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K. C. Hodde  
*Academic Medical Center*

D. A. Steeber  
*University of Wisconsin*

R. M. Albrecht  
*University of Wisconsin*

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ADVANCES IN CORROSION CASTING METHODS

K.C. Hodde, D.A. Steeber and R.M. Albrecht

Laboratory for Experimental Surgery,
Academic Medical Center,
1105 AZ Amsterdam, The Netherlands.

*Department of Veterinary Science,
University of Wisconsin,
Madison, WI 53706, USA.

(Received for publication June 20, 1990, and in revised form July 26, 1990)

Abstract

This paper briefly discusses the concept of corrosion cast preparation (primarily of blood vessels), the use of the scanning electron microscope (SEM) to study these casts and the observations which can be made, together with the merits and the limitations in various applications. A number of reviews and surveys are quoted in which the different injection media, injection methods, animal preparations and corrosion procedures are described. A new procedure of cleaning the corrosion casts with sodium hydroxide and Triton X-100 is described. The observations which can be made are listed and illustrated both on the cellular level as well as in organ systems as a whole. The discussion centers around some common misconceptions, the feasibility in various applications and the limitations of the method.

The conclusion is that the method has proven to be useful especially in conjunction with other methods. Moreover, while the concept of the method may be very straightforward the approach and the interpretation often need careful consideration and might not be as straightforward as one tends to expect from the simple sounding principle.

Key Words: Vascular Corrosion Casting; Scanning Electron Microscopy; Corroding Procedures; Blood Vessels; Lymph Node.

Introduction

Soon after the scanning electron microscope (SEM) was available commercially, the long existing and extensively used concept of corrosion casting was introduced in the field of scanning electron microscopy. Corrosion casts are produced by filling an internal luminal system or space with a liquid medium which becomes solid in situ. The surrounding tissue is then corroded and removed. The resulting replica is then dried, rendered conductive and examined in the SEM. In the first publications synthetic rubber (Nowell et al., 1970) and acrylic resin (Murakami, 1971) were used. Various modifications have led to the now routinely used low viscosity proprietary and non-proprietary rubbers and resins.

A number of surveys have reviewed the literature on corrosion casting. Because the number is still restricted and because each group of authors has specific areas of interest it is advisable to refer to each of these to get an overview. A guiding selection could be: Christofferson and Nilsson, 1988; Hodde and Nowell, 1980; and Lametschwandtner et al., 1984 and 1990. The last two mentioned references contain complete bibliographies.

A number of papers have dealt with the casting materials in particular: Murakami, 1971 (first resin replica); Frasca et al., 1978 (very low viscosity latex); Gannon, 1981 (very low viscosity resin); Hanstede and Gerrits, 1982 (araldite as injection medium); Murakami et al., 1984 (hydrophilic resin), whereas others have established physical properties of the media (Weiger et al., 1986b) and compared results of the various media (Christofferson and Nilsson, 1988).

Some of the above authors and others have dealt with the method as such: Schraufnagel and Schmid, 1988a and 1988b; Christofferson, 1988; Schraufnagel, 1987; Lametschwandtner et al., 1984 and 1990; Castenholz, 1983 (light microscopy of cast in situ in fixed tissue); Castenholz et al., 1982 (very complete treatment of leakage artifacts), and Hodde and Nowell, 1980 (the history of the method and discussion of different approaches).

At present there is no generally accepted procedure for the corrosion of the specimens into clean
Materials and Methods

Animals

For these studies inbred Balb/c mice between the ages of eight and twelve weeks were used.

Corrosion Casting

Mice were anesthetized with a subcutaneous injection of 0.15 ml ketamine hydrochloride (100 mg/ml) containing Rompun (0.25 mg/ml). The thoracic cavities were opened and the liver was dissected out for subsequent corrosion. Casts were formed by manually injecting 10 ml of a 1:1 mixture of Mercox CL-2B® to methylmethacrylate monomer (final viscosity of 3-4 cps) containing 0.16 g catalyst/ml of mixture (all from Ladd, Burlington, VT). Polymerization was completed by incubating the casts for 12 hour intervals, and graded and photographed at 24 hour intervals as described above.

SEM Preparation

After corrosion the casts were thoroughly rinsed. The left ventricle was punctured and a polyethylene cannula (Intramedic PE 100) was inserted into the aorta and held in place with a silk thread. The serial blocks were examined under a dissection microscope. The right atrium was opened to serve as an efferent port. The animals were then perfused manually with 20 ml of a modified Krebs buffer, pH 7.4, containing calcium (1.5 mM), dextrose (0.17%), heparin (10 U/ml), sodium nitrite (3 mM) and BSA (1%) warmed to 37°C. For unfixed samples, mice were further perfused with 0.1 M phosphate buffer (pH 7.4) using a Harvard infusion/withdrawal pump (South Natick, MA) at a rate of 5 ml/min for 15 minutes. When fixed samples were desired the 0.1 M phosphate buffer was supplemented with glutaraldehyde (0.5% final concentration) and the animals perfused as above. Following perfusion, all animals were cast by manually injecting 10 ml of a 1:1 mixture of Mercox CL-2B® to methylmethacrylate monomer (final viscosity of 3-4 cps) containing 0.16 g catalyst/ml of mixture (all from Ladd, Burlington, VT). Polymerization was completed by incubating the animals in a 50°C water bath for 2 hours. The desired tissues were dissected out for subsequent corrosion.

Corrosion

The various corrosion treatments used on the cast tissues (either whole or dissected pieces) were as follows:

Sodium hydroxide. Pieces of unfixed liver were placed in solutions of 5, 6, 7, 7.5, 8, 9, 10, and 40% sodium hydroxide and were compared for their efficiency of tissue corrosion. Tissues were corroded in petri dishes in an incubator at 45°C. All samples were placed into fresh solutions at 12 hour intervals. The samples were evaluated at 24 hour intervals using a dissecting microscope. At this time each sample was awarded a number from 0-4, representing the degree of clearing or digestion that had occurred (a value of 4 was awarded to a cast that appeared to be 100% cleared of tissue). Light micrographs were also taken at this time.

Sodium hydroxide with Triton X-100. Unfixed whole kidneys were placed in 7.5% sodium hydroxide solution containing either 5% or 20% Triton X-100 (v/v; Sigma, St. Louis, MO). The samples were corroded in petri dishes at 45°C. The samples were placed into fresh solutions at 12 hour intervals, and graded and photographed at 24 hour intervals as described above.

Fixed versus unfixed tissue. Pieces of fixed and unfixed lungs were corroded in 7.5% sodium hydroxide for 24 hours then placed into 5% Triton X-100 for 24 hours. All incubations were carried out at 45°C. The samples were placed into fresh solutions at 12 hour intervals, and graded and photographed at 24 hour intervals as described above.

Results

The concentration of sodium hydroxide used in the clearing process was found to have an effect upon the efficiency of corrosion. The 40% solution showed minimal tissue corrosion even after nine days, perhaps due to protein cross-linking and lipid saponification. This was especially true of tissues which contained or were surrounded by large amounts of fat (e.g. lymph nodes within fatpad). All concentrations of sodium hydroxide in the range of 5-10% appeared
detergent. Figure 1 shows the increased digestion of liver tissue that occurs in 7.5% sodium hydroxide (Fig. 1a) as compared to 40% sodium hydroxide (Fig. 1b) after 4 days of corrosion.

Both alternate 24 hour washes in 7.5% sodium hydroxide followed by Triton X-100 and the direct addition of Triton X-100 detergent to the sodium hydroxide (even though the Triton X-100 was not completely soluble in the sodium hydroxide) were effective in reducing the time to clear the casts of tissue. The 5% Triton X-100 solution was found to be equally effective as the 20% solution. Total clearing of the casts was often achieved within 48 hours. Sonication did not appear to damage the casts and was effective in cleaning the small amount of residual material which is often present from the casts.

Recently it has been described by Schraufnagel and Schmid (1988) that light microscopy of wet samples is not the optimal means of assessing the extent of corrosion. Our results further substantiate this claim. The kidney samples shown in figure 2 appear as though the corrosion had progressed differentially when the sodium hydroxide was supplemented with 5% (Fig. 2a) or 20% (Fig. 2b) Triton X-100 after nine days of treatment. When the same samples are observed by scanning electron microscopy after drying they are both seen to be completely cleared of tissue (Figs. 3a, 3b respectively). We attribute the apparent difference noted by light microscopy to be due to a difference in the number of vessels per unit area (i.e., vessel density) in the casts.

We were unable to detect any difference in the corrosion time of fixed versus unfixed tissue. Figure 4 illustrates lung tissue that had been cast without prior fixation (Fig. 4a), or fixed by perfusion with 0.5% glutaraldehyde prior to casting (Fig. 4b). After corrosion in 7.5% sodium hydroxide for 24 hours, followed by treatment with 5% Triton X-100 for 24 hours, the casts of both fixed and unfixed tissues are observed to be completely cleared of tissue. We noted similar results with all tissues examined.

Typical tissue remnants are observed surrounding a cast vessel following incomplete corrosion (Fig. 5). This pattern is distinctly different from that of resin leakage as shown in figure 6. In this case typical ring-shaped profiles surrounding the cast vessel are seen (see Castenholz et al., 1982 for an extensive discussion on the subject). Veins and arteries show different and consistently recognizable imprint patterns from the endothelial lining (Miodonski et al., 1976; Hodde et al., 1977) (Figs. 7-10). These patterns can be distinguished from other features such as trapped and/or attached blood cells (Fig. 11). The typical round versus oval endothelial cell nuclei imprints and the polygonal versus oval endothelial cell border imprints retain their shapes even in a contracted or distended condition (Figs. 9, 10 respectively).

The endothelial cell cushion configuration described previously by many authors (Wagenvoort, 1954; Fourman and Moffat, 1961) show up very clearly in cast preparations of arterial branching sites as demonstrated in figure 12. These have also been described by Hodde, 1981a; Casellas et al., 1982; Kardon et al., 1982 and Murata et al., 1982. Fourman and Moffat (1961) concluded from in vivo observations that the cushion profile protruding into the lumen samples the cell-rich axial stream, thus producing "cell-skimming."

Arteriovenous anastomosis, when filled with this method, can be established beyond any doubt. Figures 13 and 14 show an example in the nasal circulation of the rat. The direct arteriovenous connections are visible (Fig. 13) with diameters exceeding those of capillaries at least five-fold. Endothelial cell imprint patterns of arteriovenous anastomoses display an intermediate appearance from that of arteries and veins (Fig. 14).

Another pattern that is typically found in the lymph node vasculature is that of the high endothelial venules (Figs. 15, 16). These venules can be identified in the cast by the characteristic pattern of deep impressions which match the dimensions of the endothelial cells (Steeber et al., 1987; Castenholz, 1990). Retrogradely filled vessels show the shut venous valve contours as shown in figure 17. This feature has

Figure 5: SEM of cast blood vessel surrounded by incompletely digested tissue remnants. Bar = 10 µm.

Figure 6: Idem, surrounded by plastic rings resulting from leakage of the injection medium through the vessel wall (reprinted from Hodde and Nowell, 1980). Bar = 20 µm.

Figure 7: Cast of a vein. Round imprints of endothelial cell nuclei (foreshortened in perspective) and cell border imprints forming polygonal patterns are seen. Bar = 10 µm.

Figure 8: Cast of an artery. Endothelial cell nuclei imprinted as oval shapes oriented in the length axis of the vessel. Cell borders form similar patterns which are seen as grooves surrounding the nuclei in a parallel fashion. Bar = 10 µm.

Figure 9: Cast of a constricted vein in lung parenchyma with band-shaped constrictions. Nuclear imprints are still recognizable beyond doubt as venous. Bar = 20 µm.

Figure 10: Cast of a rat stapedial artery. Left half: part of vessel cast previously running through Stapes. Right part: free, distended part. Endothelial cell nuclear imprints and cell border imprints are still oval in shape. Bar = 100 µm.
been used to check the (in-) competence of experimentally damaged valves (Van Bemmelen et al., 1986)(Fig. 18).

For some studies it is necessary to keep skeletal parts intact as a reference to the spacial distribution of the blood vessels. In figure 19 this was obtained by enzymatic digestion in a detergent for one week at 37°C. The viscosity of the resin injected was such that only the arterial part (Fig. 20) or the venous part was filled and thus further dissection was not necessary.

The complete blood vasculature of a whole organ or part thereof can be demonstrated with this method of corrosion casting (Figs. 21-28) either with little dissection, e.g. incisor teeth (Hodde et al., 1983) (Figs. 21, 22) and pineal gland (Figs. 23 and 24) or with much time-consuming preparatory dissection, e.g. rat hypophysis (Figs. 25, 26) and the rat choroid plexus of the lateral ventricle (Hodde, 1979b; Weiger et al., 1986a) (Figs. 27, 28).

Discussion

As is obvious from the extensive bibliography (Lametschwandtner et al., 1984 and 1990) the corrosion casting method used with SEM has been applied in various areas to address questions of "functional anatomy" in developmental anatomy (Caduff et al., 1986), hormone transport in the Pineal gland, (Hodde, 1979a), hypophysis (Bergland and Page, 1978; Hodde, 1981b), nasal circulation and temperature regulation (Hodde, 1986), hypertension (Hodde et al., 1984), immunology (Steeber et al., 1987), pathophysiology of venous valves (Van Bemmelen et al., 1986), pharmacology (Olson, 1980), vasoproliferation (Burger et al., 1984; Christofferson, 1988) and neovascularization (Grunt et al., 1986; Walmsley et al., 1987; Christofferson, 1988). Closed systems have also been studied with this method such as milk ducts (Shenkman et al., 1985) and bile canaliculi (Yamamoto and Phillips, 1984; Gaudio et al., 1988; Schellens et al., 1988).

Questions concerning the degree to which the vascular corrosion casts reflect the in vivo situation, before the injection of medium, have been raised because vasomotor activity has been demonstrated to occur with this method (Olson, 1980; Motti et al., 1987). These questions are difficult to answer at this time because many factors may play a role such as the injection pressure and the subsequent intraluminal pressure, the viscosity, the shear stress of the medium with the wall of the vessel, and the direct vasoactive influence of the medium upon the vessel wall. Prefixation of the blood vessels might preclude a number of these factors but has some unknowns of its own in terms of vasomotor reaction to the fixative (Beringer, 1979; Smith and Reese, 1980; Bachofen et al., 1982; Walter et al., 1983; Wisse et al., 1984).

One merit of the corrosion casting method is the fact that the cast shows the real 3-D spatial measurements since the injection medium becomes solid in situ. Although most media shrink measurably (Weiger et al., 1986b) this does not cause volume shrinkage of the preparation in toto. This has to be kept in mind when comparing the results with measurements from fixed and processed (dehydrated, CPD etcetera) tissue with 50% shrinkage or more (Boyde and Macconnachie, 1979).

Sometimes the study of vascular cast preparations gives much information with a small effort in time and energy; mostly however, this is not the case and patience becomes another important resource.

References


Figure 11: Cast of a vein in the rat nasal mucosa. Trapped and/or attached blood cell imprints recognized as such. Bar = 10 µm.

Figure 12: Cast of a branched artery, flow direction was from right to left. Arterial endothelial nuclear and cell border imprints are seen. At branching sites imprints or arterial endothelial cell cushions are seen as grooves deepening into the lumen of the parent vessel towards the left. Bar = 50 µm.

Figure 13: Cast of rat nasal septum vessels. Top: arteries (A). Bottom: veins (V). Middle: arteriovenous anastomoses (AVA). Bar = 50 µm.

Figure 14: Detail of previous figure, note the shape of the endothelial cell nuclear imprints are round and elongated cell border imprints are seen. Bar = 10 µm.

Figure 15: Scanning electron micrograph of a freeze-fractured mouse lymph node showing the lumen of a high endothelial venule (HEV). The prominent endothelial cells (E) are clearly seen. Bar = 5 µm.

Figure 16: Corrosion cast of a high endothelial venule (HEV). The large bulging cuboidal endothelial cells, as shown in figure 15, produce a characteristic and consistently recognizable pattern which can be used to identify the vessel as a HEV. Bar = 10 µm.


Figure 17: Cast of a retrogradely filled vein showing the profile of a shut bicuspid valve with recognizable blood vessel lumenal surface imprinted on the surface of the cast. Bar = 50 µm.

Figure 18: Cast of a retrogradely filled vein with an experimentally caused incompetent bicuspid valve profile on the right side (compare to figure 17). Bar = 10 µm.

Figure 19: Light micrograph of a cat skull (ventral aspect), with the arterial part of the carotid rete only, injected with high viscosity resin. Tissue corroded completely but the inorganic component of the bone remaining intact. Bar = 10 mm.

Figure 20: Cast of the arterial part of a cat carotid rete. Left: maxillary artery (branch of external carotid). Right: rete vessels joining the internal carotid artery. Bar = 500 µm.

Figure 21: Rat mandibular incisor tooth. Cast of the complete vascular bed of the enamel organ (ventrolateral aspect). Incisor end to the left. Bar = 1 mm.

Figure 22: Detail of previous figure (ventral aspect). Supplying arteriole (A) with particular branching pattern feeding into the capillary network of the enamel organ. Indicator A situated at the site of arterial passage through the mandibular foramen. The (V) indicates part of the extensive overlying venous plexus. Bar = 100 µm.


Figure 23: Cast of the rat pineal gland vascular bed which has been dissected free from the surrounding brain vessel cast (lateral aspect). Note the prominently visible draining veins, connected with the left lateral vein (V), which in turn connects with the Confluent Sinuus (CS). Bar= 500 μm.

Figure 24: Detail of the previous figure showing supplying arteriole (A) and draining venule (V) at the surface of the pineal gland brain vessel cast. Bar = 100 μm.

Figure 25: Cast of rat hypophyseal blood vasculature seen from dorsal. Median Eminence (ME), Infundibulum (I), Neurohypophysis (NH), Adenohypophysis (AH), and veins (v) draining the neurohypophysis exclusively. Bar = 500 μm.

Figure 26: Detail of previous figure. Short portal veins (spv) connecting the neurohypophyseal capillary bed (bottom) with the adenohypophyseal sinusoids (top). Bar = 100 μm.

Figure 27: Cast of the rat choroid plexus, left lateral ventricle. Seen in fronto-dorsal direction. Bar = 1 mm.

Figure 28: Detail of previous figure. Various capillary patterns shown: forming a flat bed in the free margin (right half picture); in villous formation (left half); and garland type (-) surrounding an artery (a). Veins are prominently seen (v). In the rat, these are never accompanied by the capillaries except for incidental drainage as seen in the center of the picture. Bar = 10 μm.


Editor's Note: All of the reviewer's concerns were appropriately addressed by text changes, hence there is no Discussion with Reviewers.