualising elastic fibres.

We also found that substituting neutral buffered formalin for glutaraldehyde did not alter the result with either method. This suggests that routinely formalin fixed tissue can potentially be used for electron microscopic observation of elastin development in fetal organs and also for studying elastosis in soft tissue pathology such as granular cell tumours.

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