

琥珀酸脱氢酶染色试剂盒(四唑盐法)

货号: G2000

规格: 60mL

保存: 2-8°C, 避光保存, 有效期 6 个月。

产品组成:

名称	60mL	保存
试剂(A): NBT 孵育液	50mL	2-8°C, 避光
试剂(B): NBT 对照液	10mL	2-8°C, 避光

产品介绍:

琥珀酸脱氢酶(succinate dehydrogenase, SDH)是此琥珀酸氧化酶系的第一个酶, 位于线粒体内。琥珀酸脱氢酶是黄素蛋白酶, 分子内含有-SH, 决定着酶的活性, 故封闭-SH 者皆可作为抑制剂。此酶活性最适 pH 为 7.6~8.5。此酶参与三羧酸循环, 在组织化学上, 常以此酶活性作为三羧循环的代表, 亦作为线粒体的标志酶之一。含此酶活性高的组织为心肌、肾小管上皮和肝细胞。此酶对固定剂敏感, 故需用新鲜组织切片。

我司琥珀酸脱氢酶染色试剂盒(四唑盐法)以琥珀酸为底物, 在酶作用下脱氢, 硝基蓝四唑(NBT)为受氢体, 接受氢后被还原为甲噁, 呈蓝紫色, 用以代表琥珀酸脱氢酶的活性。NBT 孵育液含有特殊的中间递氢体, 可使定位更加准确, 染色更加清晰。

自备材料:

蒸馏水、恒温箱或水浴锅

操作步骤: (仅供参考)

1. 冰冻切片, 厚 6 μ m, 无需固定。
2. 切片入 NBT 孵育液中, 置于 37°C 温箱, 浸染约 5-30min。蒸馏水冲洗。
3. 甘油明胶封固。

染色结果:

酶活性部位	蓝紫色沉淀
线粒体	蓝紫色颗粒

阴性对照(可选):

1. 相同切片滴加 NBT 对照液, 置于 37°C 温箱, 孵育约 5-30min, 作为对照。其余步骤同正常步骤, 结果为阴性。
2. (可选)相同切片经 10%福尔马林浸泡 30~60min, 再入 NBT 孵育液, 结果为阴性。

注意事项:

1. 本染色液适用于冰冻切片。
2. 对冰冻切片染色时, 应减少切片在室温暴露的时间。
3. 为了您的安全和健康, 请穿实验服并戴一次性手套操作。

Succinate Dehydrogenase Stain Kit(Tetrazole Salt Method)

Cat: G2000

Size: 60mL

Storage:2-8°C, avoid light, valid for 6 months.

Kit Components

Reagent	60mL	Storage
Reagent(A): NBT Incubation Solution	50mL	2-8°C, avoid light
Reagent(B): NBT Control Solution	10mL	2-8°C, avoid light

Introduction

Succinate dehydrogenase (SDH) is the first enzyme of succinate oxidase system, which is located in mitochondria. Succinate dehydrogenase is a flavin protease, containing -SH in the molecule, which determines the activity of the enzyme, so can inhibit it by blocking -SH. The optimum pH of this enzyme was 7.6-8.5. This enzyme is involved in the tricarboxylic acid cycle. In histochemistry, it is often used as the representative of the tricarboxylic acid cycle and one of the marker enzymes of mitochondria. The tissues with high activity of this enzyme are cardiac muscle, renal tubular epithelium and liver cells. This enzyme is sensitive to fixative, so fresh tissue sections are needed.

Succinate Dehydrogenase Stain Kit(Tetrazole Salt Method) uses succinic acid as substrate, dehydrogenates under the action of enzyme, nitro blue tetrazole (NBT) as hydrogen acceptor, after receiving hydrogen, it is reduced to methoxate, which is blue purple, to represent the activity of succinate dehydrogenase. NBT Incubation Solution contains special intermediate transmitters, which can make the location more accurate and the staining more clear.

Self Provided Materials

Distilled water, Incubator or Water bath

Protocol(for reference only)

1. For frozen section,cut into 6 μm thick without fixation.
2. Put the section into NBT Incubation Solution, place in 37°C incubator, and soak for about 5-30min.
3. Rinse with distilled water.
4. Glycerin gelatin sealing.

Result

Positive Site of Enzyme	Blue Purple Precipitate
mitochondrion	Blue Purple Particles

Negative Control (Optional)

Place the same sections in NBT control solution directly in 37 °C incubator for about 5-30min. The other steps are the same as the normal steps, and the result is negative.

(optional)Soak the same section in 10% formalin for 30-60min and then put into NBT incubation solution, the result is also negative.

Note

1. The staining solution is suitable for frozen section.
2. The exposure time of frozen sections at room temperature should be reduced.
3. For your safety and health, please wear experimental clothes and disposable gloves.