Early prenatal investigation of a pregnancy at risk of adenosine deaminase deficiency using chorionic villi

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SUMMARY A pregnancy at risk for adenosine deaminase deficiency and severe combined immunodeficiency disease was investigated using samples of chorionic villi obtained during the eighth week of pregnancy. Adenosine deaminase levels suggested that the fetus was a probable carrier and that a diagnosis of severe combined immunodeficiency disease could be excluded. Enzyme and chromosome results were available within 24 hours of the chorionic villous sampling procedure, and were confirmed on amniotic fluid cell cultures after amniocentesis at 17 weeks’ gestation and on cord blood at delivery.

The basic biochemical defect in most forms of severe combined immunodeficiency disease (SCID) is unknown, but a deficiency of the purine catabolic pathway enzyme adenosine deaminase (ADA, E.C.3.5.4.4.) is found in just over 20% of cases with the autosomally inherited form of the disorder.1 Treatment of patients with SCID by enzyme replacement has been attempted2 but a successful bone marrow transplant remains the only effective therapy, an option which is available to fewer than 20% of patients who have an HLA compatible sibling.3 Because of the generally poor prognosis, most at risk couples request prenatal diagnosis and selective termination of affected fetuses. Where the ADA deficient form of SCID is identified, prenatal diagnosis can be made by assay of the enzyme in cultured amniotic fluid cells after amniocentesis at 16 to 18 weeks’ gestation,4 5 or by analysis of fetal blood samples by enzymology or fetal lymphocyte typing.6 7

We report here the early prenatal investigation of a pregnancy at risk for SCID and ADA deficiency using uncultured cells from chorionic villi obtained at eight weeks’ gestation. Within 24 hours of the procedure, a deficiency of ADA in the fetus was excluded and the patient reassured.

Case report

The patient, a 25 year old P1+1, gave birth to her first child in 1981. The child died aged 1 year with bronchopneumonia and leucopenia. A combined immunodeficiency was suspected and adenosine deaminase (ADA) activity was found to be absent in the erythrocytes. Cultured skin fibroblasts and red cells from both parents had intermediate levels of ADA confirming their carrier status. Four months later the patient conceived again and amniocentesis was arranged for 16 weeks’ gestation. When the pregnancy was 11 weeks advanced (by dates) the fetus was aborted spontaneously. Fetal cell cultures were found to have normal levels of ADA activity, but a 45,X/46.X,+7 chromosome constitution.

Before the couple’s third pregnancy the presence of ADA activity had been demonstrated in samples of chorionic villi obtained at early terminations. The parents elected to have the pregnancy monitored by this method with confirmatory amniocentesis at 16 weeks in the event of an unaffected pregnancy being predicted by examination of chorionic villi.

Methods

Samples of chorionic villi were obtained transcervically during the eighth week of pregnancy under direct vision by suction through an intravenous cannula attached to an endoscope (Olympus arthroscope). Three sampling attempts were made and villi obtained at the second and third attempts were collected in Ham’s F10. For ADA assay, three groups of villi (approximately 5 to 10 mg, free of blood and adhering maternal decidua) were selected using a dissection microscope. The remaining villi...
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were used for fetal sexing (Y fluorescence) and short term (24 hours) cultures for chromosome analysis. For the estimation of ADA activity, the villi were washed in 0.9% sodium chloride, resuspended in deionised water, and reduced to small fragments by chopping with scissors. The intracellular contents were released by sonication and the resulting suspension cleared by centrifugation at 10,000 g at 4°C for 10 minutes. The protein content of the supernatant was estimated by the Lowry method and adjusted to 0.3 to 0.5 mg/ml with deionised water.

Amniotic fluid cells were grown in Ham's F10 supplemented with 20% fetal calf serum and antibiotics. Cells were harvested by trypsinisation, washed in Hank's BSS and 0.9% sodium chloride, and the cell pellet resuspended in deionised water and disrupted by sonication. Protein content of the centrifuged supernatant was adjusted to 0.2 to 0.3 mg/ml before assay.

Packed red cells from cord blood collected in heparin were lysed with deionised water and the haemolysate further diluted to a haemoglobin concentration of 0.5 to 1.0 g/dl.

Lysates of chorionic villi, amniotic fluid cells or red cells were assayed for ADA activity under identical conditions, using a modification of the radioassay method described previously. A total of 5 μl of supernatant was mixed with 20 μl of 0.25 mol/l Tris-HCl, pH 7.4, containing 0.35 mmol/l 14C-adenosine. The reaction solution was covered with two drops of oil (paraffin:hexadecane, 80:20) to prevent evaporation and the tubes incubated at 37°C for one hour. The reaction was stopped by adding 5 μl of cold 20% trichloroacetic acid and the precipitated proteins removed by high speed centrifugation. The labelled products and unused substrate in 2 μl of the clear supernatant were separated by ascending thin layer chromatography on Polyethylenimine cellulose-F (PEI-F, Merck) with 5% Na2HPO4 as solvent. The inosine and adenosine spots were identified under short wave ultraviolet light, cut out, and the radioactivity quantified by heterogeneous liquid scintillation counting. Results were expressed as nmol adenosine deaminated per hour per mg of homogenate protein (chorionic villi and amniotic fluid cells) or nmol per hour per mg haemoglobin (red cells).

Results

ADA activity in the chorion biopsy from the pregnancy at risk and in 23 controls is presented in Table 1. Control villi were obtained at elective terminations between eight and 12 weeks’ gestation. Two of three bloodstained samples had raised levels of ADA activity reflecting contamination by the higher specific activity red cell enzyme. No difference in activity was found for samples assayed immediately after collection and after storage at −20°C for several weeks. ADA activity in chorionic villi from the pregnancy at risk was 50% of the mean of the control group, suggesting heterozygosity in the fetus.

Chromosome analysis after a 24 hour culture of cells from the chorion biopsy showed a normal male karyotype. The presence of a Y chromosome was confirmed by fluorescence analysis of interphase nuclei.

The results obtained in the chorionic villi were confirmed in AF cell cultures after amniocentesis at 17 weeks’ gestation and in cord blood at delivery (Table 2). Definite carrier status could not be ascribed to the infant from ADA activities determined in cultured AF cells or cord blood, since both tissues had activities which were just within the lower limits of the respective normal ranges.

Table 2. ADA activity in cultured amniotic fluid cells from the at risk pregnancy and in cord blood.

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<tr>
<th>ADA activity*</th>
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<tbody>
<tr>
<td>AF cell cultures</td>
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<tr>
<td>At risk pregnancy</td>
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<tr>
<td>Controls (n=9)</td>
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<tr>
<td>Range</td>
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<tr>
<td>Erythrocytes</td>
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<tr>
<td>Baby M (cord blood)</td>
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<td>Controls (venous blood)</td>
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<td>(n=22)</td>
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*Nmol/h/mg protein for AF cells, nmoVh/mg haemoglobin for red cells.
Discussion

The established method of prenatal diagnosis of chromosome abnormalities and most metabolic diseases depends upon culture of fetal cells obtained by amniocentesis after the 15th week of pregnancy. Considerable anxiety is experienced by parents during the period required for cell culture before results can be obtained (up to four weeks) and where a positive diagnosis is made there is the additional distress of a mid-trimester termination. Microscale methods of analysis which require only small numbers of cultured cells, and thus a correspondingly reduced cultivation period, have been applied to the prenatal diagnosis of a variety of metabolic diseases \( ^{11} \) including ADA deficiency, \( ^{10} \) and this approach can substantially reduce the period between amniocentesis and diagnosis. Alternatively, where the metabolic error is expressed in fetal blood, as is the case with ADA deficiency and SCID, sufficient material for direct analysis can be obtained by fetoscopy and fetal blood sampling. \( ^{6,7} \) However, these approaches, although providing a more rapid diagnosis, cannot be undertaken until 16 to 20 weeks' gestation owing to the need to obtain fetal tissue by amniocentesis or fetoscopy.

In the present communication we report the prenatal exclusion of ADA deficiency and SCID by the eighth week of pregnancy. Direct analysis on samples of chorionic villi eliminates the delay and uncertainty associated with long term cell culture and enzyme results were available within 24 hours of the sampling procedure, providing early reassurance for the patient. There was good agreement in ADA activity, measured as a percentage of the mean of the appropriate control group, between chorionic villi, cultured amniotic fluid cells, and cord blood. Thus, it is likely that the status of the fetus with respect to ADA activity is accurately reflected in samples of uncultured chorionic villi and that a deficiency of ADA activity could be readily identified, as has already been demonstrated for other inborn errors of metabolism. \( ^{12} \) Where intervention is indicated, termination of a pregnancy at eight to ten weeks is a safer, simpler, and much less traumatic experience for the patient than the corresponding procedure at 20 weeks' gestation.

As yet the risk to a continuing pregnancy after chorionic villi sampling cannot be accurately defined because of the limited experience obtained with a small number of diagnostic cases and the added complication of a high rate of spontaneous abortion in early pregnancy. However, provided the true risk is low and fetal tissue samples of adequate size uncontaminated with blood or maternal decidua can be obtained, then chorionic villi sampling is likely to become the method of choice for the prenatal diagnosis of a variety of metabolic diseases.

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References


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