Fixation and Fixatives: Roles and Functions—A Short Review

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Abstract

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► osmium tetroxide
► picric acid

Fixation is considered as physicochemical process where cells or tissues are fixed chemically. Fixatives perform various functions such as prevention of autolysis and tissue putrefaction. Various fixative agents include formaldehyde, glutaraldehyde, osmium tetroxide, glyoxal, picric acid, and so on. A detailed search on PubMed, Google scholar, and Scopus database showed very few articles on “fixation” and “fixative.” Keeping this fact in mind, a comprehensive review on fixation and fixatives was prepared. The main aim of this review is to make pathologists and laboratory technicians familiar with the basic aspects and different types of fixatives.

Introduction

Fixation is known as a physicochemical process in which cells or tissues are fixed chemically. As a result, the tissue or cell can combat the successive treatment by different reagents with negligible disfigurement of morphology.1

An ideal fixation involves complicated progression of chemical episodes. An ideal fixative is presumed to transmit mechanical toughness to tissue so that it resists destruction due to further processing steps. It prevents the autolysis, putrefaction of tissue as well as tissue component degradation.1,2 Fixation should be able to preserve the cellular structure and tissue architecture in life-like manner.2

For the purpose of tissue processing in the histopathology, fixation of tissue is considered as necessary and essential step. Fixation amends the physio-chemical state of tissues so that it remodels the reactivity of cellular components for stains.3 Fixatives can be classified in different ways, as shown in Tables 1 – 3.

Functions of Fixative

Fixatives perform various functions. The primary function of fixatives is to prevent autolysis (enzymes attack) as well as putrefaction (bacterial attack) of tissues. Autolysis seems to be a frequent issue in enzyme-rich tissues, and rigorously autolyzed tissue does not get stained properly for microscopic examination. On the other hand, bacterial invasion can be blocked by following the strict antiseptic methods.

Another important function is conserving the association in between cells and extracellular substances. Fixatives stabilize the cell component by making them insoluble, thereby reducing the alteration by subsequent treatment and also preventing osmotic damage of tissue, which may cause shrinkage or swelling, thus preserving the cellular and tissue structure in life-like state.

Fixation also performs various other functions such as making tissue firm, so that gross cutting becomes much easier. Also, fixatives help make the tissue more easily permeable for subsequent reagents and play a role in emphasizing the dissimilarity in refractive index and thus help in increased visibility of different elements of tissue.1,2,4

Factors Affecting Fixation and Fixatives

Length of Fixation

The ideal time of fixation is experimentally determined for different types of tissue. If time period for fixation is longer,
Table 1 Classification of fixatives based on chemical composition

<table>
<thead>
<tr>
<th>Fixatives</th>
<th>Examples</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Physical agents</td>
<td>Heat, microwaves</td>
</tr>
<tr>
<td>2. Aldehydes</td>
<td>Formaldehyde, acrolein, glutaraldehyde</td>
</tr>
<tr>
<td>3. Coagulants</td>
<td>Methyl alcohol, ethyl alcohol, acetic acid</td>
</tr>
<tr>
<td>4. Oxidizing agents</td>
<td>Osmium tetroxide</td>
</tr>
<tr>
<td>5. Miscellaneous</td>
<td>Picric acid, mercuric chloride</td>
</tr>
</tbody>
</table>

Table 2 Classification of fixatives based on number of structures fixed

<table>
<thead>
<tr>
<th>Fixatives</th>
<th>Examples</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Simple fixatives</td>
<td>e.g., Formaldehyde, picric acid, osmium tetroxide</td>
</tr>
<tr>
<td>2. Compound fixatives</td>
<td>e.g., Bouin’s fluid, formol saline, Zenker’s fluid</td>
</tr>
</tbody>
</table>

Table 3 Classification of fixatives based on type of structures fixed

<table>
<thead>
<tr>
<th>Fixatives</th>
<th>Examples</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Histochemical fixatives</td>
<td>Formaldehyde, glutaraldehyde, vapor fixatives</td>
</tr>
<tr>
<td>2. Microanatomical fixatives</td>
<td>Bouin’s fluid, 10% formalin, Zenker’s fluid, formol calcium, Heidenhain’s susa, Helly’s fluid, Rossman’s fluid,</td>
</tr>
<tr>
<td>3. Cytologic fixatives</td>
<td>Champy’s fluid, glacial acetic acid, alcohol, formol saline, Carney’s fluid, Clarke’s fluid, Newcomer’s fluid, Flemming’s fluid</td>
</tr>
</tbody>
</table>

it results in over-cross-linking, and samples become brittle. If time period for fixation is short, sufficient amount of penetration in tissues and cross-linking will not occur.

For oral soft tissue, overnight fixation is sufficient.

Temperature
Temperature of fixative during fixation may affect the tissue architecture. Rate of fixation is increased with increase in temperature, but increased temperature will also increase autolysis rate. If temperature is low or decreased, the diffusion rate also decreases, which results in extended penetration time.

For electron microscopic studies, 0° to 4°C is appraise as ideal temperature.

Concentration
Fixative agents need prolonged time for fixation if concentration is low. If concentrations of fixing agent are high, it results in damaging of cellular structures as well as obliterated enzyme activities.

Different fixatives have different ideal concentration that is determined experimentally; for example, ideal fixative for oral soft tissue is formalin used in 10% concentrated solution.

Size
Tissue thickness is one of the important factors for fixation. If the sample size is large, it is unfavorable for the fixative to penetrate and reach to the deeper part of the tissue, which would result in autolysis of epithelium. Ideally 4- to 6-mm-thick specimen is best suited for complete penetration by fixatives.

Osmolarity
If osmolarity of tissue as well as fixative is same, it will prevent swelling or shrinkage of the tissue.

Various Fixating Agents Used in Histopathology

Formaldehyde or Formalin
Formaldehyde was discovered in 1859 by Butlerov. In 1889 Ttilat was the first who manufactured formaldehyde commercially as industrial reagent. In 1892, Ferdinand Blum recognized that formalin could give benefit when used as a fixative.

The most routinely used solution for fixation of tissue—10% formalin solution v/v—is nothing but an aqueous suspension of formaldehyde. In 10% neutral buffered form, formaldehyde is found to be the most commonly used fixative in pathology. Reaction between the formaldehyde and macromolecules of tissue seems to be complex. Formaldehyde reacts with nucleic acids as well as proteins, and it penetrates between nucleic acids and proteins and forms stabilized shell of nucleic acid-protein complex. As compared with other fixatives, formaldehyde causes lesser tissue shrinkage, with exceptions being acetone and ethanol. Formaldehyde seems to harden tissue more when compared with other fixatives. The lipids are conserved, but carbohydrates are not fixed by formaldehyde.

Formalin comprises 37 to 40% formaldehyde and 60 to 63% water by weight. After continuous storage for long periods, accumulations of white deposits are observed in the solution. These are the precipitates of paraformaldehyde. By storing formalin at low temperature, these white deposits can be avoided. Also, 10% methanol may be added into the formalin to minimize the polymerization reaction that produces paraformaldehyde precipitate. It also contains a slight amount of formate ions. These are obtained from Cannizzaro reaction. In this reaction, two molecules of formaldehyde react together. One molecule condenses to form methanol and second molecule gets oxidized to form formic acid. The solution is acidic in reaction because of formic acid, but acidic nature of solution can be counterbalanced with incorporation of magnesium carbonate in little proportion.

Glutaraldehyde
Glutaraldehyde was found in 1963 by Sabatini et al as particular fixative for ultrastructural researches. Glutaraldehyde comprises two aldehyde groups that are divided by three methylene bridges. Although penetration rate of glutaraldehyde is found to be slower when compared with formaldehyde,
Glyoxal is considered as an alternative fixative to formalin because it is a dialdehyde in nature. It is a bifunctional aldehyde. Its individual aldehyde groups are potentially reactive, and also cross-links can be established. Glyoxal fixed tissues may demonstrate precise cellular details, lysed erythrocytes, and disintegrated microcalcifications.26,28

According to Harke and Hoffler, glyoxal does not evaporate from the solution, and as per Henry law constant, glyoxal is virtually nonvolatile with consideration to the aqueous phase. For microwave fixation, glyoxal is the chief component used in the fixatives. It does not produce vapors at room temperature, so it is considered as less dangerous in use than formaldehyde.29 By molecular weight, glyoxal is the third smallest aldehyde after formaldehyde and acetaldehyde. It contains two carbon atoms. Glyoxal is commercially manufactured as aqueous solution that contains hydrates such as trimers, dimmers, and ring structures.30

Picric Acid
Picric acid is an example of a coagulant fixative. It forms picrates with basic protein groups, which causes coagulation. For the purpose of demonstration of DNA or RNA, picric acid fixatives are not used as picric acid and can hydrolyze nucleic acids. Also, picric acid is seen to disintegrate calcium deposits in samples. Although picric acid is not able to fix most carbohydrates and lipids, picric acid is the most advised fixative to preserve glycogen. Brighter staining is seen by picric acid fixatives.4,18,31

Picric acid is an acidic solution. Therefore, sometimes it gets washed out by alcohol. To avoid this, lithium carbonate is added, which acts as a neutralizer. Luna reported that if picric acid is present in the tissue or not completely removed, distortion or obliteration of cellular structures will occur as outcome.4,32

Ethanol and Methanol
For ethanol and methanol, fixation initiates at 50 to 60% concentration and greater than 80% concentration, respectively. They are known to be coagulants that cause protein denaturation. They cause interruption in hydrogen and hydrophobic bonding by substituting water in tissue environment, which results in change in tertiary structure.

Ethanol causes mispresentation of cytoplasmic as well as nuclear details, but sometimes it can be used for preservation of glycogen. Methanol is more commonly used for fixation of exfoliative cytology smears and blood films.18,33,34

Acetone
Acetone is another fixative agent used in histopathology. It acts as an efficacious lipid solvent that results in tissue brittleness. Apart from tissue fixation, they are primarily used as an agent for dehydration in tissue processing. Because of extremely volatile as well as flammable nature, they are not recommended for use in automatic tissue processor.18,33
Acetic Acid
Acetic acid is considered as a noncoagulative fixative agent. It acts by causing nuclear proteins coagulation. Incidentally, it stabilizes and assists to prevent nucleic acids loss. Acetic acid, when combined with ethanol, is used as an effective cytological fixative that helps in conservation of nucleic acids, but if it is used singly, it results in swelling of cells. Time required for fixation by acetic acid is less as penetration of acetic acid is faster into tissues.35

Potassium Dichromate
Potassium dichromate is also a noncoagulant fixative, but if used in combination with acid solution, it acts as a coagulant fixative. It is seldom used alone for fixation because chromate ions will link with few lipids and makes them insoluble.

Chromium seems to react with hydroxyl as well as carboxyl groups. By increasing the amount of reactive basic groups, the affinity of tissues for eosin staining will boost up. It conserves mitochondria but dissolves DNA. It is suggested that tissues that are fixed with chromate fixatives have to be washed completely in water before processing of tissues any further. This step is important as it avoids establishment of chromate suboxide that is insoluble.4,38

Bouin’s Fixative
Bouin’s fixative is known as noncoagulant picroc fixative solution and was explained by Pol Andre Bouin in 1897. Bouin’s fixative is considered as good fixative for conserving delicate as well as soft tissue structures. The major portion of Bouin’s fixative contains picric acid with little quantity of acetic acid as well as formaldehyde. In the samples that have to be undertaken in situ hybridization, Bouin’s solution cannot be used because it decreases the severity of hybridization.36-38

Acrolein
Acrolein was introduced by Luft as a primary fixative agent, and it is a three carbon αβ unsaturated monoaldehyde. Acrolein provides magnificent preservation of structural detail and conserves the virus antigenicity.16,39,40 It is also known as acrylic aldehyde. It reacts with macromolecules that result in formation of cross-links that are reversible.

Acrolein is not commonly used because it is unstable at alkaline pH and forms insoluble polymers. Acrolein is highly reactive and is found to penetrate tissues rapidly. Acrolein fixatives are chiefly used in enzyme histochemistry.2,18,33

Genipin
Genipin is a glycone derivative and is outcome of geniposide. Genipin is considered as a cross-linking agent used for biological tissue fixation. It initiates the cross-linking of free amino groups simultaneously with hydroxylysine and lysine.41,42 Genipin reacts with amino acids and results in formation of dark blue color pigment. The tissue resistance against collagenase degradation rises after tissue is fixed with genipin. It has been observed that genipin acts by forming intra- and intermolecular cross-linking with cyclic structure inside the collagen in tissues.43,44

Genipin is available as a crystalline white powder that is soluble in acetone, methanol, and ethanol. Genipin demonstrates its cross-linking properties at pH 7.4–8.5.41,42 It is a known cross-linking agent and has proved its potential in various biomedical application such as dentistry, articular cartilage tissue engineering applications, nerve regeneration, and so on.45-47

Conclusion
Fixation is considered as key step in histopathology procedure. Each and every fixative has its own advantage and disadvantage. Various different fixatives perform various functions, and various factors such as size, temperature, and osmolarity have direct effect on fixation procedure.

Note
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Conflict of Interest
None declared.

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