Comparison of a time-resolved fluoroimmunoassay with a solid phase immunoradiometric assay for the measurement of alphafetoprotein in amniotic fluid

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Introduction

A new commercially available dissociation enhanced lanthanide fluoroimmunoassay (DELFIA) for the measurement of amniotic fluid α fetoprotein has been evaluated against a monoclonal antibody-based solid-phase immunoradiometric assay (IRMA) which has been the routine assay used by a Regional Amniotic Fluid Alphafetoprotein (AFP) Screening Laboratory for two years.

The DELFIA (LKB Wallac) AFP assay is a non-isotopic technique which utilizes a lanthanide metal (Europium) chelate label possessing highly fluorescent properties (Soini and Koijola [1]; Hemmila et al. [2]). The assay incorporates two monoclonal antibodies directed against two different immunogenic sites on the AFP molecule one of which is solid phase immobilized to the well of a microtitration strip.

The aim of this study was to assess the laboratory and clinical performance of this DELFIA.

Materials and methods

For μ fetoprotein analyses, amniotic fluid μ fetoprotein (AFP) was estimated by two techniques.

DELFIA AFP 1244-004 (LKB Wallac, PO Box 10, SF20101, Turku 10, Finland)

This product is a second-generation DELFIA AFP kit. AFP standards (range 1–1000 KU/l), QC and unknown samples (25 μl) are added to individual wells of microtitration strips (12 wells per strip, eight strips per plate) containing immobilized monoclonal antibody. The microtitration wells act as both reaction chambers and as measurement cuvettes. 200 μl of a europium labelled second monoclonal antibody is pipetted into the wells. The strips of microtitration wells are shaken vigorously on an automatic shaking device (Varishaker, Dynatech) for 2 min then incubated for 1 h at room temperature with continuous gentle shaking. The liquid contents of the wells are aspirated and the wells washed six times with wash buffer (plate-washing step is also the separation step), using an automatic plate-washer (LKB 1296-022). 200 μl of enhancement solution (forms a new chelate with the europium ion, thus amplifying the fluorescence) is added to each well and the strips shaken gently for 5 min. The microtitration strips are allowed to stand for 15 min and then the fluorescence measured on an automated LKB Wallac 1230 Arcus fluorometer with automated data handling. A microtitration plate of 96 wells can be fully processed with full print-out of patient results in 10 min. AFP results were calculated in kU/l corrected for the sample dilution factor and expressed as MU/l. The shelf life is reported to be at least five months (manufacturer’s information).

IRMA (SUCROSEP*) AFP IRMA, Boots-Celltech Diagnostics Ltd, Slough, UK

This immunoradiometric assay (IRMA) was performed using the protocol detailed by the manufacturer, α fetoprotein standard (range 1-25–712 kU/l) or unknown sample (50 μl) is incubated for 2 h at room temperature (15–30 °C) with 1251-labelled monoclonal antibody to α fetoprotein (100 μl) and solid phase anti α fetoprotein immunoglobulin (100 μl). During this incubation the assay tubes are shaken vigorously (300–350 min⁻¹) using an agitator (Sucroagitator, Boots-Celltech Diagnostics Ltd).

Following incubation the free and bound fractions are separated by the SUCROSEP sucrose layering non-centrifugation technique (Wright and Hunter [3]) using a semi-automatic instrument (Sucroseparator, Boots-Celltech Diagnostics Ltd). This allows assay tubes to be rapidly separated in 20-tube batches with a mean assay separation time of approximately 50 min.

The radioactivity in the assay tubes was counted on a multihead gamma counter (LKB 1260 multigamma 11) over a 120 s period. α fetoprotein values for clinical specimens were calculated as kU/l by spline function. The values were then corrected for the sample dilution factor and expressed as MU/l α fetoprotein.

* SUCROSEP is a trademark of Boots-Celltech Diagnostics Ltd.
Specimen analysis

Before analysis each amniotic fluid was diluted with a serum pool of zero α fetoprotein free serum. Normally a 1 in 200 dilution was used, but occasionally a higher dilution of the amniotic fluid sample was necessary. The batches for the two techniques were commenced on the same day with each amniotic fluid assayed in duplicate.

Amniotic fluid samples: 174 aliquots of amniotic fluid were investigated. These were collected by transabdominal amniocentesis during the second trimester of pregnancy for a variety of clinical indications. These samples were centrifuged and the supernatant either used immediately or stored at -20 °C until analysis.

Normal pregnancies: 115 liquors from normal pregnancies were investigated. Ultrasound examination of these pregnancies showed no obvious fetal abnormality. The amniotic fluid AFP by IRMA was within normal laboratory reference limits and there was no evidence of the acetyl cholinesterase isoenzyme band on polyacrylamide gel electrophoresis (Chubb et al. [4]; Smith et al. [5]).

Abnormal pregnancies: 55 samples were classified as 'abnormal'. These pregnancies had been terminated and the fetuses confirmed to have a severe fetal abnormality (27 open spina bifida, 28 anencephaly).

Equivocal pregnancies: Four samples were also studied from 'equivocal pregnancies' for which the outcome of the pregnancy was a normal infant but the amniotic fluid α fetoprotein concentration by IRMA was significantly elevated.

Statistical methods

Regression analysis was performed using standard techniques.

Results

Assay reproducibility

The intra- and inter-assay coefficients of variation (CV) for the DELFIA and IRMA were similar (table 1). The CV of the IRMA appeared to be better than the DELFIA at the highest AFP control pool concentration (sample 3).

The relationship between amniotic fluid AFP levels measured by DELFIA and IRMA

Regression analysis of results from 174 amniotic fluid samples (figure 1) falling in the range of AFP values from 3·4 to 584·0 MU/l analysed by both methods gave an excellent correlation coefficient of 0·997 (IRMA = 1·155 DELFIA + 0·946).

Table 1. Assay reproducibility of DELFIA and IRMA (intra- and inter-assay) based on repeated analyses of control pools. Prior to assays samples 1 and 2 were diluted 1/200, sample 3 was diluted 1/600.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Mean AFP concentration MU/l</th>
<th>Intra-assay %CV (N=10)</th>
<th>Inter-assay %CV (N=10)</th>
</tr>
</thead>
<tbody>
<tr>
<td>DELFIA</td>
<td>IRMA</td>
<td>DELFIA</td>
<td>IRMA</td>
</tr>
<tr>
<td>1</td>
<td>26·1</td>
<td>27·5</td>
<td>1·98</td>
</tr>
<tr>
<td>2</td>
<td>75·9</td>
<td>86·6</td>
<td>1·70</td>
</tr>
<tr>
<td>3</td>
<td>163·1</td>
<td>183·9</td>
<td>6·04</td>
</tr>
</tbody>
</table>

Table 2. α fetoprotein concentration of 115 amniotic fluids from normal pregnancies as measured by DELFIA.

<table>
<thead>
<tr>
<th>Gestational age</th>
<th>Number of samples</th>
<th>Range of AFP values (MU/l)</th>
<th>Median AFP concentration</th>
<th>Recommended cutoff [6] level of AFP (MU/l)</th>
</tr>
</thead>
<tbody>
<tr>
<td>15</td>
<td>16</td>
<td>8·3–20·8</td>
<td>13·8</td>
<td>32·5</td>
</tr>
<tr>
<td>16</td>
<td>20</td>
<td>7·2–15·6</td>
<td>11·85</td>
<td>35·6</td>
</tr>
<tr>
<td>17</td>
<td>18</td>
<td>5·8–21·6</td>
<td>9·9</td>
<td>29·7</td>
</tr>
<tr>
<td>18</td>
<td>20</td>
<td>3·6–12·6</td>
<td>8·15</td>
<td>24·5</td>
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<tr>
<td>19</td>
<td>20</td>
<td>2·8–12·0</td>
<td>5·1</td>
<td>17·9</td>
</tr>
<tr>
<td>20</td>
<td>21</td>
<td></td>
<td></td>
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</tbody>
</table>
In 53 of the 55 cases of open spina bifida and anencephaly the amniotic fluid AFP results as measured by DELFIA were greater than the multiple of the median cut-off limits. In two cases of open spina bifida the AFP level fell below the cut-off line (classified as false negative). The clinical classification of all 55 cases was exactly similar using either the DELFIA or the IRMA. In the remaining four amniotic fluids from the 'equivocal pregnancies' in which the outcome of the pregnancy was a normal infant, the AFP result as measured by DELFIA was elevated again, exactly similar to the results of the reference IRMA technique.

Discussion

Excellent agreement was shown between the amniotic fluid AFP results ($r = 0.997$) of the DELFIA and IRMA techniques. Furthermore the clinical performance of the two assays was exactly similar.

Both the DELFIA and IRMA are relatively quick techniques allowing a same day assay service if required. The DELFIA assay was more rapid, largely due to the need for a 1 h incubation period compared to a 2 h period for the IRMA.

The equipment requirements of the two assays were similar, i.e. orbital shakers and a separating system (a plate-washer or the Sucroseparator), followed by a radioactivity counter or a fluorimeter. Both techniques do not require centrifugation steps.

The DELFIA offers a number of analytical advantages, a longer shelf life of reagents than conventional radioimmunoassays and IRMAs, a greater working range (up to 1000 kU/l), a smaller sample volume and a non isotopic technique which therefore dispenses with the complications surrounding the safe handling and disposal of radioactive material.

In summary, the DELFIA proved to be a sensitive, convenient, easy to use non isotopic alternative to the measurement of amniotic fluid α fetoprotein in the routine clinical chemistry laboratory.

References
