Invasive Prenatal Diagnosis: Amniocentesis

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ABSTRACT

In this article, we will review the history and the evolution of the technique of amniocentesis and the indications of the most common invasive diagnostic and therapeutic procedure. Moreover, the most common complications of amniocentesis will be presented. Finally, we will try to establish if the use of concurrent ultrasound had any effect on the prevalence on these complications.

Keywords: Amniocentesis, Invasive prenatal diagnosis.


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HISTORY OF THE PROCEDURE

Amniocentesis is the first invasive procedure used in fetal medicine for both prenatal diagnosis and therapy. Although amniotic fluid withdrawal has been practiced for more than 150 years, the first reported cases of transabdominal evacuative amniocentesis were those of Prochownick et al in 1877, Schatz in 1882 and Hinkel in 1919 describing release of amniotic fluid from a patients with polyhydramnios. Menees et al reported removal of amniotic fluid by transabdominal needling using a radio-opaque contrast to outline the fetus and placenta.

At the beginning of the 1950s, it was used to determine the amniotic composition in cases of Rhesus isoimmunization and to correlate it with the severity of the condition of the newborn. Later on, Liley published the well-known correlation between the deviation of the spectral absorption curve of amniotic fluid resulting from bilirubin and the severity of rhesus isoimmunization. Since Liley’s studies, the practice of amniocentesis in pregnancies complicated by Rhesus disease was the standard procedure in obstetric practice until Mary described the use of middle cerebral artery (MCA) peak velocity in predicting fetal anemia and the need for in-utero blood transfusion in a noninvasive way.

Amniocentesis was used for diagnostic reasons in the ’50s, as a method for sex determination by the identification of Barr bodies in the noncultured amnion. About 10 years later, Steele and Breg reported in their paper in Lancet that the karyotype of the embryo was determined through an amniotic fluid cell culture. During the same year Thiede et al published similar findings later. The first case of prenatal diagnosis of Trisomy 21 (Down syndrome) was reported in 1968 by Nadler. In 1970, Nadler and Gerbie published the ‘Role of amniocentesis in the intrauterine diagnosis of genetic defects’ in the New England Journal of Medicine. This article was an innovation concerning genetic amniocentesis and diagnosis and since then genetic laboratories for analysis of amniotic fluid had become prevalent and indications for genetic amniocentesis included the detection of chromosomal abnormalities, gene disorders, X-linked conditions, inborn errors of metabolism, and the neural tube defects.

Amniocentesis has been established as a basic invasive method for the prenatal diagnosis of various pregnancy related conditions, such as fetal karyotyping, diagnosis of metabolic or enzymatic diseases, assessment of the severity hemolytic disease, establishment of lung maturity, diagnosis of fetal infections. Additionally, amniocentesis is used for the infusion of various drugs into the amniotic cavity, determination of the composition of the amniotic fluid and finally for evacuation of hydramnion.

EVOLUTION OF THE TECHNIQUE

During the 30s removal of amniotic was done by transabdominal needling following injection of a radio-opaque contrast in order to outline the fetus and placenta. (Menees et al reported in 1930). Later on amniocenteses were performed ‘blindly’ and the puncture site located merely by external palpation of the uterus in the abdomen. In 1967, Hofmann and Hollander in Germany stated the importance of placental localization using ultrasound before amniocentesis. Amniocentesis under ultrasound guidance was started to be implemented in 1972, with reports from Bang and Northeved from Hans Hendrik Holm at the Gentofte Hospital in Copenhagen. In the mid 1970s to mid 80s, amniocentesis was performed with the assistance of a static or realtime B-scan. A scan was first performed to locate a feasible pocket of amniotic fluid for aspiration.
fluid before a tap, the skin on top of that area was marked and the puncture was done without actually seeing the needle tip going into the fluid pocket.

Improvement of ultrasound real time scanners in the late 1970s, a small number of centers started to perform amniocentesis with the simultaneous visualization of the puncture needle tip on the scanner monitor. One such pioneer was the Birnholz group at Harvard who used an early phased array for the purpose. Needle-guide adapters soon became available from ultrasound manufacturers which could be coupled to the linear array or phased array sector probes where the needle passed through a fixed path either parallel or at an angle to the ultrasonic beam. These were difficult to use, however, particularly in a busy setting. They also had serious problem of keeping the equipment sterile. The adapters may also increase the risk of traumatization as it did not allow for the ‘desired’ and sensitive placement of needles.

Many centers started to do a freehand technique with an assistant holding onto the transducer probe that was commonly wrapped in a sterile adhesive drape. In 1984, Holzgreve in Basel, Switzerland described a large series of over 3000 ‘freehand’ amniocenteses with low complication rate. Similar experience was also reported by Platt in Los Angeles, who emphasized on the need for the transducer probe to be manipulated by the same operator which resulted in better hand-eye co-ordination. In the following year, Romero formalized the single operator two-hands technique in amniocentesis and the reduction in the number of multiple taps and bloody taps associated with the procedure. Most centers soon adopted the single operator technique, which had become popular because of its convenience and effectiveness. Newer needles were marketed with special external coating and echoluminance to enhance needle placement. Complication rates were reportedly lower with each successive improvement in technique.

INDICATIONS

Amniocentesis is used for both prenatal diagnosis and fetal therapy. In Table 1, we summarize the main indications for amniocentesis.

COMPLICATIONS OF AMNIOCENTESIS

During amniocentesis, the fetomaternal unit is injured, and thus a number of complications may occur. Maternal complications are rare. They include perforation of the intra-abdominal viscera with subsequent intra-abdominal infection, sepsis, bleeding, blood group sensitization and uterine contractions. The use of ultrasound guidance during amniocentesis has minimized the risk of maternal injury. Also aseptic technique and the use of anti-D immunoglobulin has eliminated the risk of maternal sepsis and rhesus sensitization.

Fetal complications, though, are the main concern. They include fetal loss, placental abruption, preterm labor and preterm rupture of membranes. Needle puncture injuries of the fetus and injury due to withdrawal of amniotic fluid (e.g. amniotic bands) were rare especially since amniocentesis is being performed under ultrasound guidance. Amniocentesis may also cause intra-amniotic infections through the introduction of microorganisms into the amniotic cavity via the needle. In order to eliminate the possibility of an infectious complication, rules of aseptic procedures which apply in all surgical procedures must be applied during amniocentesis, such as the aseptic cleaning of the skin and using sterilized medical gloves and needles. However, endometrial infections are not always related to amniocentesis, but may exist before the prenatal intervention.

FETAL LOSS

Fetal loss is the ultimate risk of genetic amniocentesis. When amniocentesis was first introduced in the clinical practice the risk of miscarriage due to the procedure could not be estimated accurately because there was lack of ultrasound guidance and lack of determination of fetal viability before the procedure.

Since fetal miscarriage does not occur only in association with amniocentesis, the background loss rate which is associated with the gestation age, parity and any other underlying risk factors is important to be
determined before estimating the procedure loss rate. For instance gestational age at the procedure is an important determinant of the observed fetal loss rate, since the earlier the pregnancy the greater is the pre-procedure risk of miscarriage.

The risk of fetal loss after amniocentesis has been evaluated by several series from single centers and a number of multicenter studies. The controversial results of these studies merely imply the difficulties in evaluating the procedure related loss rate and the changes in practice of the procedure.

In 1976, the National Institute of Child Health and Human Development (NICHD) published a case-control, cohort study to evaluate the safety and accuracy of second trimester amniocentesis for prenatal diagnosis. The authors reported that the group who underwent amniocentesis had a 3.5% total loss rate while the controls had 3.2%. The authors of this study also reported that the procedure loss risk was unrelated to previous pregnancy loss, volume of aspirated fluid or the number of attempts.

During the same year, the Canadian Collaborative Study presented an descriptive paper in which the total loss rate after amniocentesis was 3.2%. In this study, an increased risk of loss was reported in cases of failed attempts and large bore needles (>19 gauge). The third collaborative case control study was reported by the British Working Party on Amniocentesis in 1978. The authors concluded that the risk for fetal loss after amniocentesis was 1.2%. This risk was estimated since the total fetal loss before 28 weeks was 2.4% in cases which underwent amniocentesis and 1.2% in cases without amniocentesis (controls). This study, however, was strongly criticized mainly on the selection of control individuals who were recruited in the study later in gestation and on the replacement of some of the matched controls who had aborted.

The only randomized trial for the estimation of miscarriage risk related to amniocentesis was done in Denmark in 1986. This study randomized 4,606 low-risk women aged 24 to 35 years to have or not to have an amniocentesis, which was carried out using a 20-gauge needle under real-time ultrasound guidance. Most procedures were performed between 16 and 18 weeks of gestation. The amniocentesis group had a loss rate which exceeded the control group by 1% (1.7% and 0.7% respectively), a figure which has often been used in counseling couples undergoing amniocentesis. This study has also been criticized due to the very low loss rate in the controls.

Since then a lot of studies reported amniocentesis related fetal loss rate between 0.2 to 0.9%. In 2001, JW Seeds performed a systematic review of 29 controlled and uncontrolled studies with a total of 68,119 amniocenteses. In this review, Seeds included both reports that described amniocentesis with only preprocedure ultrasound and reports which described amniocentesis with concurrent use of ultrasound guidance.

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The goal of this review study was to estimate the procedure-related risk of pregnancy loss after mid trimester amniocentesis. The impact of concurrent ultrasound guidance on this fetal loss and fetal injuries were also examined.

He concluded that amniocentesis with concurrent ultrasound guidance in controlled studies appears to be associated with a procedure-related rate of excess pregnancy loss of 0.6% (95% CI, 0.31, 0.90). When comparing all studies, the risk of pregnancy loss after the procedure and before 28 weeks was eliminated with the use of concurrent real time ultrasound. When only controlled studies were included this trend remained although not significant.

In 2006, Eddleman et al in order to quantify the procedure-related loss rate after midtrimester amniocentesis a database generated from patients who were recruited to the first and second trimester evaluation of risk for aneuploidy (FASTER) trial was used. A total of 35,003 unselected patients from the general population with viable singleton pregnancies were enrolled in this trial and 3.0% of them underwent midtrimester amniocentesis. The total spontaneous fetal loss rate less than 24 weeks of gestation in the study group was 1.0% and was not statistically different from the background 0.94% rate seen in the control group (P—0.74, 95% CI—0.26, 0.49). The researchers stated that the odds of pregnancy loss were actually lower in patients who underwent amniocentesis because of advanced maternal age or screen positive results compared with those who did not. They concluded that this is likely due to the fact that spontaneous pregnancy loss is so strongly associated with aneuploidy, and patients who have an amniocentesis would presumably terminate aneuploid fetuses in most cases before a spontaneous loss could occur. They stated that it is not surprising that patients in these categories who elected not to undergo amniocentesis, had a higher rate of spontaneous loss because they were carrying a higher proportion of aneuploid fetuses.

Mujezinovic F and Alfirevic Z reviewed a number of studies in order to provide data for counseling concerning complications after amniocentesis and chorionic villus sampling (CVS). Twenty-nine articles of amniocentesis were included. Total pregnancy loss within 14 days was estimated to be 0.6% [95% confidence interval (CI) 0.5–0.7], rising to 0.9% (95% CI 0.6–1.3) for pregnancy loss before 24 weeks and 1.9% (95% CI 1.4–2.5) for total pregnancy loss. However, the authors noted that since most of the reviewed studies do not take into account the background
Vaginal bleeding during current pregnancy increase the risk of spontaneous abortion and pregnancy loss after amniocentesis in most of the series. In our recent series, we also found that bleeding during first and early second trimester of current pregnancy increase the risk of pregnancy loss after amniocentesis by two to three times.

Finally, factors associated with the procedure itself may alter the risk after amniocentesis. The number of needle placements, the aspiration of bloody fluid, and especially the observation of green or murky fluid are seen to be associated with a significantly increased risk of pregnancy loss after amniocentesis.

The number of needle insertions and bloody fluid are related directly to procedure technique, whereas murky or green fluid is not. Murky of green fluid is usually the result of previous intra-amniotic hemorrhage and this is the reason why it is definitely increasing the risk of fetal loss after amniocentesis.

In our series, when the aspired amniotic fluid was found to be green or brownish blood-stained indicating an old intra-amniotic hemorrhage, the risk of fetal loss is almost six times higher. On the contrary, in cases where amniotic fluid was mixed with fresh blood [e.g. in cases of placenta perforation (Fig. 1)] the fetal loss was not higher than cases when clear amniotic fluid was aspirated. Additionally, the risk of fetal loss is increased when more than one needle insertion in needed although the difference was not statistically significant.

Transplacental needle placement (Fig. 2) was found by some authors, but not others, to be associated with an increased risk of pregnancy loss.

The use of concurrent ultrasound help the operator to reduce the number of needle insertions, reduce bloody amniotic fluid by avoiding placenta insertion (Fig. 3) and aspiration and avoid fibromas. In this way it may have played a role in reducing the risk of fetal loss after amniocentesis.

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**Fig. 1:** Intra-amniotic bleeding after placental perforation (transplacental amniocentesis)
amniocentesis group and 9.1% for the control group, this difference was not statistically significant (p = 0.223).

**FETAL TRAUMA**

Theoretically, the risk of direct fetal needle injury should be reduced with the use of simultaneous ultrasound guidance. Nevertheless several case reports document or describe cases of fetal injury to the amniocentesis needle, despite the use of simultaneous guidance for the procedure.40-51 This association though is not based on direct evidence. In addition, there are reports of fetal skin marks in cases without amniocentesis and these suggest that these marks may not be the result of needle puncture of the fetus.52

**ALLOIMMUNIZATION AND VIRAL DISEASE TRANSMISSION**

Fetomaternal hemorrhage occurs in approximately 50% of all women53 and during amniocentesis on one out of six.54 The attributed risk is 1% greater than the background risk of 1.5%.55 As for preventing the Rhesus sensitization of the mother and when the husband is Rhesus positive, every Rhesus negative mother must be given 300 μg anti-D Immunoglobulin after amniocentesis, provided of course the indirect Coombs is negative.56,57 Other means to minimize the risk is the use of small gauge needles and to avoid transplacental approach.57

In the same way, HIV and hepatitis virus positive pregnant women are safe to undergo amniocentesis provided that viral load of patients is low and the transplacental route is avoided.58

The use of real time ultrasound guidance during the procedure helps the operator to avoid the transplacental route which is crucial in cases of Rhesus negative mothers and HIV or hepatitis virus positive pregnant women.

**CONCLUSION**

Real time ultrasound guidance during amniocentesis has reduced the number of needle insertion and the aspiration of bloody amniotic fluid. It has also reduced the cases of placental puncture which although does not associate with increased risk of pregnancy loss, it increases the risk of Rhesus sensitization and transmission of viral diseases (HIV, Hepatitis) for mother to the fetus.

Using concurrent ultrasound guidance seems to decrease the risk of fetal loss after amniocentesis although this decrease is not statistically significant in all series. Finally, the risk of direct fetal trauma is also eliminated when real time ultrasound is used during the needle insertion.
REFERENCES