Introduction

The glycolytic enzyme enolase (2-phosph-D-glycerate hydrolyase) exists as three dimeric isoenzymes (α, αβ, αγ) composed of three distinct subunits: α, β, and γ. Three isoenzymes are found in human brain: α, γ-α, and γ-γ. The α-γ and α-γ-γ isoenzymes are also known as neuron-specific enolase (NSE) as these isoenzymes initially were detected in neurons and neuronendocrine cells. The NSE levels are low in health and benign subjects. Elevated levels are commonly found in patients with malignant tumors with neuronendocrine differentiation, especially small cell lung cancer and neuroblastoma.

Lung cancer is one of the most spread cancer forms with incidences about 50-100 per 100,000 population. Approximately 20% of the lung cancer is small cell lung cancer. Patients with small cell lung cancer show various proportions of α-γ-γ, α-γ isoenzyme. The determination of NSE should detect the γ isoforms with the same sensitivity. The antibodies for this particular assay are specific for the γ-subunit without cross reactivity with α or β subunits.

NSE are reported to be useful diagnostic marker for lung cancer, neuroblastoma, melanoma, seminoma(S) and in injury of central nervous system. In addition to the above, NSE are reported to be useful diagnostic marker for lung cancer and neuroblasloma. Neuroendocrine differentiation, especially small cell lung cancer, and other related diseases.

Materials and components

Materials provided with the test kits:
- Monoclonal anti-NSE antibody coated microtiter plate with 96 wells.
- Sample diluent (12 mL).
- Enzyme conjugate reagent, 12 mL.
- Lyophilized NSE reference standards containing: 0, 5, 15, 40, 100, and 200 ng/mL of NSE. 1 set.
- 50 X Wash Buffer Concentrate, 15 mL.
- TMB Substrate, 12 mL.
- Stop Solution 12mL.

Materials required but not provided:
- Precision pipettes and tips, 0.025mL, 0.05 mL, 0.10 mL, and 1.0 mL.
- Disposable pipette tips.
- Distilled water.
- Glass tubes or flasks to prepare wash buffer.
- Vortex mixer.
- Absorbent paper or paper towel.
- Microtiter plate reader.
- Graph paper.

Specimen collection and preparation

1. Blood should be drawn using standard venipuncture techniques and the serum should be separated from the red blood cells as soon as practical.
2. Plasma samples collected in tubes containing EDTA, heparin, or oxalate may interfere with test procedures and should be avoided.
3. For prolong storage of whole blood can cause release of NSE from the blood cells. If sera can not be assayed immediately, they can be stored at 2-8°C for one day or frozen at -20°C for up to 30 days prior to assay. Sample should not be refrozen. Repeated freezing and thawing is not recommended. Do not store in self-defrosting freezer.
4. Do not use hyperlipemic, hemolized, Plasma is not recommended since significant amounts of NSE can be released from the thrombocytes.
5. Avoid turbid and contaminated samples.

Test principle

The NSE Quantitative Test Kit is based on a solid phase enzyme-linked immunosorbent assay. The assay system utilizes one monoclonal anti-NSE antibody for solid phase (microtiter wells) immobilization and another monoclonal anti-NSE antibody in the antibody-enzyme (horseradish peroxidase) conjugate solution. The standards and test specimen (serum) are added to the antibody coated microtiter wells. During the incubation, specific NSE bound to anti-NSE antibody on the wells. Unbound NSE antigen is removed by washing the wells with buffer. Enzyme conjugate is then added to each well. After another incubation, unbound enzyme conjugate is washed off and the amount of bound peroxidase is proportional to the concentration of the NSE present in each sample. Upon addition of the substrate and chromogen, the intensity of blue color will develop in proportion to the concentration of NSE antigen in the samples.

Storage of test kits and instrumentation

1. Unopened test kits should be stored at 2-8°C upon receipt and the microtiter plate should be kept in a sealed bag with desiccants to minimize exposure to damp air. The test kit may be used throughout the expiration date of the kit (One year from the date of manufacture). Refer to the package label for the expiration date.
2. Opened test kits will remain stable until the expiring date shown, provided it is stored as prescribed above.
3. A microtiter plate reader with a bandwidth of 10nm or less and an optical density range of 0-3 OD or greater at 450nm wavelength is acceptable for use in absorbance measurement.

Reagent preparation

1. All reagents should be brought to room temperature (18-25°C) and mixed by gently inverting or swirling prior to use. Do NOT induce foaming.
2. Add 0.5 mL of distilled water to reconstitute the lyophilized standards. Allow the reconstituted materials to stand for at least 20 minutes. Mix gently. The reconstituted standards should be stored sealed at 2-8°C.
3. Dilute 1 volume of Wash Buffer Concentrate (50x) with 49 volumes of distilled water. For example, Dilute 15 mL of Wash Buffer (50x) into 735 mL of distilled water to prepare 750 mL of washing buffer (1x). Mix well before use.

Assay procedure

1. Secure the desired number of coated wells in the holder.
2. Dispense 25μL of standard, specimens, and controls into appropriate wells.
3. Dispense 100μL of Sample diluent into each well.
4. Thoroughly mix for 10 seconds. It is very important to have complete mixing in this setup.
5. Incubate at room temperature (18-25°C) for 30 minutes.
6. Remove the incubation mixture by flicking plate content into a waste container.
7. Rinse and flick the microtiter wells 5 times with washing buffer (1X).
8. Strike the wells sharply onto absorbent paper or paper towels to remove all residual water droplets.
9. Dispense 100µL of Enzyme Conjugate Reagent into each well. Gently mix for 5 seconds.
10. Incubate at room temperature for 30 minutes.
11. Remove the incubation mixture by flicking plate contents into a waste container.
12. Rinse and flick the microtiter wells 5 times with washing buffer (1X).
13. Strike the wells sharply onto absorbent paper to remove residual water droplets.
14. Dispense 100µL TMB substrate into each well. Gently mix for 5 seconds.
15. Incubate at room temperature for 20 minutes.
16. Stop the reaction by adding 100µL of stop solution into each well.
17. Gently mix for 30 seconds to make sure that the blue color changes to yellow color completely.
18. Read optical density at 450nm with a microtiter reader within 30 minutes.

**Important Note**

1. The wash procedure is critical. Insufficient washing will result in poor precision and falsely elevated absorbance readings.
2. It is recommended that no more than 32 wells be used for each assay run if manual pipetting is used since pipetting of all standards, specimens and controls should be completed within 5 minutes. A full plate of 96 wells may be used if automated pipetting is available.
3. Duplication of all standards and specimens, although not required, is recommended.
4. If a serum specimen contains greater than 200 ng/mL of NSE the sample must be diluted with sample diluent and re-assayed as described in the assay procedure.

**Calculation of results**

Calculate the mean absorbance value for each set of NSE reference standards, specimens and controls. Construct a standard curve by plotting the mean absorbance obtained from each reference standard against its concentration in ug per mL on linear graph paper, with absorbance values on the vertical or Y axis and concentrations on the horizontal or X axis. Use the mean absorbance values for each specimen to determine the corresponding concentration of NSE in ug per mL from the standard curve. Any diluted specimens must be corrected by the appropriate dilution factor.

**Example of standard curve**

Results of a typical standard run with optical density reading at 450nm shown in the Y axis against NSE concentrations shown in the X axis.

<table>
<thead>
<tr>
<th>NSE Values (ng/mL)</th>
<th>Absorbance (450nm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>0.010</td>
</tr>
<tr>
<td>5</td>
<td>0.195</td>
</tr>
<tr>
<td>15</td>
<td>0.418</td>
</tr>
<tr>
<td>40</td>
<td>0.928</td>
</tr>
<tr>
<td>100</td>
<td>1.980</td>
</tr>
<tr>
<td>200</td>
<td>3.309</td>
</tr>
</tbody>
</table>

This standard curve is for the purpose of illustration only, and should not be used to calculate unknowns. Each user should obtain his or her own standard curve and data.

**Expected values and sensitivity**

1. It is recommended that each laboratory should determine its own normal and abnormal ranges as to account for its environmental factors such as diet, climate etc.
2. A clinical study of the NSE Quantitative kit was conducted and results are summarized as follows: Nearly all the individuals have NSE values below 15 ng/ml (95th percentile).
3. The expect ranges are representative only, and do not necessarily reflect the ranges that will be observed in a particular clinical laboratory.

**Limitations and applications**

1. For diagnostic purposes, the NSE test results must be used in conjunction with other data available to the physician.
2. The NSE test should not be used in cancer screening and should not replace any established clinical examination.
3. Samples with NSF level above 200 ng/mL should be diluted to obtain accurate value.
4. High NSE values may be found in dialysis patients with leukaemic diseases.
5. Serum should not contain visible hemolysis since erythrocytes contai significant amounts of NSE.
6. Prolonged storage of whole blood can cause release of NSE from the blood cells.

**References**


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