Current Concepts of Enzyme Histochemistry in Modern Pathology

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Abstract
Enzyme histochemistry serves as a link between biochemistry and morphology. It is based on metabolization of a substrate provided to a tissue enzyme in its orthotopic localization. Visualization is accomplished with an insoluble dye product. It is a sensitive dynamic technique that mirrors even early metabolic imbalance of a pathological tissue lesion, combined with the advantage of histotopographic enzyme localization. With the advent of immunohistochemistry and DNA-oriented molecular pathology techniques, the potential of enzyme histochemistry currently tends to be underrecognized. This review aims to draw attention to the broad range of applications of this simple, rapid and inexpensive method. Alkaline phosphatase represents tissue barrier functions in brain capillaries, duodenal enterocyte and proximal kidney tubule brush borders. Decrease in enzyme histochemical alkaline phosphatase activity indicates serious functional impairment. Enzyme histochemical increase in lysosomal acid phosphatase activity is an early marker of ischemic tissue lesions. Over the last four decades, acetylcholinesterase enzyme histochemistry has proven to be the gold standard for the diagnosis of Hirschsprung disease and is one of the most commonly applied enzyme histochemical methods today. Chloroacetate esterase and tartrate-resistant phosphatase are both resistant to formalin fixation, EDTA decalcification and paraffin embedding. Early enzyme histochemical insight into development of a pathologic tissue lesion and evaluation of function and vitality of tissue enhance our understanding of the pathophysiology of diseases. In this process, enzyme histochemistry constitutes a valuable complement to conventional histology, immunohistochemistry and molecular pathology for both diagnostic and experimental pathology.

Introduction

Enzyme histochemistry is a morphological technique applied to functional questions in histopathology (fig. 1). Enzyme histochemistry constitutes a link between biochemistry and morphology [1, 2] and provides important information complementary to conventional histology, immunohistochemistry and molecular pathology.

The father of enzyme histochemistry was A.G.P. Pearse [3, 4]. Since the initial introduction of enzyme histochemistry, reactions involving almost all enzymes known have been applied to sections of any human tissue. It was applied in the diagnosis of peracute myocardial infarction [5, 6], the diagnosis of celiac disease [7–15], of brain...
Enzyme histochemistry combines the biochemical analysis of enzyme activity with information on its topographical localization. The basic techniques of enzyme histochemistry are described in detail in the laboratory manuals of Pearse [4], Meier-Ruge et al. [38], Meier-Ruge and Bruder [40] and Lojda and Schiebler [41]. In order to briefly acquaint the reader unfamiliar with these techniques, we describe two of the most important enzyme histochemical techniques in detail.

In a dehydrogenase reaction, enzyme substrates like sodium succinate or sodium L-lactate are oxidized and a stoichiometric color indicator tetranitrotetrazolum chloride blue (TNBT) is reduced to black or blue formazan. The formazan immediately binds to local protein and permits the precise localization of the enzyme dehydrogenase in a particular tissue compartment. The enzyme histochemical reaction follows the stoichiometric principles of biochemistry. Whereas biochemistry is applied to tissue homogenates or extracts, expressing enzyme activity in turnover rates, enzyme histochemistry indicates the locus of an enzyme in the tissue section.

A second group of enzyme reactions use diazonium salt instead of tetrazolum chloride as color indicator. Enzymes stained with this kind of reaction are mainly esterases and phosphatases. The ester group or phosphate group of a naphthyl salt is split off by the enzyme reaction and the naphthyl rest couples to a diazo-salt and stains the esterase- or phosphatase-containing compartment like in a formazan color reaction [38, 41–43].

Almost all enzyme histochemical investigations are performed with frozen tissue, because most enzymes are inactivated by formalin fixation. In fact, formalin fixation is even employed to stop enzyme histochemical reactions like lactic dehydrogenase or succinic dehydrogenase reactions [40]. With the requirement of frozen tissue, transportation and storage of biopsies or surgical specimens become an issue and lead to frequently asked questions in daily routine practice. It is recommended to freeze specimens on dry ice (CO_2) at –80°C or in isopentane stored in a –25°C freezer. Liquid nitrogen carries the disadvantage of cracking freezing artifact in large tissue specimens [43]. After freezing as described, the tissue can be stored without loss of enzymatic activity for longer periods at –25°C in a small Eppendorf tube (biopsies) or plastic bag (surgical resections).

Similarly, an important technical point to stress is the minimal section thickness. It is recommendable to cut
cryostat sections at 15 μm in order to overcome the minimal enzyme activity threshold for the enzymatic starting reaction [43]. It is important to be aware that a 15-μm-thick cryostat section loses 70% of its thickness by thawing, spreading and drying, resulting in a final thickness of 4 μm, whereas an originally 4-μm-thick native cryostat section has a final thickness of 1.2 μm, so that in thin sections the enzyme reaction does not reliably develop because the enzyme activity may drop below the enzymatic starting reaction [43].

**Enzyme Histochemistry versus Biochemical Analysis**

Biochemistry employs tissue extracts or tissue homogenates. Analysis of such tissue specimens reveals a general increase or decrease in enzyme activity measured in turnover rates. In contrast, enzyme histochemistry localizes enzyme activity to cells or tissue compartments but does not provide the exact quantitative data of biochemical analysis. The capability to localize enzyme histochemical activity on a cellular level justifies the Greek prefix ‘-topo’ in the term ‘enzyme histotopochemistry’, distinguishing enzyme histochemistry from biochemistry.

Overall, there is a good correlation between histochemically judged enzyme activity per volume (area × thickness) and biochemically determined activity per milligram tissue [1].

The strength of biochemistry lies in the potential for quantification: biochemistry permits exact quantification of enzymatic turnover rates, whereas enzyme histochemistry has the disadvantage of a lack of reliable quantification, so that only semiquantitative assessment is achieved if microspectrophotometry is applied. However, the elective staining of a particular tissue structure in enzyme histochemistry permits precise morphometric size measurement by means of optic electronic morphometry equipment, e.g. the ASBA3 system [44] (compare fig. 15 and 16). The level of monochromatic light absorption permits semiautomatic calculation, e.g. of the extension of an ischemic brain infarct. Another example of morphometric analysis in conjunction with enzyme histochemistry is the determination of nerve cell size, nerve cell number and nerve cell distance of the myenteric plexus stained with lactic dehydrogenase enzyme histochemistry [40] (compare fig. 5).

Biochemistry incurs the disadvantage of tissue dilution effect by tissue surrounding the structure of interest. Chromatography, electrophoresis or density gradient centrifugation is applied to compensate for this disadvantage. Compared to enzyme histotopochemistry, the biochemical procedure is therefore time-consuming but nevertheless falls short to localize the altered structure, which is more easily demonstrated by enzyme histotopochemical means.

**Applications of Enzyme Histochemistry as a Tool in Diagnostic Pathology and Pathobiology**

Enzyme histochemistry can be used in pathohistological routine diagnosis as it has simple technical requirements. Enzyme histotopochemical demonstration of acetylcholinesterase activity in frozen colon mucosal biopsies has proven a reliable tool and is the current gold standard in the diagnosis of Hirschsprung disease today [2, 40, 45–48] (fig. 2). In particular, in the diagnosis of ultrashort Hirschsprung disease and aganglionosis limited to the internal sphincter, the so-called sphincter achalasia, the acetylcholinesterase reaction is the choice technique [49–52].

Another attractive application of enzyme histochemistry is the verification of a peracute myocardial infarction during autopsy. This can be performed macroscopic-
cally by a succinic dehydrogenase reaction in a Petri dish with succinate (13 g in 100 ml phosphate buffer at pH 7.4) and neotetrazolium chloride (100 mg dissolved in phosphate buffer). The reduced succinic dehydrogenase activity in the ischemic area of the myocardial infarction can quickly and reliably be recognized macroscopically [5, 6].

The enzyme alkaline phosphatase conveys a range of important information. On the one hand, alkaline phosphatase is eliminated by bile secretion. An increase or accumulation of alkaline phosphatase in bile capillaries of the liver indicates a disturbed mechanism of secretion [53–57] (fig. 3). Toxic lesions of the liver, infectious hepatitis [58–60] or seriously congested liver [57, 61] show similar pictures. These observations demonstrate that a simple enzyme histochemical reaction for alkaline phosphatase indicates a functional disturbance of the liver in biopsy or autoptic liver tissue. In addition to the transport function of bile capillaries for alkaline phosphatase, this enzyme represents a barrier function and is observed in structures with directed transport function. Alkaline phosphatase of brain capillaries represents the blood–brain barrier [39, 62, 63].

The alkaline phosphatase of the brush border of the proximal renal tubules is responsible for absorption of metabolites from the primary urine and is quite sensitive to toxic or ischemic injury [64–66] (fig. 4).

Of particular diagnostic value is alkaline phosphatase in the brush border of the duodenal mucosa in the diagnosis of celiac disease. Alkaline phosphatase activity is high in young adults and low in postweaning children [7]. This enzyme is a fairly sensitive parameter in the diagnosis and therapy of celiac disease. Similarly, malabsorption
syndromes show loss of alkaline phosphatase in the brush border of enterocytes [8–12]. The reappearance of an alkaline phosphatase brush border of enterocytes in duodenal mucosal biopsies is a clear indicator of compensated celiac disease. It is a parameter superior to the histological findings of villous atrophy. With the progression of celiac disease, the dehydrogenase activity of enterocytes decreases [46]. The tremendous number of publications illustrates the diagnostic and therapeutic difficulties of celiac disease [13–15].

Examination of seromuscular biopsies or resected specimens with lactic, succinic dehydrogenase and nitroide synthetase reactions permitted the morphometric characterization of hypoganglionosis as a cause of chronic constipation [67, 68]. Proximal to the aganglionic segment, often, a hypoganglionic segment is observed with scanty nerve cells and ganglia with increased interganglionic distances [40, 69] (fig. 5). The hypoganglionic segment shows a decreased acetylcholinesterase activity in the muscularis propria, indicating disturbed motility (fig. 6). Over recent years, enzyme histotopchemistry has contributed to remarkably improved understanding of the pathophysiology of chronic constipation [40].

Dehydrogenase reactions are of interest to examine the vitality of a particular tissue. In kidney biopsies, not only alkaline phosphatase is of interest but also the dehydrogenase activity of the tubulus epithelium [64–66] (fig. 4). The latter demonstrates the vitality of the kidney, which may be of interest in transplanted kidneys.

In oncology, dehydrogenase activity of a tumor demonstrates its vitality with the impact on the efficiency of cytostatic therapy or radiotherapy. It has been clearly shown that tumor tissue with high glycolytic activity is particularly resistant to radiotherapy [20, 70, 71]. High glycolytic tumor activity is revealed in enzyme histochemistry for succinic and lactic dehydrogenase (fig. 7). Enzyme histochemistry for dehydrogenase provides a quick, efficient and inexpensive orientation on the actual energy metabolism of a particular tumor and on the expected efficacy of the therapeutic strategy [21, 22]. This opens a fascinating field of research in monitoring cancer treatment. Glycolytic inhibitors, depriving cancer cells of their energy supply, are under discussion for a new class of anticancer drugs [23–25], and it has recently been demonstrated that cancer cells with high glycolytic rate [26–28] are characterized by high drug resistance [27–30].

Craniocerebral trauma and temporary ischemia is often hard to prove in a hemalum-eosin stain. Enzyme histochemistry shows a significant decrease in lactic and succinic dehydrogenase activity in Purkinje cells of the cerebellum under these conditions (fig. 8). Lesions of blood-brain barrier are characterized by focal or diffuse decrease in alkaline phosphatase in brain capillaries [72, 73] (fig. 9). This kind of brain lesions can be observed in...
boxers [16]. Only few papers in the literature deal with these particular lesions [17–19]. The brain is particularly suited for a postmortem enzyme histochemical examination, because autolytic processes start fairly late in the adult brain.

Senile plaques in the brain can be selectively highlighted by their intense acid phosphatase activity (fig. 10). Acid phosphatase is a key enzyme of lysosomal activity.

The combination of an acid phosphatase reaction with a Timm staining reveals a decrease of synapses in the surroundings of senile plaques in the hippocampus (fig. 11). Lysosomal enzymes may be responsible for the formation of senile plaques [74].
Applications of Enzyme Histochemistry in Experimental Pathology

Enzyme histochemical investigations on causes of intoxication, drug side effects or a particular disease are usually performed in laboratory animals. Enzyme histochemistry permits monitoring of a dose-dependent intoxication and time-related changes in different organs of rats, mice, dogs, cats and other laboratory animals.

Cats as twilight animals with nocturnal activity have a retina rich in rods. Such retinas have a particularly high metabolism of the visual cells and a high sensitivity to toxic side effects. The thioridazine analogue piperidylchlorophenothiazine caused night blindness and pigmentation of the retina in clinical trials with high dosage of the tranquilizer. In laboratory cats, it was possible to demonstrate that piperidylchlorophenothiazine inhibits glycolytic enzyme activity of rods (fig. 12), accompanied by an increased desquamation of peripheral rod segments. Phagocytosis and storage of these light absorbing structures from the rods in the pigment epithelium causes pigmentation of the cat retina [75]. Elucidation of this drug side effect was rendered possible by enzyme histochemical analysis.

The chloroquinone retinopathy is characterized by a toxic edema of the pigment epithelium [76, 77] (fig. 13). This implies a defect of the visual field of the retina [77], because pigment epithelium is not further able to isomerize all-trans retinine to 11-cis retinine which is necessary for rhodopsin synthesis. The reason of this localized side effect of chloroquine in the retina is its accumulation on choroid melanin and high chloroquine levels in the vitreous fluid damaging the pigment epithelium.

The development of an ischemic brain infarct by experimental occlusion of the middle cerebral artery in primates was studied with the lysosomal enzyme acid phosphatase. In this trial, the time-dependent increase in the ischemic brain infarct was studied [78] (fig. 14). A circulatory arrest in neonatal pigs demonstrated the earliest morphological changes after 90 min [78]. In primates, the release of acid phosphatase starts immediately after clipping the medial cerebral artery. The extension of the ischemic brain infarct becomes obvious 20–30 min later. Acid phosphatase permits a morphometric, time-dependent follow-up of the extension of the infarct size [78–80] (fig. 15). Morphometric measurements of alkaline phosphatase-positive capillaries in combination with the size of the ischemic infarct show that the therapeutic window for an effective infarct treatment is limited to the first 4 h with declining time-dependent effectiveness [78, 81].
Acid phosphatase offers the possibility to objectify the extension of the penumbra by morphometric techniques, which is actually much smaller than expected by X-ray investigation (fig. 16).

Performance of Enzyme Histochemistry on Formalin-Fixed and Paraffin-Embedded Tissue

Enzyme histochemistry is also frequently applied in hematology. Chloroacetate esterase is commonly used as a marker of myelopoiesis. A major advantage of this esterase is its stability to formalin fixation and paraffin embedding. Low levels of chloroacetate esterase in bone marrow smears of patients with promyelocytic leukemia are considered to indicate a bad prognosis [31]. As a nonspecific marker [32], in hematological laboratories it is routinely applied in conjunction with a series of different enzymes as β-glucuronidase, β-glucosaminidase, acid phosphatase, peroxidase and a number of dehydrogenases of the citric cycle [33].

Tartrate-resistant acid phosphatase is an enzyme localized to mature osteoclasts [34, 35]. Tartrate-resistant acid phosphatase is similarly resistant to formalin fixation, paraffin embedding and even EDTA decalcification, and is applied to visualization of osteoclasts in bone biopsies and osteopetrosis after bone marrow transplantation [35]. Serum levels of tartrate-resistant acid phosphatase are determined for monitoring bone metastases
Enzyme histochemistry as a link between biochemistry and morphology. Prog Histochem Cyberm 1976;8:1–68.


References

[36] and in patients with prosthetic hip replacement to predict periprosthetic osteopenia [37]. However, it should be stated that none of these two reactions applicable to formalin-fixed, paraffin-embedded tissue equals the dynamics and stoichiometry of classical enzyme histochemical reactions like lactic dehydrogenase, succinic dehydrogenase, alkaline phosphatase, acid phosphatase, ATPase, glucuronidase and other enzymes.

**Conclusion**

Enzyme histotopochemistry is a functional technique providing insights into the pathophysiology and pathology of a particular pathological process. It has the potential to demonstrate the vitality of cells or a particular tissue. Enzyme histochemistry serves to detect early metabolic changes in biopsies and autopsy tissue before manifestation on HE staining or immunohistochemistry. As such, it constitutes a valuable complement to other special techniques, i.e. immunohistochemistry and molecular pathology.

An exclusive diagnostic domain of enzyme histochemical analysis is the aganglioneosis of the distal rectum mucosa, which cannot reliably be diagnosed by conventional histology alone. Apart from current diagnostic applications in muscle biopsy analysis and the diagnosis of Hirschsprung disease, enzyme histochemistry provides access to a wide range of investigations in experimental pathology and toxicology as a pathophysiological supplement to conventional histology.


